42
Parts 400 to 429
Revised as of October 1, 2000

Public Health

Containing a Codification of documents
of general applicability and future effect

As of October 1, 2000

With Ancillaries

Published by
Office of the Federal Register
National Archives and Records Administration

As a Special Edition of the Federal Register
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To cite the regulations in this volume use title, part and section number. Thus, 42 CFR 400.200 refers to title 42, part 400, section 200.
Explanation

The Code of Federal Regulations is a codification of the general and permanent rules published in the Federal Register by the Executive departments and agencies of the Federal Government. The Code is divided into 50 titles which represent broad areas subject to Federal regulation. Each title is divided into chapters which usually bear the name of the issuing agency. Each chapter is further subdivided into parts covering specific regulatory areas.

Each volume of the Code is revised at least once each calendar year and issued on a quarterly basis approximately as follows:

Title 1 through Title 16..............................................................as of January 1
Title 17 through Title 27.................................................................as of April 1
Title 28 through Title 41.............................................................as of July 1
Title 42 through Title 50.............................................................as of October 1

The appropriate revision date is printed on the cover of each volume.

LEGAL STATUS

The contents of the Federal Register are required to be judicially noticed (44 U.S.C. 1507). The Code of Federal Regulations is prima facie evidence of the text of the original documents (44 U.S.C. 1510).

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To determine whether a Code volume has been amended since its revision date (in this case, October 1, 2000), consult the “List of CFR Sections Affected (LSA),” which is issued monthly, and the “Cumulative List of Parts Affected,” which appears in the Reader Aids section of the daily Federal Register. These two lists will identify the Federal Register page number of the latest amendment of any given rule.

EFFECTIVE AND EXPIRATION DATES

Each volume of the Code contains amendments published in the Federal Register since the last revision of that volume of the Code. Source citations for the regulations are referred to by volume number and page number of the Federal Register and date of publication. Publication dates and effective dates are usually not the same and care must be exercised by the user in determining the actual effective date. In instances where the effective date is beyond the cutoff date for the Code a note has been inserted to reflect the future effective date. In those instances where a regulation published in the Federal Register states a date certain for expiration, an appropriate note will be inserted following the text.

OMB CONTROL NUMBERS

The Paperwork Reduction Act of 1980 (Pub. L. 96-511) requires Federal agencies to display an OMB control number with their information collection request.
Many agencies have begun publishing numerous OMB control numbers as amendments to existing regulations in the CFR. These OMB numbers are placed as close as possible to the applicable recordkeeping or reporting requirements.

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Provisions that become obsolete before the revision date stated on the cover of each volume are not carried. Code users may find the text of provisions in effect on a given date in the past by using the appropriate numerical list of sections affected. For the period before January 1, 1986, consult either the List of CFR Sections Affected, 1949-1963, 1964-1972, or 1973-1985, published in seven separate volumes. For the period beginning January 1, 1986, a “List of CFR Sections Affected” is published at the end of each CFR volume.

INCORPORATION BY REFERENCE

What is incorporation by reference? Incorporation by reference was established by statute and allows Federal agencies to meet the requirement to publish regulations in the Federal Register by referring to materials already published elsewhere. For an incorporation to be valid, the Director of the Federal Register must approve it. The legal effect of incorporation by reference is that the material is treated as if it were published in full in the Federal Register (5 U.S.C. 552(a)). This material, like any other properly issued regulation, has the force of law.

What is a proper incorporation by reference? The Director of the Federal Register will approve an incorporation by reference only when the requirements of 1 CFR part 51 are met. Some of the elements on which approval is based are:

(a) The incorporation will substantially reduce the volume of material published in the Federal Register.

(b) The matter incorporated is in fact available to the extent necessary to afford fairness and uniformity in the administrative process.

(c) The incorporating document is drafted and submitted for publication in accordance with 1 CFR part 51.

Properly approved incorporations by reference in this volume are listed in the Finding Aids at the end of this volume.

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An index to the text of “Title 3—The President” is carried within that volume.

The Federal Register Index is issued monthly in cumulative form. This index is based on a consolidation of the “Contents” entries in the daily Federal Register.

A List of CFR Sections Affected (LSA) is published monthly, keyed to the revision dates of the 50 CFR titles.
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For a legal interpretation or explanation of any regulation in this volume, contact the issuing agency. The issuing agency's name appears at the top of odd-numbered pages.

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RAYMOND A. MOSLEY,
Director,
Office of the Federal Register.

October 1, 2000.
THIS TITLE

Title 42—Public Health is composed of three volumes. The parts in these volumes are arranged in the following order: Parts 1-399, parts 400-429 and part 430 to end. The first volume (parts 1-399) contains current regulations issued under chapter I—Public Health Service (HHS). The second volume (parts 400-429) includes regulations issued under chapter IV—Health Care Financing Administration (HHS) and the third volume (part 430 to end) contains the remaining regulations in chapter IV and the regulations issued under chapter V by the Office of Inspector General—Health Care (HHS). The contents of these volumes represent all current regulations codified under this title of the CFR as of October 1, 2000.

The OMB control numbers for the Health Care Financing Administration appear in §400.310 of chapter IV. For the convenience of the user subpart C consisting of §§400.300-400.310 is reprinted in the Finding Aids section of the third volume.

Redesignation tables appear in the Finding Aids section of all volumes.

For this volume, Linda L. Jones was Chief Editor. The Code of Federal Regulations publication program is under the direction of Frances D. McDonald, assisted by Alomha S. Morris.
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Title 42—Public Health

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## CHAPTER IV—HEALTH CARE

FINANCING ADMINISTRATION,

DEPARTMENT OF HEALTH AND HUMAN SERVICES


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PART 400—INTRODUCTION; DEFINITIONS

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Subpart B—Definitions

Sec.
400.200 General definitions.
400.202 Definitions specific to Medicare.
400.203 Definitions specific to Medicaid.

Subpart C—OMB Control Numbers for Approved Collections of Information

400.300 Scope.
400.310 Display of currently valid OMB control numbers.

AUTHORITY: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh) and 44 U.S.C. Chapter 35.

Subpart A—[Reserved]

Subpart B—Definitions

§ 400.200 General definitions.

In this chapter, unless the context indicates otherwise—

Act means the Social Security Act, and titles referred to are titles of that Act.

Administrator means the Administrator, Health Care Financing Administration.

ALJ stands for administrative law judge.

Area means the geographical area within the boundaries of a State, or a State or other jurisdiction, designated as constituting an area with respect to which a Professional Standards Review Organization or a Utilization and Quality Control Peer Review Organization has been or may be designated.

CMP stands for competitive medical plan.

Conditions of participation includes requirements for participation as the latter term is used in part 483 of this chapter.

Condition level deficiencies includes deficiencies with respect to "level A requirements" as the latter term is used in parts 442 and 483 of this chapter.

CORF stands for comprehensive outpatient rehabilitation facility.


CY stands for calendar year.

DAB stands for Departmental Appeals Board.

Department means the Department of Health and Human Services (HHS), formerly the Department of Health, Education, and Welfare.

ESRD stands for end-stage renal disease.

FDA stands for the Food and Drug Administration.

FQHC means Federally qualified health center.

FR stands for Federal Register.

FY stands for fiscal year.

HCFA stands for Health Care Financing Administration.

HCPP stands for health care prepayment plan.

HHS stands for the Department of Health and Human Services.

HHA stands for home health agency.

HMO stands for health maintenance organization.

ICF stands for intermediate care facility.

ICF/MR stands for intermediate care facility for the mentally retarded.

Medicaid means medical assistance provided under a State plan approved under title XIX of the Act.

Medicare means the health insurance program for the aged and disabled under title XVIII of the Act.

NCD stands for national coverage determination.

OASDI stands for the Old Age, Survivors, and Disability Insurance program under title II of the Act.

OIG stands for the Department’s Office of the Inspector General.

Peer review organization means an organization that has a contract with HCFA, under part B of title XI of the Act, to perform utilization and quality control review of the health care furnished, or to be furnished, to Medicare beneficiaries.

PRO stands for peer review organization.

QDWI stands for Qualified Disabled and Working Individual.

QMB stands for Qualified Medicare Beneficiary.
§ 400.202 Definitions specific to Medicare.

As used in connection with the Medicare program, unless the context indicates otherwise—

Beneficiary means a person who is entitled to Medicare benefits.

Carrier means an entity that has a contract with HCFA to determine and make Medicare payments for Part B benefits payable on a charge basis and to perform other related functions.

Critical access hospital (CAH) means a facility designated by HFCA as meeting the applicable requirements of section 1820 of the Act and of subpart F of part 485 of this chapter.

Entitled means that an individual meets all the requirements for Medicare benefits.

Essential access community hospital (EACH) means a hospital designated by HCFA as meeting the applicable requirements of section 1820 of the Act and of subpart G of part 412 of this chapter, as in effect on September 30, 1997.

GME stands for graduate medical education.

Hospital insurance benefits means payments on behalf of, and in rare circumstances directly to, an entitled individual for services that are covered under Part A of title XVIII of the Act.

Intermediary means an entity that has a contract with HCFA to determine and make Medicare payments for Part A or Part B benefits payable on a cost basis and to perform other related functions.

Medicare Part A means the hospital insurance program authorized under Part A of title XVIII of the Act.

Medicare Part B means the supplementary medical insurance program authorized under Part B of title XVIII of the Act.

National coverage determination (NCD) means a national policy determination

Qualified Disabled and Working Individual means an individual who—

1. Is eligible to enroll for Medicare Part A under section 1818A of the Act.

2. Has income, as determined in accordance with SSI methodologies, that does not exceed 200 percent of the Federal poverty guidelines (as defined and revised annually by the Office of Management and Budget) for a family of the size of the individual’s family;

3. Has resources, as determined in accordance with SSI methodologies, that do not exceed twice the maximum amount established for SSI eligibility, for an individual or for an individual and his or her spouse; and

4. Is not otherwise eligible for Medicaid.

Qualified Medicare Beneficiary means an individual who—

1. Is entitled to Medicare Part A, with or without payment of premiums, but is not entitled solely because he or she is eligible to enroll as a QDWI;

2. Has resources, as determined in accordance with SSI methodologies, that do not exceed twice the maximum amount established for SSI eligibility; and

3. Has income, as determined in accordance with SSI methodologies, that does not exceed 100 percent of the Federal poverty guidelines.

Regional Administrator means a Regional Administrator of HCFA.

Regional Office means one of the regional offices of HCFA.

RHC stands for rural health clinic.

RRB stands for Railroad Retirement Board.

Secretary means the Secretary of Health and Human Services.

SNF stands for skilled nursing facility.

Social security benefits means monthly cash benefits payable under section 202 or 223 of the Act.

SSA stands for Social Security Administration.

United States means the fifty States, the District of Columbia, the Commonwealth of Puerto Rico, the Virgin Islands, Guam, American Samoa, and the Northern Mariana Islands.


regarding the coverage status of a particular service, that HCFA makes under section 1862(a)(1) of the Act, and publishes as a FEDERAL REGISTER notice or HCFA Ruling. (The term does not include coverage changes mandated by statute.)

Nonparticipating supplier means a supplier that does not have an agreement with HCFA to participate in Part B of Medicare in effect on the date of the service.

Participating supplier means a supplier that has an agreement with HCFA to participate in Part B of Medicare in effect on the date of the service.

Payment on an assignment-related basis means payment for Part B services—

(1) To a physician or other supplier that accepts assignment from the beneficiary, in accordance with §424.55 or §424.56 of this chapter;

(2) To a physician or other supplier after the beneficiary’s death, in accordance with §424.64(c)(1) of this chapter; or

(3) To an entity that pays the physician or other supplier under a health benefit plan, in accordance with §424.66 of this chapter.

Provider means a hospital, a CAH, a skilled nursing facility, a comprehensive outpatient rehabilitation facility, a home health agency, or a hospice that has in effect an agreement to participate in Medicare, or a clinic, a rehabilitation agency, or a public health agency that has in effect a similar agreement but only to furnish outpatient physical therapy or speech pathology services, or a community mental health center that has in effect a similar agreement but only to furnish partial hospitalization services.


Services means medical care or services and items, such as medical diagnosis and treatment, drugs and biologicals, supplies, appliances, and equipment, medical social services, and use of hospital, CAH, or SNF facilities.

Supplementary medical insurance benefits means payment to or on behalf of an entitled individual for services covered under Part B of title XVIII of the Act.

Supplier means a physician or other practitioner, or an entity other than a provider, that furnishes health care services under Medicare.


§ 400.203 Definitions specific to Medicaid.

As used in connection with the Medicaid program, unless the context indicates otherwise—

Applicant means an individual whose written application for Medicaid has been submitted to the agency determining Medicaid eligibility, but has not received final action. This includes an individual (who need not be alive at the time of application) whose application is submitted through a representative or a person acting responsibly for the individual.

Federal financial participation (FFP) means the Federal Government’s share of a State’s expenditures under the Medicaid program.

FMAP stands for the Federal medical assistance percentage, which is used to calculate the amount of Federal share of State expenditures for services.

Medicaid agency or agency means the single State agency administering or supervising the administration of a State Medicaid plan.

Nursing facility (NF), effective October 1, 1990, means an SNF or an ICF participating in the Medicaid program.

Provider means any individual or entity furnishing Medicaid services under a provider agreement with the Medicaid agency.

Recipient means an individual who has been determined eligible for Medicaid.

Services means the types of medical assistance specified in section 1905(a) of the Act and defined in subpart A of part 440 of this chapter.
§ 400.300 Display of currently valid OMB control numbers.

**State means the several States, the District of Columbia, the Commonwealth of Puerto Rico, the Virgin Islands, Guam, American Samoa and the Northern Mariana Islands.**

State plan or the plan means a comprehensive written commitment by a Medicaid agency, submitted under section 1902(a) of the Act, to administer or supervise the administration of a Medicaid program in accordance with Federal requirements.


Subpart C—OMB Control Numbers for Approved Collections of Information

**SOURCE:** 49 FR 4477, Feb. 7, 1984, unless otherwise noted.

§ 400.300 Scope.

This subpart collects and displays control numbers assigned by the Office of Management and Budget (OMB) to collections of information contained in HCFA regulations, in accordance with OMB’s regulations for controlling paperwork burdens on the public, 5 CFR part 1320. HCFA intends that the subpart comply with the requirements of section 3507(f) of the Paperwork Reduction Act of 1980, 44 U.S.C. chapter 35 which requires that agencies shall not engage in a “collection of information,” without obtaining a control number from OMB.

§ 400.310 Display of currently valid OMB control numbers.

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§ 401.101 Purpose and scope.

(a) The regulations in this subpart:

(1) Implement section 1106(a) of the Social Security Act as it applies to the Health Care Financing Administration (HCFA). The rules apply to information obtained by officers or employees of HCFA in the course of administering title XVIII of the Social Security Act (Medicare), information obtained by Medicare intermediaries or carriers in the course of carrying out agreements...
§ 401.102 Definitions.

For purposes of this subpart:

Act means the Social Security Act.

Freedom of Information Act rules mean the substantive mandatory disclosure provisions of the Freedom of Information Act, 5 U.S.C. 552 (including the exemptions from mandatory disclosure, 5 U.S.C. 552(b), as implemented by the Department's public information regulation, 45 CFR part 5, subpart F and by §§ 401.106 to 401.152 of this subpart.

Person means a person as defined in the Administrative Procedure Act, 5 U.S.C. 551(2). This includes State or local agencies, but does not include Federal agencies or State or Federal courts.

Record has the same meaning as that provided in 45 CFR 5.5.

Subject individual means an individual whose record is maintained by the Department in a system of records, as the terms "individual," "record," and "system of records" are defined in the Privacy Act of 1974, 5 U.S.C. 552a(a).

§ 401.105 Rules for disclosure.

(a) General rule. The Freedom of Information Act rules shall be applied to every proposed disclosure of information. If, considering the circumstances of the disclosure, the information would be made available in accordance with the Freedom of Information Act rules, then the information may be disclosed regardless of whether the requester or recipient of the information has a statutory right to request the information under the Freedom of Information Act, 5 U.S.C. 552, or whether a request has been made.

(b) Application of the general rule. Pursuant to the general rule in paragraph (a) of this section,

(1) Information shall be disclosed—

(i) To a subject individual when required by the access provision of the Privacy Act, 5 U.S.C. 552a(d), as implemented by the Department Privacy Act regulation, 45 CFR part 5b; and

(ii) To a person upon request when required by the Freedom of Information Act, 5 U.S.C. 552;

(2) Unless prohibited by any other statute (e.g., the Privacy Act of 1974, 5 U.S.C. 552a(b), the Tax Reform Act of 1976, 26 U.S.C. 6103, or section 1106(d) and (e) of the Social Security Act), information may be disclosed to any requester or recipient of the information, including another Federal agency or a State or Federal court, when the information would not be exempt from mandatory disclosure under Freedom of Information Act rules or when the information nevertheless would be made available under the Department's public information regulation's criteria for disclosures which are in the public interest and consistent with obligations of confidentiality and administrative necessity, 45 CFR part 5, subpart F, as supplemented by §§ 401.106 to 401.152 of this subpart.

§ 401.106 Publication.
(a) Methods of publication. Materials required to be published under the provisions of The Freedom of Information Act, 5 U.S.C. 552 (a)(1) and (2) are published in one of the following ways:
(1) By publication in the Federal Register of HCFA regulations, and by their subsequent inclusion in the Code of Federal Regulations;
(2) By publication in the Federal Register of appropriate general notices;
(3) By other forms of publication, when incorporated by reference in the Federal Register with the approval of the Director of the Federal Register; and
(4) By publication of indexes of precedential orders and opinions issued in the adjudication of claims, statements of policy and interpretations which have been adopted but have not been published in the Federal Register, and of administrative staff manuals and instructions to staff that affect a member of the public.
(b) Availability for inspection. Those materials which are published in the Federal Register pursuant to 5 U.S.C. 552(a)(1) shall, to the extent practicable and to further assist the public, be made available for inspection at the places specified in § 401.128.
§ 401.108 HCFA rulings.
(a) After September 1981, a precedent final opinion or order or a statement of policy or interpretation that has not been published in the Federal Register as a part of a regulation or of a notice implementing regulations, but which has been adopted by HCFA as having precedent, may be published in the Federal Register as a HCFA Ruling and will be made available in the publication entitled HCFA Rulings.
(b) Precedent final opinions and orders and statements of policy and interpretation that were adopted by HCFA before October, 1981, and that have not been published in the Federal Register are available in HCFA Rulings.
(c) HCFA Rulings are published under the authority of the Administrator, HCFA. They are binding on all HCFA Components, and on the Social Security Administration to the extent that components of the Social Security Administration adjudicate matters under the jurisdiction of HCFA.
[48 FR 22924, May 23, 1983]
§ 401.110 Publications for sale.
The following publications containing information pertaining to the program, organization, functions, and procedures of HCFA may be purchased from the Superintendent of Documents, Government Printing Office, Washington, DC 20402.
(a) Titles 20, 42, and 45 of the Code of Federal Regulations.
(b) Federal Register issues.
(c) Compilation of the Social Security Laws.
(d) HCFA Rulings.
(e) Social Security Handbook. The information in the Handbook is not of precedent or interpretative force.
(f) Medicare/Medicaid Directory of Medical Facilities.
§ 401.112 Availability of administrative staff manuals.
All HCFA administrative staff manuals and instructions to staff personnel which contain policies, procedures, or interpretations that affect the public are available for inspection and copying. A complete listing of such materials is published in HCFA Rulings. These manuals are generally not printed in a sufficient quantity to permit sale or other general distribution to the public. Selected material is maintained at Social Security Administration district offices and field offices and may be inspected there. See §§ 401.130 and 401.132 for a listing of this material.
§ 401.116 Availability of records upon request.
(a) General. In addition to the records made available pursuant to §§ 401.106, 401.108, 401.110 and 401.112, HCFA will, upon request made in accordance with this subpart, make identified records available to any person, unless they are exempt from disclosure under the provisions of section 552(b) of title 5, United States Code (see § 401.126), or any other provision of law.
§ 401.118 Misappropriation, alteration, or destruction of records.

No person may remove any record made available to him for inspection or copying under this part, from the place where it is made available. In addition, no person may steal, alter, mutilate, obliterate, or destroy in whole or in part, such a record. See sections 641 and 2071 of title 18 of the United States Code.

§ 401.118 Deletion of identifying details.

When HCFA publishes or otherwise makes available an opinion or order, statement of policy, or other record which relates to a private party or parties, the name or names or other identifying details will be deleted.

§ 401.120 Creation of records.

Records will not be created by compiling selected items from the files, and records will not be created to provide the requester with such data as ratios, proportions, percentages, per capita, frequency distributions, trends, correlations, and comparisons. If such data have been compiled and are available in the form of a record, the record shall be made available as provided in this subpart.

§ 401.126 Information or records that are not available.

(a) Specific exemptions from disclosure. Pursuant to paragraph (b) of 5 U.S.C. 552, certain classes of records are exempt from disclosure. For some examples of the kinds of materials which are exempt, see subpart F of the public information regulation of the Department of Health and Human Services (45 CFR part 5) and the appendix to that regulation.

(b) Materials exempt from disclosure by statute. Pursuant to paragraph (b)(3) of 5 U.S.C. 552, as amended, which exempts from the requirement for disclosure matters that are exempted from disclosure by statute, provided that such statute requires that the matters be withheld from the public in such a manner as to leave no discretion on the issue, or establishes particular criteria for withholding or refers to particular types of matter to be withheld.

(1) Reports described in sections 1106 (d) and (e) of the Social Security Act shall not be disclosed, except in accordance with the provisions of sections 1106 (d) and (e). Sections 1106 (d) and (e) provide for public inspection of certain official reports dealing with the operation of the health programs established by titles XVIII and XIX of the Social Security Act (Medicare and Medicaid), but require that program validation survey reports and other formal evaluations of providers of services shall not identify individual patients, individual health care practitioners, or other individuals. Section 1106(e) further requires that none of the reports shall be made public until the contractor or provider whose performance is being evaluated has had a reasonable opportunity to review that report and to offer comments. See §401.133 (b) and (c).

(2)(i) Except as specified in paragraph (b)(2)(ii) of this section, HCFA may not disclose any accreditation survey or any information directly related to the survey (including corrective action plans) made by and released to it by the Joint Commission on Accreditation of Healthcare Organizations, the American Osteopathic Association or any other national accreditation organization that meets the requirements of §488.6 or §493.506 of this chapter. Materials that are confidential include accreditation letters and accompanying recommendations and comments prepared by an accreditation organization concerning the entities it surveys.

(ii) Exceptions.

(A) HCFA may release the accreditation survey of any home health agency; and

(B) HCFA may release the accreditation survey and other information directly related to the survey (including corrective action plans) to the extent the survey and information relate to an enforcement action (for example, denial of payment for new admissions, civil money penalties, temporary management and termination) taken by HCFA; and

(3) Tax returns and return information defined in section 6103 of the Internal Revenue Code, as amended by the Tax Reform Act of 1976, shall not be disclosed except as authorized by the Internal Revenue Code.
(c) Effect of exemption. Neither 5 U.S.C. 552 nor this regulation directs the withholding of any record or information, except to the extent of the prohibitions in paragraph (b) of this section. Except for material required to be withheld under the statutory provisions incorporated in paragraph (b) of this section or under another statute which meets the standards in 5 U.S.C. 552(b)(3), materials exempt from mandatory disclosure will nevertheless be made available when this can be done consistently with obligations of confidentiality and administrative necessity. The disclosure of materials or records under these circumstances in response to a specific request, however, is of no precedent force with respect to any other request.

§ 401.128 Where requests for records may be made.

(a) General. Any request for any record may be made to—

(1) Any HCFA component;

(2) Director, Office of Public Affairs, HCFA 313-H, Hubert H. Humphrey Building, 200 Independence Avenue, Washington, DC 20201; or

(3) Director of Public Affairs in any Regional Office of the Department of Health and Human Services.

The locations and service areas of these offices are as follows:


Region IV—101 Marietta Street, Atlanta, GA 30323. Alabama, Florida, Georgia, Kentucky, Mississippi, North Carolina, South Carolina, Tennessee.

Region V—300 South Wacker Drive, Chicago, IL 60606. Illinois, Indiana, Michigan, Minnesota, Ohio, Wisconsin.

Region VI—1200 Main Tower Building, Dallas, TX 75202. Arkansas, Louisiana, New Mexico, Oklahoma, Texas.

Region VII—601 East 12th Street, Kansas City, MO 64106. Iowa, Kansas, Missouri, Nebraska.


(b) Records pertaining to individuals. HCFA maintains some records pertaining to individuals. Disclosure of such records is generally prohibited by section 1106 of the Social Security Act (42 U.S.C. 1306), except as prescribed in §401.105 (See also §401.126(b)). Requests for records pertaining to individuals may be addressed to:

Director, Office of Research, Demonstrations and Statistics, HCFA, Baltimore, Maryland 21235, when information is sought from the record of a person who has participated in a research survey conducted by or for HCFA, Office of Research, Demonstrations and Statistics; or whose records have been included by statistical sampling techniques in research and statistical studies authorized by the Social Security Act in the field of health care financing.

(c) Requests for materials listed in §401.130 or §401.132 or indexed in the HCFA Rulings. A request to inspect and copy materials listed in §401.130 or §401.132 or indexed in HCFA Rulings may be made to any district or branch office of the Social Security Administration. If the specific material requested is not available in the office receiving the request, the material will be obtained and made available promptly.

§ 401.130 Materials available at social security district offices and branch offices.

(a) Materials available for inspection. The following are available or will be made available for inspection at the social security district offices and branch offices:

(1) Compilation of the Social Security Laws.

(2) The Public Information Regulation of the Department of Health and Human Services (45 CFR part 5).
(3) Medicare Program regulations issued by the Health Care Financing Administration. 42 CFR chapter IV.
(4) HCFA Rulings.
(b) Materials available for inspection and copying. The following materials are available or will be made available for inspection and copying at the social security district offices and branch offices:
(1) Claims Manual of the Social Security Administration.
(2) Department Staff Manual on Organization, Department of Health and Human Services, Part F, HCFA.
(3) Parts 2 and 3 of the Part A Intermediary Manual (Providers Services under Medicare HCFA Pub. 13-2 and 13-3).
(4) Parts 2 and 3 of the Part B Intermediary Manual (Physician and Supplier Services).
(5) Intermediary Letters Related to Parts 2 and 3 of the Part A and Part B Intermediary Manuals.
(6) State Buy-In Handbook (State Enrollment of Eligible Individuals under the Supplementary Medical Insurance Program) and Letters.
(7) Group Practice Prepayment Plan Manual (HIM-8) and Letters.
(8) State Operations Manual (HIM-7).
HCFA Letters to State Agencies on Medicare.
(10) Skilled Nursing Facility Manual (HCFA Pub. 12).
(11) Hearing Officers Handbook (Supplementary Medical Insurance Program—HIM-21).
(12) Hospital Manual (HIM-10).
(13) Home Health Agency Manual (HIM-11).
(14) Outpatient Physical Therapy Provider Manual (HIM-9).
(16) Audit Program Manuals for Hospital (HIM-16), Home Health Agency (HIM-17), and Extended Care Facilities (HIM-18).
(17) Statements of deficiencies based upon survey reports of health care institutions or facilities prepared after January 31, 1973, by a State agency, and such reports (including pertinent written statements furnished by such institution or facility on such statements of deficiencies), as set forth in
§ 401.132. Except as otherwise provided for at § 401.133 and 488.325 of this chapter for SNFs, such statements of deficiencies, reports, and pertinent written statements shall be available or made available only at the social security district office and regional office servicing the area in which the institution or facility is located, except that such statements of deficiencies and pertinent written statements shall also be available at the local public assistance offices servicing such area.
(b) Indexes to the materials listed in paragraph (a) of this section and in this paragraph (b) and an index to the Bureau of Hearings and Appeals Handbook.
§ 401.132 Materials in field offices of the Office of Hearings and Appeals, SSA.
(a) Materials available for inspection. The following materials are available for inspection in the field offices of the Office of Hearings and Appeals, SSA:
(1) Title 45 of the Code of Federal Regulations (including the public information regulation of the Department of Health and Human Services).
(2) Regulations of the Social Security Administration and HCFA.
(3) Title 5, United States Code.
(4) Compilation of the Social Security Laws.
(5) HCFA Rulings.
(b) Handbook available for inspection and copying. The Office of Hearings and Appeals Handbook is available for inspection and copying in the field offices of the Office of Hearings and Appeals.
§ 401.133 Availability of official reports on providers and suppliers of services, State agencies, intermediaries, and carriers under Medicare.
Except as otherwise provided for in § 488.325 of this chapter for SNFs, the following must be made available to the public under the conditions specified:
(a) Statements of deficiencies and survey reports on providers of services prepared by State agencies. (1) Statements
§ 401.134 Release of Medicare information to State and Federal agencies.

(a) Except as provided in paragraph (b) of this section, the following information may be released to an officer or
§ 401.135 Release of Medicare information to the public.

The following shall be made available to the public under the conditions specified:

(a) Information as to amounts paid to providers and other organizations and facilities for services to beneficiaries under title XVIII of the Act: Provided, That no information identifying any particular beneficiaries shall be disclosed under this paragraph.

(b) The name of any provider of services or other person furnishing services to Medicare beneficiaries who—

(1) Has been found by a Federal court to have been guilty of submitting false claims in connection with Medicare; or

(2) Has been found by a carrier or intermediary, after consultation with a professional medical association functioning external to program administration, to have provided unnecessary services.

(c) The following information may be released to any officer or employee of an agency of the Federal or a State government lawfully charged with the administration of a program receiving grants-in-aid under title V and XIX of the Social Security Act for the purpose of administration of those titles, or to any officer or employee of the Department of Army, Department of Defense, solely for the administration of its Civilian Health and Medical Program of the Uniformed Services (CHAMPUS):

(1) Information, including the identification number, concerning charges made by physicians, other practitioners, or suppliers, and amounts paid under Medicare for services furnished to beneficiaries by such physicians, other practitioners, or suppliers, to enable the agency to determine the proper amount of benefits payable for medical services performed in accordance with those programs or to

(2) Information as to physicians or other practitioners that has been disclosed under § 401.105.

(3) Information relating to the qualifications and certification status of hospitals and other health care facilities obtained in the process of determining whether, and certifying as to whether, institutions or agencies meet or continue to meet the conditions of participation of providers of services or whether other entities meet or continue to meet the conditions for coverage of services they furnish.

(b) The release of such information shall not be authorized by a fiscal intermediary or carrier.

(c) The following information may be released to any officer or employee of an agency of the Federal or a State government lawfully charged with the administration of a program receiving grants-in-aid under title V and XIX of the Social Security Act for the purpose of administration of those titles, or to any officer or employee of the Department of Army, Department of Defense, solely for the administration of its Civilian Health and Medical Program of the Uniformed Services (CHAMPUS), provided that the agency has filed an agreement with HCFA that the information will be released only to the agency's enforcement branch and that the agency will preserve the confidentiality of the information received and will not disclose that information for other than program purposes:

(1) The name and address of any provider of medical services, organization, or other person being actively investigated for possible fraud in connection with Medicare, and the nature of such suspected fraud. An active investigation exists when there is significant evidence supporting an initial complaint but there is need for further investigation.

(2) The name and address of any provider of medical services, organization, or other person found, after consultation with an appropriate professional association or program review team, to have provided unnecessary services, or of any physician or other individual found to have violated the assignment agreement on at least three occasions.

(3) The name and address of any provider of medical services, organization or other person released under paragraph (c)(1) or (2) of this section concerning which an active investigation is concluded with a finding that there is no fraud or other prosecutable offense.
medical authority, to have been engaged in a pattern of furnishing services to beneficiaries which are substantially in excess of their medical needs; except that the name of any provider or other person shall not be disclosed pursuant to a finding under this paragraph (b)(2) of this section, unless that provider or other person has first been afforded a reasonable opportunity to offer evidence on his behalf.

(c) Upon request in writing, cost reports submitted by providers of services pursuant to section 1815 of the Act to enable the Secretary to determine amounts due the providers.

§ 401.136 Requests for information or records.

(a) A request should reasonably identify the requested record by brief description. Requesters who have detailed information which would assist in identifying the records requested are urged to provide such information in order to expedite the handling of the request. Envelopes in which written requests are submitted should be clearly identified as Freedom of Information requests. The request should include the fee or request determination of the fee. When necessary, a written request will be promptly forwarded to the proper office, and the requester will be advised of the date of the receipt and identification and address of the proper office.

(b) Determinations of whether records will be released or withheld will be made within 10 working days from date of receipt of the request in the office listed in § 401.128 except where HCFA extends this time and sends notice of such extension to the requester. Such extension may not exceed 10 additional working days and shall apply only where the following unusual circumstances exist:

(1) The need to search for and collect the requested records from field facilities or other establishments that are separate from the office processing the requests;

(2) The need to search for, collect, and appropriately examine a voluminous amount of separate and distinct records which are requested in a single request; or

(3) The need for consultation, which shall be conducted with all practicable speed, with another agency having a substantial interest in the request or among two or more components of HCFA having a substantial interest in the subject matter of the request.

(c) If an extension is made, the requester will be notified in writing before the expiration of 10 working days from receipt of the request and will be given an explanation of why the extension was necessary and the date on which a determination will be made.

(d) Authority to extend the time limit with respect to any request for information or records is granted to the Director, Office of Public Affairs, HCFA and to the Director of Public Affairs in any HHS Regional Office. Those officers and employees of HCFA who are listed in § 401.144(a) as having authority to deny requests for information from records maintained on individuals are granted authority to extend the time limit for responding to requests for information from such records.

§ 401.140 Fees and charges.

(a) Statement of policy. It is HCFA's policy to comply with certain requests for information services without charge. Except as otherwise determined pursuant to paragraph (c) of this section, fees will be charged for the following services with respect to all other requests for information from records which are reasonably identified by the requesters:

(1) Reproduction, duplication, or copying of records;

(2) Searches for records; and

(3) Certification or authentication of records.

(b) Fee schedules. The fee schedule is as follows:

(1) Search for records. Three dollars per hour: Provided, however, That no charge will be made for the first half hour.

(2) Reproduction, duplication, or copying of records. Ten cents per page where such reproduction can be made by commonly available photocopier machines. The cost of reproducing records which cannot be so photocopied will be determined on an individual basis at actual cost.
§ 401.144 Certification or authentication of records. Three dollars per certification or authentication.

§ 401.148 Administrative review.

(a) Review by the Administrator. A person whose request has been denied may initiate a review by filing a request for review with the Administrator of HCFA, 700 East High Rise Building, 6401 Security Boulevard, Baltimore, Maryland 21235, within 30 days of receipt of the determination to deny or within 30 days of receipt of records which are in partial response to his request if a portion of a request is granted and a portion denied, whichever is later. Upon receipt of a timely request for review, the Administrator will review the decision in question and the findings upon which it was based. Upon the basis of the data considered in connection with the decision and whatever other evidence and written argument is submitted by the person requesting the review or which is otherwise obtained, the Administrator or his designee will affirm or revise in whole or in part the findings and decision in question. A decision to affirm the denial will be made only upon concurrence of the Assistant Secretary for Public Affairs, or his designee, after consultation with the General Counsel or his or her designee, and the appropriate program policy official. Written notice of the decision of the Administrator will be mailed to the person who requested the review. A written decision will be made within 20 working days from receipt of the request for review. Extension of the time limit may be granted under the circumstances listed in § 401.136(b) to the extent that the maximum 10 days limit on extensions has not been exhausted on the initial determination. The decision will include the basis for it and will advise the requester of his right to judicial review.

(b) Failure of the Administrator to comply with the time limits. Failure of the Administrator to comply with the time limits set forth in § 401.136 and this section constitutes an exhaustion of the requester's administrative remedies.

§ 401.152 Court review.

Where the Administrator upon review affirms the denial of a request for records, in whole or in part, the requester may seek court review in the district court of the United States pursuant to 5 U.S.C. 552(a)(4)(B).
§ 401.601 Basis and scope.

(a) Basis. This subpart implements for HCFA the Federal Claims Collection Act (FCCA) of 1966 (31 U.S.C. 3711), and conforms to the regulations (4 CFR parts 101-105) issued jointly by the General Accounting Office and the Department of Justice that generally prescribe claims collection standards and procedures under the FCCA for the Federal government.

(b) Scope. Except as provided in paragraphs (c) through (f) of this section, the regulations in this subpart describe HCFA's procedures and standards for the collection of claims in any amount, and the compromise of, or the suspension or termination of collection action on, all claims for money or property that do not exceed $100,000 or such higher amount as the Attorney General may from time to time prescribe, exclusive of interest, arising under any functions delegated to HCFA by the Secretary.

(c) Amount of claim. HCFA refers all claims that exceed $100,000 or such higher amount as the Attorney General may from time to time prescribe, exclusive of interest, arising under any functions delegated to HCFA by the Secretary.

(d) Related regulations—(1) Department regulations. DHHS regulations applicable to HCFA that generally implement the FCCA for the Department are located at 45 CFR part 30. These regulations apply only to the extent HCFA regulations do not address a situation.

(2) HCFA regulations. The following regulations govern specific debt management situations encountered by HCFA and supplement this subpart:

(i) Claims against Medicare beneficiaries for the recovery of overpayments are covered in 20 CFR 404.515.

(ii) Adjustments in Railroad Retirement or Social Security benefits to recover Medicare overpayments to individuals are covered in §§ 405.350-405.358 of this chapter.

(iii) Claims against providers, physicians, or other suppliers of services for overpayments under Medicare and for assessment of interest are covered in §§ 405.377 and 405.378 of this chapter, respectively.

(iv) Claims against beneficiaries for unpaid hospital insurance or supplementary medical insurance premiums under Medicare are covered in § 408.110 of this chapter.

(v) State repayment of Medicaid funds by installments is covered in § 430.48 of this chapter.

(e) Collection and compromise under other statutes and at common law. The regulations in this subpart do not—

(1) Preclude disposition by HCFA of claims under statutes, other than the FCCA, that provide for the collection or compromise of a claim, or suspension or termination of collection action.

(2) Affect any rights that HCFA may have under common law as a creditor.

(f) Fraud. The regulations in this subpart do not apply to claims in which there is an indication of fraud, the presentation of a false claim, or misrepresentation on the part of a debtor or any other party having an interest in the claim. HCFA forwards these claims to the Department of Justice for disposition under 4 CFR 105.1.

(g) Enforced collection. HCFA refers claims to the Department of Justice for enforced collection through litigation in those cases which cannot be compromised or on which collection action cannot be suspended or terminated in accordance with this subpart or the regulations issued jointly by the Attorney General and the Comptroller General.

§ 401.603 Definitions.

For purposes of this subpart—

Claim means any debt owed to HCFA.

Debtor means any individual, partnership, corporation, estate, trust or other legal entity against which HCFA has a claim.
§ 401.605 Omissions not a defense.

The failure of HCFA to comply with the regulations in this subpart, or with the related regulations listed in §401.601(d), is not available as a defense to a debtor against whom HCFA has a claim for money or property.

§ 401.607 Claims collection.

(a) General policy. HCFA recovers amounts of claims due from debtors, including interest where appropriate, by—

(1) Direct collections in lump sums or in installments; or

(2) Offsets against monies owed to the debtor by the Federal government where possible.

(b) Collection in lump sums. Whenever possible, HCFA attempts to collect claims in full in one lump sum. However, if HCFA determines that a debtor is unable to pay the claim in one lump sum, HCFA may instead enter into an agreement to accept regular installment payments.

(c) Collection in installments. Generally, HCFA requires that all claims to be satisfied by installment payments must be liquidated in three years or less. If unusual circumstances exist, such as the possibility of debtor insolvency, an installment agreement that extends beyond three years may be approved.

(1) Debtor request. If a debtor desires to repay a claim in installments, the debtor must submit—

(i) A request to HCFA; and

(ii) Any information required by HCFA to make a decision regarding the request.

(2) HCFA decision. HCFA will determine the number, amount and frequency of installment payments based on the information submitted by the debtor and on other factors such as—

(i) Total amount of the claim;

(ii) Debtor's ability to pay; and

(iii) Cost to HCFA of administering an installment agreement.

(d) Collection by offset. (1) HCFA may offset, where possible, the amount of a claim against the amount of pay, compensation, benefits or other monies that a debtor is receiving or is due from the Federal government.

(2) Under regulations at §405.350—405.358 of this chapter, HCFA may initiate adjustments in program payments to which an individual is entitled under title II of the Act (Federal Old Age, Survivors, and Disability Insurance Benefits) or under the Railroad Retirement Act of 1974 (45 U.S.C. 231) to recover Medicare overpayments.

§ 401.613 Compromise of claims.

(a) Amount of compromise. HCFA requires that the amount to be recovered through a compromise of a claim must—

(1) Bear a reasonable relation to the amount of the claim; and

(2) Be recoverable through enforced collection procedures.

(b) General factors. After considering the bases for a decision to compromise a claim under paragraph (c) of this section, HCFA may further consider factors such as—

(1) The age and health of the debtor if the debtor is an individual;

(2) Present and potential income of the debtor; and

(3) Whether assets have been concealed or improperly transferred by the debtor.

(c) Basis for compromise. Bases on which HCFA may compromise a claim include the following—

(1) Inability to pay. HCFA may compromise a claim if it determines that the debtor, or the estate of a deceased debtor, does not have the present or prospective ability to pay the full amount of the claim within a reasonable time.

(2) Litigative probabilities. HCFA may compromise a claim if it determines that it would be difficult to prevail in a case before a court of law as a result of the legal issues involved or inability of the parties to agree to the facts of the case. The amount that HCFA accepts in compromise under this provision will reflect—

(i) The likelihood that HCFA would have prevailed on the legal question(s) involved;

(ii) Whether and to what extent HCFA would have obtained a full or partial recovery of a judgment, depending on the availability of witnesses, or
other evidentiary support for HCFA's claim; and

(iii) The amount of court costs that would be assessed to HCFA.

(3) Cost of collecting the claim. HCFA may compromise a claim if it determines that the cost of collecting the claim does not justify the enforced collection of the full amount. In this case, HCFA may adjust the amount it accepts as a compromise to allow an appropriate discount for the costs of collection it would have incurred but for the compromise.

(d) Enforcement policy. HCFA may compromise statutory penalties, forfeitures, or debts established as an aid to enforcement or to compel compliance, if it determines that its enforcement policy, in terms of deterrence and securing compliance both present and future, is adequately served by acceptance of the compromise amount.

§ 401.615 Payment of compromise amount.

(a) Time and manner of compromise. Payment by the debtor of the amount that HCFA has agreed to accept as a compromise in full settlement of a claim must be made within the time and in the manner prescribed by HCFA. Accordingly, HCFA will not settle a claim until the full payment of the compromise amount has been made.

(b) Effect of failure to pay compromise amount. Failure of the debtor to make payment, as provided by the compromise agreement, reinstates the full amount of the claim, less any amounts paid prior to the default.

(c) Prohibition against grace periods. HCFA will not agree to inclusion of a provision in an installment agreement that would permit grace periods for payments that are late under the terms of the agreement.

§ 401.617 Suspension of collection action.

(a) General conditions. HCFA may temporarily suspend collection action on a claim if the following general conditions are met—

(1) Amount of future recovery. HCFA determines that future collection action may result in a recovery of an amount sufficient to justify periodic review and action on the claim by HCFA during the period of suspension.

(2) Statute of limitations. HCFA determines that—

(i) The applicable statute of limitations has been tolled, waived or has started running anew; or

(ii) Future collections may be made by HCFA through offset despite an applicable statute of limitations.

(b) Basis for suspension. Bases on which HCFA may suspend collection action on a particular claim include the following—

(1) A debtor cannot be located; or

(2) A debtor—

(i) Owns no substantial equity in property;

(ii) Is unable to make payment on HCFA's claim or is unable to effect a compromise; and

(iii) Has future prospects that justify retention of the claim.

(c) Locating debtors. HCFA will make every reasonable effort to locate missing debtors sufficiently in advance of the bar of an applicable statute of limitations to permit timely filing of a lawsuit to recover the amount of the claim.

(d) Effect of suspension on liquidation of security. HCFA will liquidate security, obtained in partial recovery of a claim, despite a decision under this section to suspend collection action against the debtor for the remainder of the claim.

§ 401.621 Termination of collection action.

(a) General factors. After considering the bases for a decision to terminate collection action under paragraph (b) of this section, HCFA may further consider factors such as—

(1) The age and health of the debtor if the debtor is an individual;

(2) Present and potential income of the debtor; and

(3) Whether assets have been concealed or improperly transferred by the debtor.

(b) Basis for termination of collection action. Bases on which HCFA may terminate collection action on a claim include the following—

(1) Inability to collect a substantial amount of the claim. HCFA may terminate collection action if it determines
§ 401.623 Joint and several liability.

(a) Collection action. HCFA will liquidate claims as quickly as possible. In cases of joint and several liability among two or more debtors, HCFA will not allocate the burden of claims payment among the debtors. HCFA will proceed with collection action against one debtor even if other liable debtors have not paid their proportionate shares.

(b) Compromise. Compromise with one debtor does not release a claim against remaining debtors. Furthermore, HCFA will not consider the amount of a compromise with one debtor to be a binding precedent concerning the amounts due from other debtors who are jointly and severally liable on the claim.

§ 401.625 Effect of HCFA claims collection decisions on appeals.

Any action taken under this subpart regarding the compromise of a claim, or suspension or termination of collection action on a claim, is not an initial determination for purposes of HCFA appeal procedures.

PART 402—CIVIL MONEY PENALTIES, ASSESSMENTS, AND EXCLUSIONS

Subpart A—General Provisions

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Subpart B—Civil Money Penalties and Assessments

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Subpart C—Exclusions [Reserved]

AUTHORITY: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh).
SOURCE: 63 FR 68690, Dec. 14, 1998, unless otherwise noted.
paragraph (c), (d), or (e) of this section; and
(2) Sets forth the appeal rights of persons subject to penalties, assessments, or exclusion and the procedures for reinstatement following exclusion.
(c) Civil money penalties. HCFA or OIG may impose civil money penalties against any person or other entity specified in paragraphs (c)(1) through (c)(30) of this section under the identified section of the Act. The authorities that also permit imposition of an assessment or exclusion are noted in the applicable paragraphs.
(1) Sections 1833(h)(5)(D) and 1842(j)(2)—Any person that knowingly and willfully, and on a repeated basis, bills for a clinical diagnostic laboratory test, other than on an assignment-related basis. This provision includes tests performed in a physician’s office but excludes tests performed in a rural health clinic. (This violation may also include an assessment and cause exclusion.)
(2) Section 1833(i)(6)—Any person that knowingly and willfully presents, or causes to be presented, a bill or request for payment for an intraocular lens inserted during or after cataract surgery for which the Medicare payment rate includes the cost of acquiring the class of lens involved.
(3) Section 1833(q)(2)(B)—Any entity that knowingly and willfully fails to provide information about a referring physician, including the physician’s name and unique physician identification number for the referring physician, when seeking payment on an unassigned basis. (This violation, if it occurs in repeated cases, may also cause an exclusion.)
(4) Sections 1834(a)(11)(A) and 1842(j)(2)—Any durable medical equipment supplier that knowingly and willfully charges for a covered service that is furnished on a rental basis after the rental payments may no longer be made (except for maintenance and servicing) as provided in section 1834(a)(7)(A). (This violation may also include an assessment and cause exclusion.)
(5) Sections 1834(a)(18)(B) and 1842(j)(2)—Any nonparticipating durable medical equipment supplier that knowingly and willfully, in violation of section 1834(a)(18)(A), fails to make a refund to Medicare beneficiaries for a covered service for which payment is precluded due to an unsolicited telephone contact from the supplier. (This violation may also include an assessment and cause exclusion.)
(6) Sections 1834(b)(5)(C) and 1842(j)(2)—Any nonparticipating physician or supplier that knowingly and willfully charges a Medicare beneficiary more than the limiting charge, as specified in section 1834(b)(5)(B), for radiologist services. (This violation may also include an assessment and cause exclusion.)
(7) Sections 1834(c)(4)(C) and 1842(j)(2)—Any nonparticipating physician or supplier that knowingly and willfully charges a Medicare beneficiary more than the limiting charge, as specified in section 1834(c)(4)(B), for mammography screening. (This violation may also include an assessment and cause exclusion.)
(8) Sections 1834(h)(3) and 1842(j)(2)—Any supplier of prosthetic devices, orthotics, and prosthetics that knowingly and willfully charges for a covered prosthetic device, orthotic, or prosthetic that is furnished on a rental basis after the rental payment may no longer be made (except for maintenance and servicing). (This violation may also include an assessment and cause exclusion.)
(9) Section 1834(j)(2)(A)(iii)—Any supplier of durable medical equipment, including a supplier of prosthetic devices, prosthetics, orthotics, or supplies, that knowingly and willfully distributes a certificate of medical necessity in violation of section 1834(j)(2)(A)(i) or fails to provide the information required under section 1834(j)(2)(A)(ii).
(10) Sections 1834(j)(4) and 1842(j)(2)—(i) Any supplier of durable medical equipment, including a supplier of prosthetic devices, prosthetics, orthotics, or supplies, that knowingly and willfully fails to make refunds in a timely manner to Medicare beneficiaries for services billed other than on an assignment-related basis if—
(A) The supplier does not possess a Medicare supplier number;
(B) The service is denied in advance under section 1834(a)(15); or
(C) The service is determined not to be medically necessary or reasonable.

(ii) These violations may also include an assessment and cause exclusion.

(11) Sections 1842(b)(18)(B) and 1842(j)(2)—Any practitioner specified in section 1842(b)(18)(C) (physician assistants, nurse practitioners, clinical nurse specialists, certified registered nurse anesthetists, certified nurse-midwives, clinical social workers, and clinical psychologists) or other person that knowingly and willfully bills or collects for any services by the practitioners on other than an assignment-related basis. (This violation may also include an assessment and cause exclusion.)

(12) Sections 1842(k) and 1842(j)(2)—Any physician who knowingly and willfully presents, or causes to be presented, a claim or bill for an assistant at cataract surgery performed on or after March 1, 1987 for which payment may not be made because of section 1862(a)(15). (This violation may also include an assessment and cause exclusion.)

(13) Sections 1842(l)(3) and 1842(j)(2)—Any nonparticipating physician who does not accept payment on an assignment-related basis and who knowingly and willfully fails to refund on a timely basis any amounts collected for services that are not reasonable or medically necessary or are of poor quality, in accordance with section 1842(l)(1)(A). (This violation may also include an assessment and cause exclusion.)

(14) Sections 1842(m)(3) and 1842(j)(2)—(i) Any nonparticipating physician, who does not accept payment for an elective surgical procedure on an assignment-related basis and whose charge is at least $500, who knowingly and willfully fails to—

(A) Disclose the information required by section 1842(m)(1) concerning charges and coinsurance amounts; and

(B) Refund on a timely basis any amounts collected for the procedure in excess of the charges recognized and approved by the Medicare program.

(ii) This violation may also include an assessment and cause exclusion.

(15) Sections 1842(n)(3) and 1842(j)(2)—Any physician who knowingly and willfully, in repeated cases, bills one or more beneficiaries, for purchased diagnostic tests, any amount other than the payment amount specified in section 1842(n)(1)(A) or section 1842(n)(1)(B). (This violation may also include an assessment and cause exclusion.)

(16) Section 1842(p)(3)(A)—Any physician who knowingly and willfully fails promptly to provide the appropriate diagnosis code or codes upon request by HCFA or a carrier on any request for payment or bill not submitted on an assignment-related basis for any service furnished by the physician. (This violation, if it occurs in repeated cases, may also cause exclusion.)

(17) Sections 1842(l)(3) and 1842(j)(2)—

(i) Any nonparticipating physician, supplier, or other person that furnishes physicians' services and does not accept payment on an assignment-related basis, that—

(A) Knowingly and willfully bills or collects in excess of the limiting charge (as defined in section 1842(l)(1)(A)) on a repeated basis; or

(B) Fails to make an adjustment or refund on a timely basis as required by section 1848(g)(1)(A)(iii) or (iv).

(ii) These violations may also include an assessment and cause exclusion.

(18) Section 1848(g)(3)(B) and 1842(j)(2)—Any person that knowingly and willfully bills for State plan approved physicians' services, as defined in section 1848(j)(3), on other than an assignment-related basis for a Medicare beneficiary who is also eligible for Medicaid (these individuals include qualified Medicare beneficiaries). This provision applies to services furnished on or after April 1, 1990. (This violation may also include an assessment and cause exclusion.)

(19) Section 1842(p)(3), 1842(l)(3), and 1842(j)(2)(A)—

(i) Any physician, supplier, or other person (except any person that has been excluded from the Medicare program) that, for services furnished after September 1, 1990, knowingly and willfully—

(A) Fails to submit a claim on a standard claim form for services provided for which payment is made under Part B on a reasonable charge or fee schedule basis; or
(B) Imposes a charge for completing and submitting the standard claims form.

(ii) These violations, if they occur in repeated cases, may also cause exclusion.

(20) Section 1862(b)(5)(C)—Any employer (other than a Federal or other governmental agency) that, before October 1, 1998, willfully or repeatedly fails to provide timely and accurate information requested relating to an employee's group health insurance coverage.

(21) Section 1862(b)(6)(B)—Any entity that knowingly, willfully, and repeatedly—

(i) Fails to complete a claim form relating to the availability of other health benefit plans in accordance with section 1862(b)(6)(A); or

(ii) Provides inaccurate information relating to the availability of other health benefit plans on the claim form.

(22) Section 1877(g)(5)—Any person that fails to report information required by HHS under section 1877(f) concerning ownership, investment, and compensation arrangements. (This violation may also include an assessment and cause exclusion.)

(23) Sections 1879(h), 1834(a)(18), and 1842(j)(2)—

(i) Any durable medical equipment supplier, including a supplier of prosthetic devices, prosthetics, orthotics, or supplies, that knowingly and willfully fails to make refunds in a timely manner to Medicare beneficiaries for services billed on an assignment-related basis if—

(A) The supplier did not possess a Medicare supplier number;

(B) The service is denied in advance under section 1834(a)(15) of the Act; or

(C) The service is determined not to be payable under section 1834(a)(17)(b) because of unsolicited telephone contacts.

(ii) These violations may also include an assessment and cause exclusion.

(24) Section 1882(a)(2)—Any person that issues a Medicare supplemental policy that has not been approved by the State regulatory program or does not meet Federal standards on and after the effective date in section 1882(p)(1)(C). (This violation may also include an assessment and cause exclusion.)

(25) Section 1882(p)(8)—Any person that sells or issues Medicare supplemental policies, on or after July 30, 1992, that fail to conform to the NAIC or Federal standards established under section 1882(p). (This violation may also include an assessment and cause exclusion.)

(26) Section 1882(p)(9)(C)—

(i) Any person that sells a Medicare supplemental policy and—

(A) Fails to make available for sale the core group of basic benefits when selling other Medicare supplemental policies with additional benefits; or

(B) Fails to provide the individual, before the sale of the policy, an outline of coverage describing the benefits provided by the policy.

(ii) These violations may also include an assessment and cause exclusion.

(27) Section 1882(q)(5)(C)—

(i) Any person that fails to—

(A) Suspend a Medicare supplemental policy at the policyholder’s request, if the policyholder applies for and is determined eligible for medical assistance, and the policyholder provides notice within 90 days of the eligibility determination; or

(B) Automatically reinstate the policy as of the date of termination of medical assistance if the policyholder loses eligibility for medical assistance and the policyholder provides notice within 90 days of loss of eligibility.

(ii) These violations may also include an assessment and cause exclusion.

(28) Section 1882(r)(6)(A)—Any person that fails to provide refunds or credits as required by section 1882(r)(1)(B). (This violation may also include an assessment and cause exclusion.)

(29) Section 1882(s)(3)—

(i) Any issuer of a Medicare supplemental policy that—

(A) Does not waive any time periods applicable to preexisting conditions, waiting periods, elimination periods, or probationary periods if the time periods were already satisfied under a preceding Medicare supplemental policy; or

(B) Denies a policy, conditions the issuance or effectiveness of the policy, or discriminates in the pricing of the policy based on health status or other
criteria as specified in section 1882(s)(2)(A).

(ii) These violations may also include an assessment and cause exclusion.

(30) Section 1882(t)(2)—

(i) Any issuer of a Medicare supplemental policy that—

(A) Fails substantially to provide medically necessary services to enrollees seeking the services through the issuer’s network of entities;

(B) Imposes premiums on enrollees in excess of the premiums approved by the State;

(C) Acts to expel an enrollee for reasons other than nonpayment of premiums; or

(D) Does not provide each enrollee at the time of enrollment with the specific information provided in section 1882(t)(1)(E)(i) or fails to obtain a written acknowledgment from the enrollee of receipt of the information (as required by section 1882(t)(1)(E)(ii)).

(ii) These violations may also include an assessment and cause exclusion.

(d) Assessments. HCFA or OIG may impose assessments in addition to civil money penalties for violations of the following statutory sections:

(1) Section 1833: Paragraph (h)(5)(D).

(2) Section 1834: Paragraphs (a)(11)(A), (a)(18)(B), (b)(5)(C), (c)(4)(C), (h)(3), and (j)(4).

(3) Section 1842: Paragraphs (k), (l)(3), (m)(3), and (n)(3).

(4) Section 1848: Paragraph (g)(1)(B).

(5) Section 1877: Paragraph (g)(5).

(6) Section 1879: Paragraph (h).

(7) Section 1882: Paragraphs (d)(2), (p)(8), (p)(9)(C), (q)(5)(C), (r)(6)(A), (s)(3), and (t)(2).

(e) Exclusions. (1) HCFA or OIG may exclude any person from participation in the Medicare program on the basis of any of the following violations of the statute:

(i) Section 1833: Paragraphs (h)(5)(D) and, in repeated cases, (q)(2)(B).

(ii) Section 1834: Paragraphs (a)(11)(A), (a)(18)(B), (b)(5)(C), (c)(4)(C), (h)(3), and (j)(4).

(iii) Section 1842: Paragraphs (b)(18)(B), (k), (l)(3), (m)(3), (n)(3), and, in repeated cases, (p)(3)(B).

(iv) Section 1848: Paragraphs (g)(1)(B), (g)(3)(B), and, in repeated cases, (g)(4)(B)(ii).

(v) Section 1877: Paragraph (g)(5).

(vi) Section 1879: Paragraph (h).

(vii) Section 1882: Paragraphs (a)(2), (p)(8), (p)(9)(C), (q)(5)(C), (r)(6)(A), (s)(3), and (t)(2).

(2) HCFA or OIG must exclude from participation in the Medicare program any of the following, under the identified section of the Act:

(i) Section 1834(a)(17)(C)—Any supplier of durable medical equipment and supplies that are covered under section 1834(a)(13) that knowingly contacts Medicare beneficiaries by telephone regarding the furnishing of covered services in violation of section 1834(a)(17)(A) and whose conduct establishes a pattern of prohibited contacts as described under section 1834(a)(17)(A).

(ii) Section 1834(h)(3)—Any supplier of prosthetic devices, orthotics, and prosthetics that knowingly contacts Medicare beneficiaries by telephone regarding the furnishing of prosthetic devices, orthotics, or prosthetics in the same manner as in the violation under section 1834(a)(17)(A) and whose conduct establishes a pattern of prohibited contacts in the same manner as described in section 1834(a)(17)(C).

(f) Responsible persons. (1) If HCFA or OIG determines that more than one person is responsible for any of the violations described in paragraph (c) or paragraph (d) of this section, it may impose a civil money penalty or a civil money penalty and assessment against any one of those persons or jointly and severally against two or more of those persons. However, the aggregate amount of the assessments collected may not exceed the amount that could be assessed if only one person were responsible.

(2) A principal is liable for penalties and assessments for the actions of his or her agent acting within the scope of the agency.

(g) Time limits. Neither HCFA nor OIG initiates an action to impose a civil money penalty, assessment, or proceeding to exclude a person from participation in the Medicare program unless it begins the action within 6 years from the date on which the claim was presented, the request for payment was made, or the incident occurred.
§ 402.3 Definitions.

For purposes of this part:

Assessment means the amount described in § 402.107 and includes the plural of that term.

Assignment-related basis means that the claim submitted by a physician, supplier or other person is paid on the basis of an assignment, whereby the physician, supplier or other person agrees to accept the Medicare payment as payment in full for the services furnished to the beneficiary and is precluded from charging the beneficiary more than the deductible and coinsurance based upon the approved Medicare fee amount. Additional obligations, including obligations to make refunds in certain circumstances, are established at section 1842(b)(3) of the Act.

Claim means an application for payment for a service for which the Medicare or Medicaid program may pay.

Covered means that a service is described as reasonable and necessary for the diagnosis or treatment of illness or injury or to improve the functioning of a malformed body member. A service is not covered if it is specifically identified as excluded from Medicare Part B coverage or is not a defined Medicare Part B benefit.

Exclusion means the temporary or permanent barring of a person or other entity from participation in the Medicare or State health care program and that services furnished or ordered by that person are not paid for under either program.

General Counsel means the General Counsel of HHS or his or her designees.

Knowingly or knowingly and willfully means that a person, with respect to information—

(1) Has actual knowledge of the information;
(2) Acts in deliberate ignorance of the truth or falsity of the information; or
(3) Acts in reckless disregard of the truth or falsity of the information; and
(4) No proof of specific intent is required.

Medicare supplemental policy means a policy guaranteeing that a health plan will pay a policyholder's coinsurance and deductible and will cover other limitations on payment imposed under title XVIII of the Act and will provide additional health plan or non-Medicare coverage for services up to a predefined benefit limit.

NAIC stands for the National Association of Insurance Commissioners.

Nonparticipating describes a physician, supplier, or other person (excluding any provider of services) that, at the time of furnishing the services to Medicare Part B beneficiaries, is not a participating physician or supplier.

Participating describes a physician or supplier (excluding any provider of services) that, before the beginning of any given year, enters into an agreement with HHS that provides that the physician or supplier will accept payment under the Medicare program on an assignment-related basis for all services furnished to Medicare Part B beneficiaries.

Penalty means the amount described in § 402.105 and includes the plural of that term.

Person means an individual, trust or estate, partnership, corporation, professional association or corporation, or other entity, public or private.

Physicians' services means the following Medicare covered professional services:

(1) Surgery, consultation, home, office and institutional calls, and other professional services performed by physicians.
(2) Services and supplies furnished "incident to" a physician's professional services.
(3) Outpatient physical and occupational therapy services.
(4) Diagnostic x-ray tests and other diagnostic tests (excluding clinical diagnostic laboratory tests).
(5) X-ray, radium, and radioactive isotope therapy, including materials and services of technicians.
(6) Antigens prepared by a physician.

Radiologist service means radiology services performed only by, or under the direction of, a physician who is certified, or eligible to be certified, by the American Board of Radiology or for whom radiology services account for at least 50 percent of the total amount of charges made under part B of title XVIII of the Act.

Request for payment means an application submitted by a person to any person for payment for a service.
§ 402.5 Right to a hearing before the final determination.

HCFA or OIG does not make a determination adverse to any person under this part until the person has been given a written notice and opportunity for the determination to be made on the record after a hearing at which the person is entitled to be represented by counsel, to present witnesses, and to cross-examine witnesses against the person.

§ 402.7 Notice of proposed determination.

(a) If HCFA or OIG proposes a penalty and, as applicable, an assessment, or proposes to exclude a respondent from participation in Medicare in accordance with this part, it sends the respondent written notice of its intent by certified mail, return receipt requested. The notice includes the following information:

(1) Reference to the statutory basis or bases for the penalty, assessment, exclusion, or any combination, as applicable.

(2)(i) A description of the claims, requests for payment, or incidents with respect to which the penalty, assessment, and exclusion are proposed; or

(ii) If HCFA or OIG is relying upon statistical sampling to project the number and types of claims or requests for payment and the dollar amount, a description of the claims and requests for payment comprising the sample and a brief description of the statistical sampling technique HCFA or OIG used.

(3) The reason why the claims, requests for payment, or incidents are subject to a penalty and assessment.

(4) The amount of the proposed penalty and of any proposed assessment.

(5) Any mitigating or aggravating circumstances that HCFA or OIG considered when it determined the amount of the proposed penalty and any applicable assessment.

(6) Information concerning response to the notice, including—

(i) A specific statement of the respondent’s right to a hearing; and

(ii) A statement that failure to request a hearing within 60 days renders the proposed determination final and permits the imposition of the proposed penalty and any assessment.

(iii) A statement that the debt may be collected through an administrative offset.

(7) In the case of a respondent that has an agreement under section 1866 of the Act, notice that imposition of an exclusion may result in termination of the provider’s agreement in accordance with section 1866(b)(2)(C) of the Act.

§ 402.9 Failure to request a hearing.

(a) If the respondent does not request a hearing within 60 days of receipt of the notice of proposed determination specified in §402.7, any civil money penalty, assessment, or exclusion becomes final and HCFA or OIG may impose the proposed penalty, assessment, or exclusion, or any less severe penalty, assessment, or suspension.

(b) HCFA or OIG notifies the respondent by certified mail, return receipt requested, of any penalty, assessment, or exclusion that has been imposed and of the means by which the respondent may satisfy the judgment.
§ 402.105 Amount of penalty.

(a) $2,000. Except as provided in paragraphs (b) through (f) of this section, HCFA or OIG may impose a penalty of not more than $2,000 for each service, bill, or refusal to issue a timely refund that is subject to a determination under this part and for each incident involving the knowing, willful, and repeated failure of an entity furnishing a service to submit a properly completed claim form or to include on the claim form accurate information regarding the availability of other health insurance benefit plans (§402.1(c)(21)).

(b) $1,000. HCFA or OIG may impose a penalty of not more than $1,000 for the following:

(c) The respondent has no right to appeal a penalty, assessment, or exclusion for which he or she has not requested a hearing.

§ 402.11 Notice to other agencies and other entities.

(a) Whenever a penalty, assessment, or exclusion becomes final, HCFA or OIG notifies the following organizations and entities about the action and the reasons for it:

(1) The appropriate State or local medical or professional association.

(2) The appropriate peer review organization.

(3) As appropriate, the State agency responsible for the administration of each State health care program (Medicaid, the Maternal and Child Health Services Block Grant Program, and the Social Services Block Grant Program).

(4) The appropriate Medicare carrier or fiscal intermediary.

(5) The appropriate State or local licensing agency or organization (including the Medicare and Medicaid State survey agencies).

(6) The long-term care ombudsman.

(b) For exclusions, HCFA or OIG also notifies the public and specifies the effective date.

§ 402.13 Penalty, assessment, and exclusion not exclusive.

Penalties, assessments, and exclusions imposed under this part are in addition to any other penalties prescribed by law.

§ 402.15 Collateral estoppel.

(a) When a final determination that the respondent presented or caused to be presented a claim or request for payment falling within the scope of §402.1 has been rendered in any proceeding in which the respondent was a party and had an opportunity to be heard, the respondent is bound by that determination in any proceeding under this part.

(b) A person who has been convicted (whether upon a verdict after trial or upon a plea of guilty or nolo contendere) of a Federal crime charging fraud or false statements is barred from denying the essential elements of the criminal offense if the proceedings under this part involve the same transactions.

§ 402.17 Settlement.

HCFA or OIG has exclusive authority to settle any issues or case, without the consent of the ALJ or the Secretary, at any time before a final decision by the Secretary. Thereafter, the General Counsel has the exclusive authority.

§ 402.19 Hearings and appeals.

The hearings and appeals procedures set forth in part 1005 of chapter V of this title are available to any person that receives an adverse determination under this part. For an appeal of a civil money penalty, assessment, or exclusion imposed under this part, either HCFA or OIG may represent the government in the hearing and appeals process.

§ 402.21 Judicial review.

After exhausting all available administrative remedies, a respondent may seek judicial review of a penalty, assessment, or exclusion that has become final. The respondent may seek review only with respect to a penalty, assessment, or exclusion with respect to which the respondent filed an exception under §1005.21(c) of this title unless the court excuses the failure or neglect to urge the exception in accordance with section 1128A(e) of the Act because of extraordinary circumstances.
§ 402.105

(1) Per certificate of medical necessity knowingly and willfully distributed to physicians on or after December 31, 1994 that—
   (i) Contains information concerning the medical condition of the patient; or
   (ii) Fails to include cost information.

(2) Per individual about whom information is requested, for willful or repeated failure of an employer to respond to an intermediary or carrier about coverage of an employee or spouse under the employer’s group health plan (§ 402.1(c)(20)).

(c) $5,000. HCFA or OIG may impose a penalty of not more than $5,000 for each violation resulting from the following:
   (1) The failure of a Medicare supplemental policy issuer, on a replacement policy, to waive any time periods applicable to pre-existing conditions, waiting periods, elimination periods, or probationary periods that were satisfied under a preceding policy (§ 402.1(c)(29)); and
   (2) Any issuer of any Medicare supplemental policy denying a policy, conditioning the issuance or effectiveness of the policy, or discriminating in the pricing of the policy based on health status or other criteria as specified in section 1882(c)(20)(A), § 402.1(c)(29).

(d) $10,000. (1) HCFA or OIG may impose a penalty of not more than $10,000 for each day that reporting entity ownership arrangements is late (§ 402.1(c)(22)).
   (2) HCFA or OIG may impose a penalty of not more than $10,000 for the following violations that occur on or after January 1, 1997:
      (i) Knowingly and willfully, and on a repeated basis, billing for a clinical diagnostic laboratory test, other than on an assignment-related basis (§ 402.1(c)(1)).
      (ii) By any durable medical equipment supplier, knowingly and willfully charging for a covered service that is furnished on a rental basis after the rental payments may no longer be made (except for maintenance and servicing) (§ 402.1(c)(4)).
      (iii) By any durable medical equipment supplier, knowingly and willfully, in violation of section 1834(a)(18)(A), failing to make a refund to Medicare beneficiaries for a covered service for which payment is precluded due to an unsolicited telephone contact from the supplier (§ 402.1(c)(5)).
      (iv) By any nonparticipating physician or supplier, knowingly and willfully charging a Medicare beneficiary more than the limiting charge, as specified in section 1834(b)(5)(B), for radiologist services (§ 402.1(c)(6)).
      (v) By any nonparticipating physician or supplier, knowingly and willfully charging a Medicare beneficiary more than the limiting charge, as specified in section 1834(c)(3), for mammography screening (§ 402.1(c)(7)).
      (vi) By any supplier of prosthetic devices, orthotics, and prosthetics, knowingly and willfully charging for a covered prosthetic device, orthotic, or prosthetic that is furnished on a rental basis after the rental payment may no longer be made (except for maintenance and servicing) (§ 402.1(c)(8)).
      (vii) By any supplier of durable medical equipment, including a supplier of prosthetic devices, orthotics, or supplies, knowingly and willfully failing to make refunds in a timely manner to Medicare beneficiaries for services billed other than on an assigned-related basis if—
         (A) The supplier does not possess a Medicare supplier number;
         (B) The service is denied in advance; or
         (C) The service is determined not to be medically necessary or reasonable (§ 402.1(c)(10)).
      (viii) Knowingly and willfully billing or collecting for any services on other than an assignment-related basis for practitioners specified in section 1842(b)(18)(B) (§ 402.1(c)(11)).
      (ix) By any physician, knowingly and willfully presenting, or causing to be presented, a claim or bill for an assistant at cataract surgery performed on or after March 1, 1987 for which payment may not be made because of section 1862(a)(15) (§ 402.1(c)(12)).
      (x) By any nonparticipating physician who does not accept payment on an assignment-related basis, knowingly and willfully failing to refund on a timely basis any amounts collected for services that are not reasonable or medically necessary or are of poor
quality, in accordance with section 1842(f)(1)(A) (§ 402.1(c)(13)).

(xi) By any nonparticipating physician, who does not accept payment for an elective surgical procedure on an assignment-related basis and whose charge is at least $500, knowingly and willfully failing to—
(A) Disclose the information required by section 1842(m)(1) concerning charges and coinsurance amounts; and
(B) Refund on a timely basis any amount collected for the procedure in excess of the charges recognized and approved by the Medicare program (§ 402.1(c)(14)).

(xii) By any physician, in repeated cases, knowingly and willfully billing one or more beneficiaries, for purchase diagnostic tests, any amount other than the payment amount specified in section 1842(n)(1)(A) or section 1842(m)(1)(B) (§ 402.1(c)(15)).

(xiii) By any nonparticipating physician, supplier, or other person that furnishes physicians' services and does not accept payment on an assignment-related basis—
(A) Knowingly and willfully billing or collecting in excess of the limiting charge (as defined in section 1843(g)(2)) on a repeated basis; or
(B) Failing to make an adjustment or refund on a timely basis as required by section 1848(g)(1)(A)(iii) or (iv) (§ 402.1(c)(17)).

(xiv) Knowingly and willfully billing for State plan approved physicians' services on other than an assignment-related basis for a Medicare beneficiary who is also eligible for Medicaid (§ 402.1(c)(18)).

(xv) By any supplier of durable medical equipment, including a supplier of prosthetic, orthotic, or supplies, knowingly and willfully failing to make refunds in a timely manner to Medicare beneficiaries for services billed on an assignment-related basis if—
(A) The supplier did not possess a Medicare supplier number;
(B) The service is denied in advance; or
(C) The service is determined not to be medically necessary or reasonable (§ 402.1(c)(23)).

(e) $15,000. HCFA or OIG may impose a penalty of not more than $15,000 if

(f) $25,000. HCFA or OIG may impose a penalty of not more than $25,000 for each of the following violations:

(1) Issuance of a Medicare supplemental policy that has not been approved by an approved State regulatory program or does not meet Federal standards on and after the effective date in section 1882(p)(1)(C) of the Act (§ 402.1(c)(23)).

(2) Sale or issuance after July 30, 1992, of a Medicare supplemental policy that fails to conform with the NAIC or Federal standards established under section 1882(p) of the Act (§ 402.1(c)(25)).

(3) Failure to make the core group of basic benefits available for sale when selling other Medicare supplemental plans with additional benefits (§ 402.1(c)(26)).

(4) Failure to provide, before sale of a Medicare supplemental policy, an outline of coverage describing the benefits provided by the policy (§ 402.1(c)(26)).

(5) Failure of an issuer of a policy to suspend or reinstate a policy, based on the policy holder's request, during entitlement to or upon loss of eligibility for medical assistance (§ 402.1(c)(27)).

(6) Failure to provide refunds or credits for Medicare supplemental policies as required by section 1882(r)(1)(B) (§ 402.1(c)(28)).

(7) By an issuer of a Medicare supplemental policy—
(i) Substantial failure to provide medically necessary services to enrollees seeking the services through the issuer's network of entities;
(ii) Imposition of premiums on enrollees in excess of the premiums approved by the State;
(iii) Action to expel an enrollee for reasons other than nonpayment of premiums; or
(iv) Failure to provide each enrollee, at the time of enrollment, with the specific information provided in section 1882(t)(1)(E)(i) or failure to obtain a written acknowledgment from the enrollee of receipt of the information (as required by section 1882(t)(1)(E)(ii) (section 1882(t)(2))).
§ 402.107 Amount of assessment.

A person subject to civil money penalties specified in § 402.1(c) may be subject, in addition, to an assessment. An assessment is a monetary payment in lieu of damages sustained by HHS or a State agency.

(a) The assessment may not be more than twice the amount claimed for each service that was a basis for the civil money penalty, except for the violations specified in paragraph (b) of this section that occur before January 1, 1997.

(b) For the violations specified in this paragraph occurring after January 1, 1997, the assessment may not be more than three times the amount claimed for each service that was the basis for a civil money penalty. The violations are the following:

(1) Knowingly and willfully billing, and on a repeated basis, for a clinical diagnostic laboratory test, other than on an assignment-related basis (§ 402.1(c)(1)).

(2) By any durable medical equipment supplier, knowingly and willfully charging for a covered service that is furnished on a rental basis after the rental payments may no longer be made (except for maintenance and servicing) as provided in section 1834(a)(7)(A) (§ 402.1(c)(4)).

(3) By any durable medical equipment supplier, knowingly and willfully failing, in violation of section 1834(a)(18)(A), to make a refund to Medicare beneficiaries for a covered service for which payment is precluded due to an unsolicited telephone contact from a supplier (§ 402.1(c)(5)).

(4) By any nonparticipating physician or supplier, knowingly and willfully charging a Medicare beneficiary more than the limiting charge as specified in section 1834(b)(5)(B), for radiologist services (§ 402.1(c)(6)).

(5) By any nonparticipating physician or supplier, knowingly and willfully charging a Medicare beneficiary more than the limiting charge as specified in section 1834(c)(3), for mammography screening (§ 402.1(c)(7)).

(6) By any supplier of prosthetic devices, orthotics, and prosthetics, knowingly and willfully charging for a covered prosthetic device, orthotic, or prosthetic that is furnished on a rental basis after the rental payment may no longer be made (except for maintenance and servicing) (§ 402.1(c)(8)).

(7) By any supplier of durable medical equipment, including a supplier of prosthetic devices, prosthetics, orthotics, or supplies, knowingly and willfully failing to make refunds in a timely manner to Medicare beneficiaries for services billed other than on an assignment-related basis if—

(i) The supplier does not possess a Medicare supplier number;

(ii) The service is denied in advance; or

(iii) The service is determined not to be medically necessary or reasonable (§ 402.1(c)(10)).

(8) Knowingly and willfully billing or collecting for any services on other than an assignment-related basis for practitioners specified in section 1842(b)(18)(B) (§ 402.1(c)(11)).

(9) By any physician, knowingly and willfully presenting, or causing to be presented, a claim or bill for an assistive device at cataract surgery performed on or after March 1, 1987 for which payment may not be made because of section 1862(a)(15) (§ 402.1(c)(12)).

(10) By any nonparticipating physician who does not accept payment on an assignment-related basis, knowingly and willfully failing to refund on a timely basis any amounts collected for services that are not reasonable or medically necessary or are of poor quality, in accordance with section 1842(l)(1)(A) (§ 402.1(c)(13)).

(11) By any nonparticipating physician, who does not accept payment for an elective surgical procedure on an assignment-related basis and whose charge is at least $500, knowingly and willfully failing to—

(i) Disclose the information required by section 1842(m)(1) concerning charges and coinsurance amounts; and

(ii) Defund on a timely basis any amount collected for the procedure in excess of the charges recognized and approved by the Medicare program (§ 402.1(c)(14)).

(12) By any physician, in repeated cases, knowingly and willfully billing one or more beneficiaries, for purchased diagnostic tests, any amount
other than the payment amount specified in section 1842(n)(1)(A) or section 1842(n)(1)(B) (§ 402.1(c)(15)).

(13) By any nonparticipating physician, supplier, or other person that furnishes physicians’ services and does not accept payment on an assignment-related basis—
(i) Knowingly and willfully billing or collecting in excess of the limiting charge (as defined in section 1843(g)(2)) on a repeated basis; or
(ii) Failing to make an adjustment or refund on a timely basis as required by section 1848(g)(1)(A) (iii) or (iv) (§ 402.1(c)(17)).

(14) Knowingly and willfully billing for State plan approved physicians’ services on other than an assignment-related basis for a Medicare beneficiary who is also eligible for Medicaid (§ 402.1(c)(18)).

(15) By any supplier of durable medical equipment, including suppliers of prosthetic devices, prosthetics, orthotics, or supplies, knowingly and willfully failing to make refunds in a timely manner to Medicare beneficiaries for services billed on an assignment-related basis if—
(i) The supplier did not possess a Medicare supplier number;
(ii) The service is denied in advance; or
(iii) The service is determined not to be medically necessary or reasonable (§ 402.1(c)(23)).

§ 402.109 Statistical sampling.

(a) Purpose. HCFA or OIG may introduce the results of a statistical sampling study to show the number and amount of claims subject to sanction under this part that the respondent presented or caused to be presented.

(b) Prima facie evidence. The results of the statistical sampling study, if based upon an appropriate sampling and computed by valid statistical methods, constitute prima facie evidence of the number and amount of claims or requests for payment subject to sanction under § 402.1.

(c) Burden of proof. Once HCFA or OIG has made a prima facie case, the burden is on the respondent to produce evidence reasonably calculated to rebut the findings of the statistical sampling study. HCFA or OIG then has the opportunity to rebut this evidence.

§ 402.111 Factors considered in determinations regarding the amount of penalties and assessments.

(a) Basic factors. In determining the amount of any penalty or assessment, HCFA or OIG takes into account the following:
(1) The nature of the claim, request for payment, or information given and the circumstances under which it was presented or given.
(2) The degree of culpability, history of prior offenses, and financial condition of the person submitting the claim or request for payment or giving the information.
(3) The resources available to the person submitting the claim or request for payment or giving the information.
(4) Such other matters as justice may require.

(b) Criteria to be considered. As guidelines for taking into account the factors listed in paragraph (a) of this section, HCFA or OIG considers the following circumstances:
(i) Aggravating circumstances of the incident. An aggravating circumstance is any of the following:
(1) The services or incidents were of several types, occurring over a lengthy period of time.
(2) There were many of these services or incidents or the nature and circumstances indicate a pattern of claims or requests for payment for these services or a pattern of incidents.
(3) The amount claimed or requested for these services was substantial.
(4) Before the incident or presentation of any claim or request for payment subject to imposition of a civil money penalty, the respondent was held liable for criminal, civil, or administrative sanctions in connection with a program covered by this part or any other public or private program of payment for medical services.
(v) There is proof that a respondent engaged in wrongful conduct, other than the specific conduct upon which liability is based, relating to government programs or in connection with the delivery of a health care service. (The statute of limitations governing
§ 402.113 When a penalty and assessment are collectible.

A civil money penalty and assessment become collectible after the earliest of the following:

(a) Sixty days after the respondent receives HCFA’s or OIG’s notice of proposed determination under §402.7, if the respondent has not requested a hearing before an ALJ.

(b) Immediately after the respondent abandons or waives his or her appeal right at any administrative level.

(c) Thirty days after the respondent receives the ALJ’s decision imposing a civil money penalty or assessment under §1005.20(d) of this title, if the respondent has not requested a review before the DAB.

(d) If the DAB grants an extension of the period for requesting the DAB’s review, the day after the extension expires if the respondent has not requested the review.

(e) Immediately after the ALJ’s decision denying a request for a stay of the effective date under §1005.22(b) of this title.

(f) If the ALJ grants a stay under §1005.22(b) of this title, immediately after the judicial ruling is completed.

(g) Sixty days after the respondent receives the DAB’s decision imposing a civil money penalty if the respondent has not requested a stay of the decision under §1005.22(b) of this title.

§ 402.115 Collection of penalty or assessment.

(a) Once a determination by HHS has become final, HCFA is responsible for the collection of any penalty or assessment.
(b) The General Counsel may compromise a penalty or assessment imposed under this part, after consultation with HCFA or OIG, and the Federal government may recover the penalty or assessment in a civil action brought in the United States district court for the district where the claim was presented or where the respondent resides.

(c) The United States or a State agency may deduct the amount of a penalty and assessment when finally determined, or the amount agreed upon in compromise, from any sum then or later owing to the respondent.

(d) Matters that were raised or that could have been raised in a hearing before an ALJ or in an appeal under section 1128A(e) of the Act may not be raised as a defense in a civil action by the United States to collect a penalty under this part.

Subpart C—Exclusions [Reserved]
§ 403.200 Basis and scope.

(a) Provisions of the legislation. This subpart implements, in part, section 1882 of the Social Security Act. The intent of that section is to enable Medicare beneficiaries to identify Medicare supplemental policies that do not duplicate Medicare, and that provide adequate, fairly priced protection against expenses not covered by Medicare. The legislation establishes certain standards for Medicare supplemental policies and provides two methods for informing Medicare beneficiaries which policies meet those standards:

(1) Through a State approved program, that is, a program that a Supplemental Health Insurance Panel determines to meet certain minimum requirements for the regulation of Medicare supplemental policies; and

(2) In a State without an approved program, through certification by the Secretary of policies voluntarily submitted by insuring organizations for review against the standards.

(b) Scope of subpart. This subpart sets forth the standards and procedures HCFA will use to implement the voluntary certification program.

§ 403.201 State regulation of insurance policies.

(a) The provisions of this subpart do not affect the right of a State to regulate policies marketed in that State.

(b) Approval of a policy under the voluntary certification program, as provided for in §403.235(b), does not authorize the insuring organization to market a policy that does not conform to applicable State laws and regulations.

§ 403.205 Medicare supplemental policy.

(a) Except as specified in paragraph (d) of this section, Medicare supplemental policy (policy) means a health insurance policy or other health benefit plan—

(1) That a private entity offers to a Medicare beneficiary; and

(2) That is primarily designed, or is advertised, marketed, or otherwise purported to provide payment for expenses incurred for services and items that are not reimbursed under the Medicare program because of deductibles, coinsurance, or other limitations under Medicare.

(b) Unless otherwise specified in this subpart, the term policy includes both policy form and policy.

(1) Policy form means the form of health insurance contract that is approved by and on file with the State agency for the regulation of insurance.

(2) Policy means the contract—

(i) Issued under the policy form; and

(ii) Held by the policyholder.

(c) Medicare supplemental policy includes the following—

(1) An individual policy.

(2) A group policy.

(d) Medicare supplemental policy does not include a Medicare+Choice plan or any of the following health insurance policies or health benefit plans:

(1) A policy or plan of one or more employers for employees, former employees, or any combination thereof.

(2) A policy or plan of one or more labor organizations for members, former members, or any combination thereof.
(3) A policy or plan of the trustees of a fund established by one or more labor organizations, one or more employers, or any combination, for any one or combination of the following—
   (i) Employees.
   (ii) Former employees.
   (iii) Members.
   (iv) Former members.
(4) A policy or plan of a profession, trade, or occupational association, if the association—
   (i) Is composed of individuals all of whom are actively engaged in the same profession, trade, or occupation;
   (ii) Has been maintained in good faith for a purpose other than obtaining insurance; and
   (iii) Has been in existence for at least two years before the date of its initial offering of a Medicare supplemental health insurance policy to its members.
(5) For purposes of the voluntary certification program, a policy issued to an employee or to a member of a labor organization as an addition to a franchise plan (a plan that enables members of the same entity to purchase an individual policy marketed to them under group underwriting procedures), if the plan is in existence on July 1, 1982.

§ 403.210 NAIC model standards.
(a) NAIC model standards means the National Association of Insurance Commissioners (NAIC) “Model Regulation to Implement the Individual Accident and Insurance Minimum Standards Act” (as amended and adopted by the NAIC on June 6, 1979, as it applies to Medicare supplemental policies). Copies of the NAIC model standards can be purchased from the National Association of Insurance Commissioners at 350 Bishops Way, Brookfield, Wisconsin 53004, and from the NIARS Corporation, 318 Franklin Avenue, Minneapolis, Minnesota 55404.
(b) The policy must comply with the provisions of the NAIC model standards, except as follows—
   (1) Policy, for purposes of this paragraph, means individual and group policy, as specified in §403.205. The NAIC model standards limit “policy” to individual policy.
   (2) The policy must meet the loss ratio standards specified in §403.215.

§ 403.215 Loss ratio standards.
(a) The policy must be expected to return to the policyholders, in the form of aggregate benefits provided under the policy—
   (1) At least 75 percent of the aggregate amount of premiums in the case of group policies; and
   (2) At least 60 percent of the aggregate amount of premiums in the case of individual policies.
(b) For purposes of loss ratio requirements, policies issued as a result of solicitation of individuals through the mail or by mass media advertising are considered individual policies.

STATE REGULATORY PROGRAMS

§ 403.220 Supplemental Health Insurance Panel.
(a) Membership. The Supplemental Health Insurance Panel (Panel) consists of—
   (1) The Secretary or a designee, who serves as chairperson, and
   (2) Four State Commissioners or Superintendents of Insurance appointed
§ 403.222 State with an approved regulatory program.

(a) A State has an approved regulatory program if the Panel determines that the State has in effect under State law a regulatory program that provides for the application of standards, with respect to each Medicare supplemental policy issued in that State, that are equal to or more stringent than those specified in section 1882 of the Social Security Act.

(b) Policy issued in that State means—

(1) A group policy, if the holder of the master policy resides in that State; and

(2) An individual policy, if the policy is—

(i) Issued in that State; or

(ii) Issued for delivery in that State.

(c) A policy issued in a State with an approved regulatory program is considered to meet the NAIC model standards in §403.210 and loss ratio standards in §403.215.

§ 403.232 Requirements and procedures for obtaining certification.

(a) To be certified by HCFA, a policy must meet—

(1) The NAIC model standards specified in §403.210;

(2) The loss ratio standards specified in §403.215; and

(3) Any State requirements applicable to a policy—

(i) Issued in that State; or

(ii) Marketed in that State.

(b) An insuring organization requesting certification of a policy must submit the following to HCFA for review—

(1) A copy of the policy form (including all the documents that would constitute the contract of insurance that is proposed to be marketed as a certified policy).

(2) A copy of the application form including all attachments.

(3) A copy of the uniform certificate issued under a group policy.

(4) A copy of the outline of coverage, in the form prescribed by the NAIC model standards.

(5) A copy of the Medicare supplement buyers’ guide to be provided to all applicants if the buyers’ guide is not the HCFA/NAIC buyers’ guide.

(6) A statement of when and how the outline of coverage and the buyers’ guide will be delivered and copies of applicable receipt forms.
(7) A copy of the notice of replacement and statement as to when and how that notice will be delivered.
(8) A list of States in which the policy is authorized for sale. If the policy was approved under a deemer provision in any State, the conditions involved must be specified.
(9) A copy of the loss ratio calculations, as specified in §403.250.
(10) Loss ratio supporting data, as specified in §403.256.
(11) A statement of actuarial opinion, as specified in §403.258.
(12) A statement that the insuring organization will notify the policyholders in writing, within the period of time specified in §403.245(c), if the policy is identified as a certified policy at the time of sale and later loses certification.
(13) A signed statement in which the president of the insuring organization, or a designee, attests that—
   (i) The policy meets the requirements specified in paragraph (a) of this section; and
   (ii) The information submitted to HCFA for review is accurate and complete and does not misrepresent any material fact.

§ 403.235 Review and certification of policies.
(a) HCFA will review policies that the insuring organization voluntarily submits, except that HCFA will not review a policy issued in a State with an approved regulatory program under §403.222.
(b) If the requirements specified in §403.232 are met, HCFA will—
   (1) Certify the policy; and
   (2) Authorize the insuring organization to display the emblem on the policy, as provided for in §403.231.
(c) If HCFA certifies a policy, it will inform all State Commissioners and Superintendents of Insurance of that fact.

§ 403.239 Submittal of material to retain certification.
(a) HCFA certification of a policy that continues to meet the requirements will remain in effect, if the insuring organization files the following material with HCFA no later than the date specified in paragraph (b) or (c) of this section—
   (1) Any changes in the material, specified in §403.232(b), that was submitted for previous certification.
   (2) The loss ratio supporting data specified in §403.256(b).
   (3) A signed statement in which the president of the insuring organization, or a designee, attests that—
      (i) The policy continues to meet the requirements specified in §403.232(a); and
      (ii) The information submitted to HCFA for review is accurate and complete and does not misrepresent any material fact.
(b) Except as specified in paragraph (c) of this section, the insuring organization must file the material with HCFA no later than June 30 of each year. The first time the insuring organization must file the material is no later than June 30 of the calendar year that follows the year in which HCFA—
   (1) Certifies a new policy; or
   (2) Certifies a policy that lost certification as provided in §403.245.
(c) If the loss ratio calculation period, used to calculate the expected loss ratio for the last actuarial certification submitted to HCFA, ends before the June 30 date of paragraph (b) of this section, the insuring organization must file the material with HCFA no later than the last day of that rate calculation period.

§ 403.245 Loss of certification.
(a) A policy loses certification if—
   (1) The insuring organization withdraws the policy from the voluntary certification program; or
   (2) HCFA determines that—
      (i) The policy fails to meet the requirements specified in §403.232(a); or
      (ii) The insuring organization has failed to meet the requirements for submittal of material specified in §403.239.
(b) If a policy loses its certification, HCFA will inform all State Commissioners and Superintendents of Insurance of that fact.
(c) If a policy that displays the emblem, or that has been marketed as a certified policy without the emblem, loses certification, the insuring organization must notify each holder of the
§ 403.248 Administrative review of HCFA determinations.

(a) This section provides for administrative review if HCFA determines—
(1) Not to certify a policy; or
(2) That a policy no longer meets the standards for certification.

(b) If HCFA makes a determination specified in paragraph (a) of this section, it will send a notice to the insuring organization containing the following information:

(1) That HCFA has made such a determination.
(2) The reasons for the determination.
(3) That the insuring organization has 30 days from the date of the notice to—
   (i) Request, in writing, an administrative review of the HCFA determination; and
   (ii) Submit additional information to HCFA for review.
(4) That, if the insuring organization requests an administrative review, HCFA will conduct the review, as provided for in paragraph (c) of this section.
(5) That, in a case involving loss of certification, the HCFA determination will go into effect 30 days from the date of the notice, unless the insurance organization requests an administrative review. If the insurance organization requests an administrative review, the policy retains its certification until HCFA makes a final determination.
(c) To calculate “present values”, the insurance organization may ignore discounting (an actuarial procedure that provides for the impact of a variety of factors, such as lapse of policies) for loss ratio calculation periods not exceeding 12 months.
§ 403.253 Calculation of benefits.

(a) General provisions. (1) Except as provided for in paragraph (a)(2) of this section, calculate the amount of “benefits” by—

(i) Adding the present values on the initial calculation date of—

(A) Expected incurred benefits in the loss ratio calculation period; and

(B) The total policy reserve at the last day of the loss ratio calculation period; and

(ii) Subtracting the total policy reserve on the initial calculation date from the sum of these values.

(2) To calculate the amount of “benefits” in the case of community or pool rated individual or group policies rerated on an annual basis, calculate the expected incurred benefits in the loss ratio calculation period.

(b) Calculation of total policy reserve—

(1) Option for calculation. The insuring organization must calculate “total policy reserve” according to the provisions of paragraph (b) (2) or (3) of this section.

(2) Total policy reserve: Federal provisions.

(i) “Total policy reserve” means the sum of—

(A) Additional reserve; and

(B) The reserve for future contingent benefits.

(ii) Additional reserve means the amount calculated on a net level reserve basis, using appropriate values to account for lapse, mortality, morbidity, and interest, that on the valuation date represents—

(A) The present value of expected incurred benefits over the loss ratio calculation period; less—

(B) The present value of expected net premiums over the loss ratio calculation period; and

(iii) Net premium means the level portion of the gross premium used in calculating the additional reserve. On the day the policy is issued, the present value of the series of those portions equals the present value of the expected incurred claims over the period that the gross premiums are computed to provide coverage.

(iv) Reserve for future contingent benefits means the amounts, not elsewhere included, that provide for the extension of benefits after insurance coverage terminates. These benefits—

(A) Are predicated on a health condition existing on the date coverage ends;

(B) Accrue after the date coverage ends; and

(C) Are payable after the valuation date.

(3) Total policy reserve: State provisions. “Total policy reserve” means the total policy reserve calculated according to appropriate State law or regulation.

§ 403.254 Calculation of premiums.

(a) General provisions. To calculate the amount of “premiums”, calculate the present value on the initial calculation date of expected earned premiums for the loss ratio calculation period.

(b) Specific provisions.

(1) Earned premium for a given period means—

(i) Written premiums for the period; plus—

(ii) The total premium reserve at the beginning of the period; less—

(iii) The total premium reserve at the end of the period.

(2) Written premiums in a period means—

(i) Premiums collected in that period; plus—

(ii) Premiums due and uncollected at the end of that period; less—

(iii) Premiums due and uncollected at the beginning of that period.

(3) Total premium reserve means the sum of—

(i) The unearned premium reserve;

(ii) The advance premium reserve; and

(iii) The reserve for rate credits.

(4) Unearned premium reserve means the portion of gross premiums due that provide for days of insurance coverage after the valuation date.

(5) Advance premium reserve means premiums received by the insuring organization that are due after the valuation date.

(6) Reserve for rate credits means rate credits on a group policy that—

(i) Accrued by the valuation date of the policy; and

(ii) Are paid or credited after the valuation date.
§ 403.256 Loss ratio supporting data.

(a) For purposes of requesting HCFA certification under §403.232, the insuring organization must submit the following loss ratio data to HCFA for review—

1. A statement of why the policy is to be considered, for purposes of the loss ratio standards, an individual or a group policy.
2. The earliest age at which policyholders can purchase the policy.
3. The general marketing method and the underwriting criteria used for the selection of applicants to whom coverage is offered.
4. What policies are to be included under the one policy form, by the dates the policies are issued.
5. The loss ratio calculation period.
6. The scale of premiums for the loss ratio calculation period.
7. The expected level of earned premiums in the loss ratio calculation period.
8. The expected level of incurred claims in the loss ratio calculation period.
9. A description of how the following assumptions were used in calculating the loss ratio.
   i. Morbidity.
   ii. Mortality.
   iii. Lapse.
   iv. Assumed increases in the Medicare deductible.
   v. Impact of inflation on reimbursement per service.
   vi. Interest.
   vii. Expected distribution, by age and sex, of persons who will purchase the policy in the coming year.
   viii. Expected impact on morbidity by policy duration of—
       A. The process used to select insureds from among those that apply for a policy; and
       B. Pre-existing condition clauses in the policy.

(b) For purposes of requesting continued HCFA certification under §403.239(a), the insuring organization must submit the following to HCFA—

1. A description of all changes in the loss ratio data, specified in paragraph (a) of this section, that occurred since HCFA last reviewed the policy.
2. The past loss ratio experience for the policy, including the experience of all riders and endorsements issued under the policy. The loss ratio experience data must include earned premiums, incurred claims, and total policy reserves that the insuring organization calculates—
   i. For all years of issue combined; and
   ii. Separately for each calendar year since HCFA first certified the policy.

§ 403.258 Statement of actuarial opinion.

(a) For purposes of certification requests submitted under §403.232(b) and subsequent review as specified in §403.239(a), statement of actuarial opinion means a signed declaration in which a qualified actuary states that the assumptions used in calculating the expected loss ratio are appropriate and reasonable, taking into account actual policy experience, if any, and reasonable expectations.

(b) Qualified actuary means—

1. A member in good standing of the American Academy of Actuaries; or
2. A person who has otherwise demonstrated his or her actuarial competence to the satisfaction of the Commissioner or Superintendent of Insurance of the domiciliary State of the insuring organization.

Subpart C—Recognition of State Reimbursement Control Systems

SOURCE: 51 FR 15492, Apr. 24, 1986, unless otherwise noted.

§ 403.300 Basis and purpose.

(a) Basis. This subpart implements section 1886(c) of the Act, which authorizes payment for Medicare inpatient hospital services in accordance with a State’s reimbursement control system rather than under the Medicare reimbursement principles as described in HCFA’s regulations and instructions.

(b) Purpose. Contained in this subpart are—

1. The basic requirements that a State reimbursement control system must meet in order to be approved by HCFA;
2. A description of HCFA’s review and evaluation procedures; and
§ 403.302 Definitions.

For purposes of this subpart—

Chief executive officer of a State means the Governor of the State or the Governor’s designee.

Existing demonstration project refers to demonstration projects approved by HCFA under the authority of section 402(a) of the Social Security Amendments of 1967 (42 U.S.C. 1395b-1) or section 222(a) of the Social Security Amendments of 1972 (42 U.S.C. 1395b-1 (note)) and in effect on April 20, 1983 (the date of the enactment of Pub. L. 98-21 (Social Security Amendments of 1983)).

Federal hospital means a hospital that is administered by, or that is under exclusive contract with, the Department of Defense, the Veterans Administration, or the Indian Health Service.

State system or system refers to a State reimbursement control system that is approved by HCFA under the authority of section 1886(c) of the Act and that satisfies the requirements described in this subpart.

§ 403.304 Minimum requirements for State systems—discretionary approval.

(a) Discretionary approval by HCFA. HCFA may approve Medicare payments under a State system, if HCFA determines that the system meets the requirements in paragraphs (b) and (c) of this section and, if applicable paragraph (d) of this section.

(b) Requirements for State system. (1) An application for approval of the system must be submitted to HCFA by the Chief Executive Officer of the State.

(2) The State system must apply to substantially all non-Federal acute care hospitals in the State.

(3) All hospitals covered by the system must have and maintain a utilization and quality control review agreement with a Peer Review Organization, as required under section 1866(a)(1)(F) of the Act and §466.78(a) of this chapter.

(4) Federal hospitals must be excluded from the State system.

(5) Nonacute care or specialty hospital (such as rehabilitation, psychiatric, or children’s hospitals) may, at the option of the State, be excluded from the State system.

(6) The State system must apply to at least 75 percent of all revenues or expenses—

(i) For inpatient hospital services in the State; and

(ii) For inpatient hospital services under the State’s Medicaid plan.

(7) Under the system, HMOs and competitive medical plans (CMPs), as defined by section 1876(b) of the Act and part 417 of this chapter, must be allowed to negotiate payment rates with hospitals.

(8) The system must limit hospital charges for Medicare beneficiaries to deductibles, coinsurance or non-covered services.

(9) Unless a waiver is granted by HCFA under §489.23 of this chapter, the system must prohibit payment, as required under section 1862(a)(14) of the Act and §405.310(m) of this chapter, for nonphysician services provided to hospital inpatients under Part B of Medicare.

(10) The system must require hospitals to submit Medicare cost reports or approved reports in lieu of Medicare cost reports as required.

(11) The system must require—

(i) Preparation, collection, or retention by the State of reports (such as financial, administrative, or statistical reports) that may be necessary, as determined by HCFA, to review and monitor the State’s assurances; and

(ii) Submission of the reports to HCFA upon request.

(12) The system must provide hospitals an opportunity to appeal errors that they believe have been made in the determination of their payment rates. The system, if it is prospective may not permit providers to file administrative appeals that would result in a retroactive revision of prospectively determined payment rates.

(c) Satisfactory assurances. The State must provide to HCFA satisfactory assurance as to the following:

(1) The system provides for equitable treatment of hospital patients and hospital employees.

(2) The system provides for equitable treatment of all entities that pay hospitals for inpatient hospital services,
including Federal and State programs. Under the requirement, the following conditions must be met:

(i) Both the Medicare and Medicaid programs must participate under the system.

(ii) The State must assure equitable and uniform treatment under the system of third-party payors of inpatient hospital services in terms of opportunity. Equitable opportunity must include, but need not be limited to, participation in the system and availability of discounts. Criteria under which discounts are made available must be equitably and uniformly applied to all payors, except for discounts negotiated by HMOs and CMPs. Discounts available to HMOs and CMPs as a result of their statutory right to negotiate payment rates independently of a State system, as described in paragraph (b)(7) of this section, need not be available to other payors.

(iii) The State must assure that all third-party payors that participate under the system share in the system’s risks and benefits.

(3) The amount of Medicare payments made under the system over 36-month periods may not exceed the amount of Medicare payment that would otherwise have been made under the Medicare principles of reimbursement for Medicare items and services had the State system not been in effect. States must submit the assurance and supporting data as required by §403.320 to document that the payment limit is not exceeded. States that have an existing Medicare demonstration project in effect on April 20, 1983, and that have requested approval of a State system under section 1886(c)(4) of the Act, may elect to have the effectiveness of the State system under this paragraph judged on the basis of the State system’s rate of increase or inflation in Medicare inpatient hospital payments as compared to the national rate of increase or inflation for such payments during the three cost reporting periods of the hospitals in the State beginning on or after October 1, 1983.

(d) Additional cost-effectiveness assurance. If the assurances and supporting data required under paragraph (c)(3) of this section are insufficient to provide assurance satisfactory to HCFA regarding the cost-effectiveness of a State system, the State may additionally submit one of the following assurances in order to meet the cost-effectiveness test:

(1) State responsibility for excess payments. The State must agree that each month Medicare intermediaries will disburse to the State’s hospital Federal funds that in the aggregate equal no more than would have been disbursed in the absence of the State system. Any additional funds necessary to pay hospitals for Medicare services required by the State system will be paid to the intermediaries by the State. These additional amounts will be refunded to the State by the intermediaries to the extent that, in subsequent months, the State system requires a smaller aggregate payment for Medicare services than would have been paid in the absence of the State system.

(2) Limitations on payments. (i) The State must agree that if its projections exceed what Medicare would pay in any particular period, the State and HCFA will establish and agreed upon payment schedule that will limit payments under the State system based on a predetermined percentage relationship between projected State payments and what payments would have been under Medicare.

(ii) If deviation from the predetermined relationship described in paragraph (d)(2)(i) of this section occurs, the State must further agree that—

(A) Medicare payments would be capped automatically at payment levels based on the rates used for the Medicare prospective payment system and the State would be required to pay the difference to individual hospitals in its system; or

(B) The State may provide by legislation or legally binding regulations that any reduced payments to hospitals under the system that result from this cost-effectiveness assurance will constitute full and final payment for hospital services furnished to Medicare beneficiaries for the period covered by these reduced payments.
§ 403.306 Additional requirements for State systems—mandatory approval.

(a) General policy—(1) Mandatory approval. HFCA will approve an application for Medicare reimbursement under a State system if the system meets all of the requirements of § 403.304 and of paragraph (b) of this section.

(2) Exception. HFCA may approve an application if the State system meets all of the requirements of § 403.304 but only some of the requirements of paragraph (b) of this section.

(b) Additional requirements—(1) Operation of system. The system must—

(i) Be operated directly by the State or by entity designated under State law;

(ii) Provide for payments to hospitals using a methodology under which—

(A) Prospectively determined payment rates are established; and

(B) Exceptions, adjustments, and methods for changes in methodology are set forth;

(iii) Provide that a change by the State in the system that has the effect of materially changing payments to hospitals can take effect only upon 60 days notice to HCFA and to the hospitals likely to be materially affected by the change and upon HCFA’s approval of the change.

(2) Satisfactory assurances—(i) Admissions practice. The State must assure that the operation of the system will not result in any change in hospital admission practices that result in—

(A) A significant reduction in the proportion of patients receiving hospital services covered under the system who have no third-party coverage and who are unable to pay for hospital services;

(B) A significant reduction in the proportion of individuals admitted to hospitals for inpatient hospital services for which payment is less, or is likely to be less, than the anticipated charges for or cost of the services;

(C) A refusal to admit patients who would be expected to require unusually costly or prolonged treatment for reasons other than those related to the appropriateness of the care available at the hospital; or

(D) A refusal to provide emergency services to any person who is in need of emergency services, if the hospital provides the services.

(ii) Consultation with local government officials. The State must provide documentation that it has consulted with local government officials concerning the impact of the system on publicly owned or operated hospitals.

§ 403.308 State systems under demonstration projects—mandatory approval.

HFCA will approve an application from a State for a State system if—

(a) The system was in effect prior to April 20, 1983 under an existing demonstration project; and

(b) The minimum requirements and assurances for approval of a State system are met under § 403.304(b)(1)–(10) and § 403.304(c), and, if appropriate § 403.304(d).

§ 403.310 Reduction in payments.

(a) General rule. If HCFA determines that the satisfactory assurances required of a State under § 403.304(c) and, if applicable, § 403.304(d) have not been met, or will not be met, with respect to any 36-month period, HCFA will reduce Medicare payments to individual hospitals being reimbursed under the State’s system or, if applicable, under the Medicare payment system, in an amount equal to the amount by which the Medicare payments under the system exceed the amount of Medicare payments to such hospitals that otherwise would have been made not using the State system. The amount of the recoupment will include, when appropriate, interest charges computed in accordance with § 405.378 of this chapter.

(b) Recoupment procedures. The amount of the overpayment will be recouped on a proportionate basis from each of those hospitals that received payments under the State system that exceeded the payments they would have received under the Medicare payment system. Each hospital’s share of the aggregate excess payment will be determined on the basis of a comparison of the hospital’s proportionate share of the aggregate payment received under the State system that is
§ 403.312 Submittal of application.

The Chief Executive Officer of the State is responsible for—

(a) Submittal of the application to HCFA for approval; and

(b) Supplying the assurances and necessary documentation as required under §§403.304 through 403.308.

§ 403.314 Evaluation of State systems.

HCFA will evaluate all State applications for approval of State systems and notify the State of its determination within 60 days.

§ 403.316 Reconsideration of certain denied applications.

(a) Request for reconsideration. If HCFA denies an application for a State system, the State may request that HCFA reconsider the denial if the State believes that its system meets all of the requirements for mandatory approval under §§403.304 and 403.306 or, in the case of a State with a system operating under an existing demonstration project, the applicable requirements of §§403.304 and 403.308.

(b) Time limit. (1) The State must submit its request for reconsideration within 60 days after the date of HCFA’s notice that the application was denied.

(2) HCFA will notify the State of the results of its reconsideration within 60 days after it receives the request for reconsideration.

§ 403.318 Approval of State systems.

(a) Approval agreement. If HCFA approves a State system, a written agreement will be executed between HCFA and the Chief Executive Officer of the State. The agreement must incorporate any terms of the State's application for approval of the system as agreed to by the parties and, as a minimum, must contain provisions that require the following:

(1) The system is operated directly by the State or an entity designated by State law.

(2) For purposes of the Medicare program, the State's system applies only to Medicare payments for inpatient, and if applicable, outpatient hospital services.

(3) The system conforms to applicable Medicare law and regulations other than those relating to the amount of reimbursement for inpatient hospital services, or for inpatient and outpatient services, whichever the State system covers. Applicable regulations include, for example, those describing Medicare benefits and entitlement requirements for program beneficiaries, as explained in parts 406 and 409 of this chapter; the requirements at part 405, subpart J of this chapter specifying conditions of participation for hospitals; the requirements at part 405, subparts A, G, and S of this chapter on Medicare program administration; and all applicable fraud and abuse regulations contained in titles 42 and 45 of the CFR.

(4) The State must obtain HCFA's approval of the State's reporting forms and of provider cost reporting forms or other forms that have not been approved by HCFA but that are necessary for the collection of required information.
(b) Effective date. An approved State system may not be effective earlier than the date of the approval agreement, which may not be retroactive.

§ 403.320 HCFA review and monitoring of State systems.

(a) General rule. The State must submit an assurance and detailed and quantitative studies of provider cost and financial data and projections to support the effectiveness of its system, as required by paragraphs (b) and (c) of this section.

(b) Required information. (1) Under § 403.304(c)(3) an assurance is required that the system will not result in greater payments over a 36-month period than would have otherwise been made under Medicare not using such system. If a State that has an existing demonstration project in effect on April 20, 1983 elects under § 403.304(c)(3) to have the effectiveness of its system judged on the basis of a rate of increase factor, the State must submit an assurance that its rate of increase or inflation in inpatient hospital payments does not exceed, for that portion of the 36-month period that is subject to this test, the national rate of increase or inflation in Medicare inpatient hospital payments. The election of the rate of increase test applies only to the three cost reporting periods beginning on or after October 1, 1983. At the end of these cost reporting periods, the State must assure, beginning with the first month after the expiration of the third cost reporting period beginning after October 1, 1983, that payments under its system will not exceed over the remainder of the 36-month period what Medicare payments would have been.

(2) Estimates and data are required to support the State's assurance, required under § 403.304(c)(3), that expenditures under the State system will not exceed what Medicare would have paid over a 36-month period. The estimates and projections of what Medicare would have otherwise paid must take into account all the Medicare reimbursement principles in effect at the time and, for any period in which payments either exceed or are less than Medicare levels, the values of interest the Medicare Trust Fund earned, or would have earned, on these amounts. Upon application for approval, the State must submit projections for each hospital for the first 12-month period covered by the assurance, in both the aggregate and on a per discharge basis, of Medicare inpatient expenditures under Medicare principles of reimbursement and parallel projections of Medicare inpatient expenditures under the State's system and the resulting cost or savings to Medicare. The State must also submit separate statewide projections for each year of the 36-month period, in both the aggregate and on a weighted average discharge basis, of inpatient expenditures under the State system and under the Medicare principles of reimbursement.

(3) The projection submitted under paragraph (b)(2) of this section must include a detailed description of the methodology and assumptions used to derive the expenditure amounts under both systems. In instances where the assumptions are different under the projections cited in paragraph (b)(2) of this section, the State must provide a detailed explanation of the reasons for the differences. At a minimum, the following separate data and assumptions are to be included in the projections for the Medicare principles and for the State's system.

(i) The State system base year and the Medicare allowable and reimbursable cost of each hospital that the State used to develop the projections, including the amount of estimated pass through costs.

(ii) The categories of costs that are included in the State system and are reimbursed differently under the State system than under the Medicare system.

(iii) The number of Medicare and total base year discharges and admissions for each hospital.

(iv) The rate of change factor (and the method of application of this factor) used to project the base year costs over the 36-month period to which the assurance would apply.

(v) Any allowance for anticipated growth in the amount of services from the base year (if applicable, the allowance must be presented in separate estimates for population increases or for
increases in rates of admissions or both).

(vi) Any adjustment in which the State is permitted by HCFA to take into account previous reductions in the Medicare payment amounts that were the result of the effectiveness of the State's system even though Medicare was not a part of that system.

(vii) Appropriate recognition and projection of the time value of trust fund expenditures for the period the State system expenditures were either less than or exceeded the Medicare system payments.

(viii) States applying under a rate of increase effectiveness test under §403.304(c)(3) must also submit data projecting the parallel rates of increase during the requisite period.

(4) The projections must include both the aggregate payments and the payments per discharge for the individual hospitals and for the State as a whole.

(5) On a case-by-case basis, HCFA may require additional data and documentation as needed to complete its review and monitoring.

(6) For existing Medicare demonstration projects in effect on April 20, 1983, the assurance and data as required by paragraphs (a) and (b) of this section, if appropriate, may be based on aggregate payments or payments per inpatient admission or discharge. HCFA will judge the effectiveness of these systems on the basis of the rate of increase or inflation in Medicare inpatient hospital payments compared to the national rate of increase or inflation for such payments during the State's hospitals' three cost reporting periods beginning on or after October 1, 1983. The data submitted by the State for this period subject to the rate of increase test must include the rate of increase projection for that particular period of time. For the subsequent period of time, the State must assure that payments under its system will not exceed what Medicare payments would have been, as described in §403.304(c)(3).

(7) If the amount of Medicare payments under the State system exceeds what would have been paid under the Medicare reimbursement principles in any given year, the State must also submit quantitative evidence that the system will result in expenditures that do not exceed what Medicare expenditures would have been over the 36 month period beginning with the first month that the State system is operating. For a State that has an existing demonstration project in effect on April 20, 1983, and that elects under §403.304(c)(3) to have a rate of increase test apply, if the State's rate of increase or inflation exceeds the national rate of increase or inflation in a given year, the State must submit quantitative evidence that, over 36 months, its payments will not exceed the national rate of increase or inflation. Furthermore, if payments under the State's system must be compared to actual Medicare expenditures, at the end of the third cost reporting period, as described in paragraph (b)(1) of this section, and payments under the State's system exceed what Medicare would have paid in a given year, the State must submit quantitative evidence that, over 36 months, payments under its system will not exceed what Medicare would have paid.

(c) Review of assurances regarding expenditures. HCFA will review the State's assurances and data submitted under this section, as a prerequisite to the approval of the State's system. HCFA will compare the State's projections of payment amounts to HCFA data in order to determine if the State's assurance is reasonable and fully supportable. If the HCFA data indicate that the State's system would result in payment amounts that would be more than that which would have been paid under the Medicare principles, the State's assurances would not be acceptable. For States applying in accordance with §403.308, if HCFA data indicate that the State's system would result in a rate of increase or inflation that would be more than the national rate of increase or inflation, the State's assurances would not be acceptable.

(d) Medicaid upper limit. In accordance with §447.253 of this chapter, the State system may not result in aggregate payments for Medicaid inpatient hospital services that would exceed the amount that would have otherwise been paid under the Medicare
(e) Monitoring of Medicare expenditures. HCFA will monitor on a quarterly basis expenditures under the State's system as compared to what Medicare expenditures would have been if the system had not been in effect. If HCFA determines at any time that the payments made under the State's system exceed the States' projections, as established by the satisfactory assurances required under §403.304(c) and, if appropriate, the predetermined percentage relationship of the payments as required under §403.304(d), HCFA will—

1. Conclude that payments under the State system over a 36-month period will exceed what Medicare would have paid;
2. Terminate the waiver; and
3. Recoup overpayments to the affected hospitals in accordance with the procedures described in §403.310.

§ 403.321 State systems for hospital outpatient services.

HCFA may approve a State's application for approval of an outpatient system if the following conditions are met:

(a) The State's inpatient system is approved.

(b) The State's outpatient application meets the requirements and assurances for an inpatient system described in §403.304(b) and (c), and §403.306(b)(1) and (b)(2)(ii).

(c) The State submits a separate application that provides separate assurances and estimates and data in further support of its assurance submitted under paragraph (b)(1) of §403.320, as follows:

1. Upon application for approval, the State must submit estimates and data that include, but are not limited to, projections for the first 12-month period covered by the assurance for each hospital, in both the aggregate and on an average cost per service and payment basis, of Medicare outpatient expenditures under Medicare principles of reimbursement; parallel projections of Medicare outpatient expenditures under the State system; and the resulting cost or savings to Medicare independent of the State system for hospital inpatient services.

2. The State must submit separate statewide projections for each year of the 36-month period of the aggregate outpatient expenditures for each system. The projections submitted under this paragraph must—

   (i) Comply with the requirements of paragraphs (b)(3) and (5) of §403.320 regarding a detailed description of the methodology used to derive the expenditure amounts;

   (ii) Include the data and assumptions set forth in paragraphs (b)(3)(i), (ii), (iii), (iv), and (v) of §403.320 and

   (iii) Include any assumption the State has adopted for establishing the number of Medicare and total base year outpatient services for each hospital.

3. The State must provide a detailed explanation of the reasons for any difference between the data or assumptions used for the separate projections.

§ 403.322 Termination of agreements for Medicare recognition of State systems.

(a) Termination of agreements. (1) HCFA may terminate any approved agreement if it finds, after the procedures described in this paragraph are followed, that the State system does not satisfactorily meet the requirements of section 1886(c) of the Act or the regulations in this subpart. A termination must be effective on the last day of a calendar quarter.

(2) HCFA will give the State reasonable notice of the proposed termination of an agreement and of the reasons for the termination at least 90 days before the effective date of the termination.

(3) HCFA will give the State the opportunity to present evidence to refute the finding.

(4) HCFA will issue a final notice of termination upon a final review and determination on the State's evidence.

(b) Termination by State. A State may voluntarily terminate a State system by giving HCFA notice of its intent to terminate. A termination must be effective on the last day of a calendar quarter. The State must notify HCFA of its intent to terminate at least 90 days before the effective date of the termination.
Subpart D—[Reserved]

Subpart E—Beneficiary Counseling and Assistance Grants

SOURCE: 59 FR 51128, Oct. 7, 1994, unless otherwise noted.

§ 403.500 Basis, scope, and definition.

(a) Basis. This subpart implements, in part, the provisions of section 4360 of Public Law 101-508 by establishing a minimum level of funding for grants made to States for the purpose of providing information, counseling, and assistance relating to obtaining adequate and appropriate health insurance coverage to individuals eligible to receive benefits under the Medicare program.

(b) Scope of subpart. This subpart sets forth the following:

(1) Conditions of eligibility for the grant.
(2) Minimum levels of funding for those States qualifying for the grants.
(3) Reporting requirements.

(c) Definition. For purposes of this subpart, the term “State” includes (except where otherwise indicated by the context) the 50 States, the District of Columbia, the Commonwealth of Puerto Rico, the Virgin Islands, Guam, and American Samoa.

§ 403.501 Eligibility for grants.

To be eligible for a grant under this subpart, the State must have an approved Medicare supplemental regulatory program under section 1882 of the Act and submit a timely application to HCFA that meets the requirements of—

(a) Section 4360 of Public Law 101-508 (42 USC 1395b-4);
(b) This subpart; and
(c) The applicable solicitation for grant applications issued by HCFA.

§ 403.502 Availability of grants.

HCFA awards grants to States subject to availability of funds, and if applicable, subject to the satisfactory progress in the State's project during the preceding grant period. The criteria by which progress is evaluated and the performance standards for determining whether satisfactory progress has been made are specified in the terms and conditions included in the notice of grant award sent to each State. HCFA advises each State as to when to make application, what to include in the application, and provides information as to the timing of the grant award and the duration of the grant award. HCFA also provides an estimate of the amount of funds that may be available to the State.

[65 FR 34985, June 1, 2000]

§ 403.504 Number and size of grants.

(a) General. For available grant funds, up to and including $10,000,000, grants will be made to States according to the terms and formula in paragraphs (b) and (c) of this section. For any available grant funds in excess of $10,000,000, distribution of grants will be at the discretion of HCFA, and will be made according to criteria that HCFA will communicate to the States via grant solicitation. HCFA will provide information to each State as to what must be included in the application for grant funds. HCFA awards the following type of grants:

(1) New program grants.
(2) Existing program enhancement grants.

(b) Grant Award. Subject to the availability of funds, each eligible State that submits an acceptable application receives a grant that includes a fixed amount (minimum funding level) and a variable amount.

(1) A fixed portion is awarded to States in the following amounts:
(i) Each of the 50 States, $75,000.
(ii) The District of Columbia, $75,000.
(iii) Puerto Rico, $75,000.
(iv) American Samoa, $25,000.
(v) Guam, $25,000.
(vi) The Virgin Islands, $25,000.

(2) A variable portion, which is based on the number and location of Medicare beneficiaries residing in the State is awarded to each State. The variable amount a particular State receives is determined as set forth in paragraph (c) of this section.

(c) Calculation of variable portion of the grant. (1) HCFA bases the variable portion of the grant on—

(i) The amount of available funds, and
(ii) A comparison of each State with the average of all of the States (except
Health Care Financing Administration, HHS

§ 403.512 Administration.

(a) General. Administration of grants will be in accordance with the provisions of this subpart, 45 CFR part 92 ("Uniform Administrative Requirements for Grants and Cooperative Agreements to State and Local Governments"), 45 CFR 74.4, the terms of the solicitation, and the terms of the notice of grant award. Except for the minimum funding levels established by §403.504(b)(1), in the event of conflict between a provision of the notice of grant award, any provision of the solicitation, or of any regulation enumerated in 45 CFR 74.4 or in part 92, the terms of the notice of grant award control.

(b) Notice. HCFA provides notice to each applicant regarding HCFA’s decision on an application for grant funding under §403.504.

(c) Appeal. Any applicant for a grant under this subpart has the right to appeal HCFA’s determination regarding its application. Appeal procedures are governed by the regulations at 45 CFR part 16 (Procedures of the Departmental Grant Appeals Board).
Subpart F—Reserved

Subpart G—Religious Nonmedical Health Care Institutions—Benefits, Conditions of Participation, and Payment

SOURCE: 64 FR 67047, Nov. 30, 1999, unless otherwise noted.

§ 403.700 Basis and purpose.

This subpart implements sections 1821, 1861(e), (y), and (ss); 1869; and 1878 of the Act regarding Medicare payment for inpatient hospital or posthospital extended care services furnished to eligible beneficiaries in religious nonmedical health care institutions.

§ 403.702 Definitions and terms.

For purposes of this subpart, the following definitions and terms apply:

Election means a written statement signed by the beneficiary or the beneficiary's legal representative indicating the beneficiary's choice to receive nonmedical care or treatment for religious reasons.

Excepted medical care means medical care that is received involuntarily or required under Federal, State, or local laws.

FFY stands for Federal fiscal year.

Medical care or treatment means health care furnished by or under the direction of a licensed physician that can involve diagnosing, treating, or preventing disease and other damage to the mind and body. It may involve the use of pharmaceuticals, diet, exercise, surgical intervention, and technical procedures.

Nonexcepted medical care means medical care (other than excepted medical care) that is sought by or for a beneficiary who has elected religious nonmedical health care institution services.

Religious nonmedical care or religious method of healing means health care furnished under established religious tenets that prohibit conventional or unconventional medical care for the treatment of a beneficiary, and the sole reliance on these religious tenets to fulfill the beneficiary's total health care needs.

RNHCI stands for “religious nonmedical health care institution,” as defined in section 1861(ss)(1) of the Act.

Religious nonmedical nursing personnel means individuals who are grounded in the religious beliefs of the RNHCI, trained and experienced in the principles of nonmedical care, and formally recognized as competent in the administration of care within their religious nonmedical health care group.

§ 403.720 Conditions for coverage.

Medicare covers services furnished in an RNHCI if the following conditions are met:

(a) The provider meets the definition of an RNHCI as defined in section 1861(ss)(1) of the Act. That is, it is an institution that:

(1) Is described in section 501(c)(3) of the Internal Revenue Code of 1986 and is exempt from taxes under section 501(a).

(2) Is lawfully operated under all applicable Federal, State, and local laws and regulations.

(3) Furnishes only nonmedical nursing items and services to beneficiaries who choose to rely solely upon a religious method of healing and for whom the acceptance of medical services would be inconsistent with their religious beliefs.

(4) Furnishes nonmedical items and services exclusively through nonmedical nursing personnel who are experienced in caring for the physical needs of nonmedical patients.

(5) Furnishes nonmedical items and services to inpatients on a 24-hour basis.

(6) Does not furnish, on the basis of religious beliefs, through its personnel or otherwise medical items and services (including any medical screening, examination, diagnosis, prognosis, treatment, or the administration of drugs) for its patients.

(7) Is not owned by, is not under common ownership with, or does not have an ownership interest of 5 percent or more in, a provider of medical treatment or services and is not affiliated with a provider of medical treatment or services or with an individual who has an ownership interest of 5 percent...
or more in a provider of medical treatment or services. (Permissible affiliations are described at §403.738(c).)

(8) Has in effect a utilization review plan that sets forth the following:
   (i) Provides for review of the admissions to the institution, the duration of stays, and the need for continuous extended duration of stays in the institution, and the items and services furnished by the institution.
   (ii) Requires that reviews be made by an appropriate committee of the institution that included the individuals responsible for overall administration and for supervision of nursing personnel at the institution.
   (iii) Provides that records be maintained of the meetings, decisions, and actions of the review committee.
   (iv) Meets other requirements as the Secretary finds necessary to establish an effective utilization review plan.

(9) Provides information HCFA may require to implement section 1821 of the Act, including information relating to quality of care and coverage decisions.

(10) Meets other requirements HCFA finds necessary in the interest of the health and safety of the patients who receive services in the institution. These requirements are the conditions of participation in this subpart.

(b) The provider meets the conditions of participation cited in §§403.730 through 403.746. (A provider may be deemed to meet conditions of participation in accordance with part 488 of this chapter.)

(c) The provider has a valid provider agreement as a hospital with HCFA in accordance with part 489 of this chapter and for payment purposes is classified as an extended care hospital.

(d) The beneficiary has a condition that would make him or her eligible to receive services covered under Medicare Part A as an inpatient in a hospital or SNF.

(e) The beneficiary has a valid election as described in §403.724 in effect for Medicare covered services furnished in an RNHCI.

§ 403.724 Valid election requirements.

(a) General requirements. An election statement must be made by the Medicare beneficiary or his or her legal representative.

1. The election must be a written statement that must include the following statements:
   (i) The beneficiary is conscientiously opposed to acceptance of nonexcepted medical treatment.
   (ii) The beneficiary acknowledges that the acceptance of nonexcepted medical treatment is inconsistent with his or her sincere religious beliefs.
   (iii) The beneficiary acknowledges that the receipt of nonexcepted medical treatment constitutes a revocation of the election and may limit further receipt of services in an RNHCI.
   (iv) The beneficiary acknowledges that the election may be revoked by submitting a written statement to HCFA.
   (v) The beneficiary acknowledges that revocation of the election will not prevent or delay access to medical services available under Medicare Part A in facilities other than RNHCIs.

2. The election must be signed and dated by the beneficiary or his or her legal representative.

3. The election must be notarized.

4. The RNHCI must keep a copy of the election statement on file and submit the original to HCFA with any information obtained regarding prior elections or revocations.

5. The election becomes effective on the date it is signed.

6. The election remains in effect until revoked.

(b) Revocation of election. (1) A beneficiary’s election is revoked by one of the following:

(i) The beneficiary receives nonexcepted medical treatment for which Medicare payment is requested.
   (ii) The beneficiary voluntarily revokes the election and notifies HCFA in writing.

(2) The receipt of excepted medical treatment as defined in §403.702 does not revoke the election made by a beneficiary.

(c) Limitation on subsequent elections. (1) If a beneficiary’s election has been made and revoked twice, the following limitations on subsequent elections apply:

1. The beneficiary is not eligible for services in an RNHCI.

2. The beneficiary is not eligible for services in an RNHCI as an inpatient.

3. The beneficiary is not eligible for services in an RNHCI as an outpatient.

4. The beneficiary is not eligible for services in an RNHCI as a skilled nursing facility.

5. The beneficiary is not eligible for services in an RNHCI as a intermediate care facility.

6. The beneficiary is not eligible for services in an RNHCI as a long-term care facility.
§ 403.730 Condition of participation: Patient rights.

An RNHCI must protect and promote each patient’s rights.

(a) Standard: Notice of rights. The RNHCI must do the following:

(1) Inform each patient of his or her rights in advance of furnishing patient care.

(2) Have a process for prompt resolution of grievances, including a specific person within the facility whom a patient may contact to file a grievance. In addition, the facility must provide patients with information about the facility’s process as well as with contact information for appropriate State and Federal resources.

(b) Standard: Exercise of rights. The patient has the right to:

(1) Be informed of his or her rights and to participate in the development and implementation of his or her plan of care.

(2) Make decisions regarding his or her care, including transfer and discharge from the RNHCI. (See §403.736 for discharge and transfer requirements.)

(3) Formulate advance directives and expect staff who furnish care in the RNHCI to comply with those directives, in accordance with part 489, subpart I of this chapter. For purposes of conforming with the requirement in §489.102 that there be documentation in the patient’s medical records concerning advanced directives, the patient care records of a beneficiary in an RNHCI are equivalent to medical records held by other providers.

(c) Standard: Privacy and safety. The patient has the right to:

(1) Personal privacy.

(2) Care in a safe setting.

(3) Freedom from verbal, psychological, and physical abuse, and misappropriation of property.

(4) Freedom from the use of restraints.

(5) Freedom from involuntary seclusion.

(d) Standard: Confidentiality of patient records. For any patient care records or election information it maintains on patients, the RNHCI must establish procedures to do the following:

(1) Safeguard the privacy of any information that identifies a particular patient. Information from, or copies of, records may be released only to authorized individuals, and the RNHCI must ensure that unauthorized individuals cannot gain access to or alter patient records. Original patient care records must be released only in accordance with Federal or State laws, court orders, or subpoenas.

(2) Maintain the records and information in an accurate and timely manner.

(3) Ensure timely access by patients to the records and other information that pertains to that patient.

(4) Abide by all Federal and State laws regarding confidentiality and disclosure for patient care records and election information.

§ 403.732 Condition of participation: Quality assessment and performance improvement.

The RNHCI must develop, implement, and maintain a quality assessment and performance improvement program.

(a) Standard: Program scope. (1) The quality assessment and performance improvement program must include, but is not limited to, measures to evaluate:

(i) Access to care.

(ii) Patient satisfaction.

(iii) Staff performance.

(iv) Complaints and grievances.

(v) Discharge planning activities.

(vi) Safety issues, including physical environment.

(2) In each of the areas listed in paragraph (a)(1) of this section, and any other areas the RNHCI includes, the RNHCI must do the following:

(i) Define quality assessment and performance improvement measures.
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§ 403.736 Condition of participation: Discharge planning.

The RNHCI must have in effect a discharge planning process that applies to all patients. The process must assure that appropriate post-institution services are obtained for each patient, as necessary.

(a) Standard: Discharge planning evaluation. (1) The RNHCI must assess the need for a discharge plan for any patient identified as likely to suffer adverse consequences if there is no planning and for any other patient upon his or her request or at the request of his or her legal representative. This discharge planning evaluation must be initiated at admission and must include the following:
   (i) An assessment of the possibility of a patient needing post-RNHCI services and of the availability of those services.
   (ii) An assessment of the probability of a patient’s capacity for self-care or of the possibility of the patient being cared for in the environment from which he or she entered the RNHCI.

(b) Standard: Discharge plan. (1) If the discharge planning evaluation indicates a need for a discharge plan, qualified and experienced personnel must develop or supervise the development of the plan.

(ii) Describe and outline quality assessment and performance improvement activities appropriate for the services furnished by or in the RNHCI.

(iii) Measure, analyze, and track performance that reflect care and RNHCI processes.

(iv) Inform all patients, in writing, of the scope and responsibilities of the quality assessment and performance improvement program.

(3) The RNHCI must set priorities for performance improvement, considering the prevalence of and severity of identified problems.

(4) The RNHCI must act to make performance improvements and must track performance to assure that improvements are sustained.

(b) Standard: Program responsibilities.

(1) The governing body, administration, and staff are responsible for ensuring that the quality assessment and performance improvement program addresses identified priorities in the RNHCI and are responsible for the development, implementation, maintenance, and performance improvement of assessment actions.

(2) The RNHCI must include all programs, departments, functions, and contracted services when developing, implementing, maintaining, and evaluating the program of quality assessment and performance improvement.

§ 403.734 Condition of participation: Food services.

The RNHCI must have an organized food service that is directed and adequately staffed by qualified personnel.

(a) Standard: Sanitary conditions. The RNHCI must furnish food to the patient that is obtained, stored, prepared, distributed, and served under sanitary conditions.

(b) Standard: Meals. The RNHCI must serve meals that furnish each patient with adequate nourishment in accordance with the recommended dietary allowances of the Food and Nutrition Board of the National Research Council, National Academy of Sciences. The RNHCI must do the following:

1. Furnish food that is palatable, attractive, and at the proper temperature and consistency.

2. Offer substitutes of similar nourishment to patients who refuse food served or desire alternative choices.

3. Furnish meals at regular times comparable to normal mealtimes in the community. There must be no more than 14 hours between a substantial evening meal and breakfast the following day.

4. The RNHCI must offer snacks at bedtime.

§ 403.736 Condition of participation: Food services.

The RNHCI must have in effect a discharge planning process that applies to all patients. The process must assure that appropriate post-institution services are obtained for each patient, as necessary.

(a) Standard: Discharge planning evaluation. (1) The RNHCI must assess the need for a discharge plan for any patient identified as likely to suffer adverse consequences if there is no planning and for any other patient upon his or her request or at the request of his or her legal representative. This discharge planning evaluation must be initiated at admission and must include the following:

(i) An assessment of the possibility of a patient needing post-RNHCI services and of the availability of those services.

(ii) An assessment of the probability of a patient’s capacity for self-care or of the possibility of the patient being cared for in the environment from which he or she entered the RNHCI.

(2) The staff must complete the assessment on a timely basis so that arrangements for post-RNHCI care are made before discharge and so that unnecessary delays in discharge are avoided.

(3) The discharge planning evaluation must be included in the patient’s rights record for use in establishing an appropriate discharge plan and must discuss the results of the evaluation with the patient or a legal representative acting on his or her behalf.

(b) Standard: Discharge plan. (1) If the discharge planning evaluation indicates a need for a discharge plan, qualified and experienced personnel must develop or supervise the development of the plan.
§ 403.738 Condition of participation: Administration.

An RNHCI must have written policies regarding its organization, services, and administration.

(a) Standard: Compliance with Federal, State, and local laws. The RNHCI must operate in compliance with all applicable Federal, State, and local laws, regulations, and codes including, but not limited to, those pertaining to the following:

(1) Protection against discrimination on the basis of race, color, national origin, age, or handicap (45 CFR parts 80, 84, and 91).

(2) Protection of human research subjects (45 CFR part 46).

(3) Application of all safeguards to protect against the possibility of fraud and abuse (42 CFR part 455).

(b) Standard: Governing body. (1) The RNHCI must have a governing body, or a person designated to function as a governing body, that is legally responsible for establishing and implementing all policies regarding the RNHCI's management and operation.

(2) The governing body must appoint the administrator responsible for the management of the RNHCI.

(c) Standard: Affiliations and disclosure. (1) An affiliation is permissible if it is between one of the following:

(i) An individual serving as an uncompensated director, trustee, officer, or staff member of an RNHCI and another individual, with whom he or she has a family relationship, who is affiliated with (or has an ownership interest in) a provider of medical treatment or services.

(ii) An individual who is a director, trustee, officer, employee, or staff member of an RNHCI and another individual, with whom he or she has a family relationship, who is affiliated with (or has an ownership interest in) a provider of medical treatment or services.

(iii) The RNHCI and an individual or entity furnishing goods or services as a vendor to both providers of medical treatment or services and RNHCIs.

(2) The RNHCI complies with the disclosure requirements of §§ 420.206 and 455.104 of this chapter.

(3) The RNHCI furnishes written notice, including the identity of each new individual or company, to HCFA at the time of a change, if a change occurs in any of the following:

(i) Persons with an ownership or control interest, as defined in §§ 420.201 and 455.101 of this chapter.

(ii) The officers, directors, agents, or managing employees.

(iii) The religious entity, corporation, association, or other company responsible for the management of the RNHCI.

(iv) The RNHCI's administrator or director of nonmedical nursing services.

§ 403.740 Condition of participation: Staffing.

The RNHCI must be staffed with qualified experienced personnel who are present in sufficient numbers to meet the needs of the patients.
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(a) Standard: Personnel qualifications. The RNHCI must ensure that staff who supervise or furnish services to patients are qualified to do so and that staff allowed to practice without direct supervision have specific training to furnish these services.

(b) Standard: Education, training, and performance evaluation. (1) The RNHCI must ensure that staff (including contractors and other individuals working under arrangement) have the necessary education and training concerning their duties so that they can furnish services competently. This education includes, but is not limited to, training related to the individual job description, performance expectations, applicable organizational policies and procedures, and safety responsibilities.

(2) Staff must demonstrate, in practice, the skills and techniques necessary to perform their duties and responsibilities.

(3) The RNHCI must evaluate the performance of staff and implement measures for improvement.

§ 403.742 Condition of participation: Physical environment.

A RNHCI must be designed, constructed, and maintained to ensure the safety of the patients, staff, and the public.

(a) Standard: Buildings. The physical plant and the overall environment must be maintained in a manner that ensures the safety and well-being of the patients. The RNHCI must have the following:

(i) Emergency power for emergency lights, for fire detection and alarm systems, and for fire extinguishing systems.

(ii) Procedures for the proper storage and disposal of trash.

(iii) Proper ventilation and temperature control and appropriate lighting levels to ensure a safe and secure environment.

(iv) A written disaster plan to address loss of power, water, sewage, and other emergencies.

(v) Facilities for emergency gas and water supply.

(vi) An effective pest control program.

(vii) A preventive maintenance program to maintain essential mechanical, electrical, and fire protection equipment operating in an efficient and safe manner.

(viii) A working call system for patients to summon aid or assistance.

(b) Standard: Patient rooms. Patient rooms must be designed and equipped for adequate care, comfort, and privacy of the patient.

(i) Patient rooms must meet the following conditions:

(1) Accommodate no more than four patients.

(2) Measure at least 80 square feet per patient in multiple patient rooms and at least 100 square feet in single patient rooms.

(3) Have direct access to an exit corridor.

(iv) Be designed or equipped to assure full visual privacy for each patient.

(v) Have at least one window to the outside.

(vi) Have a floor at or above grade level.

(2) The RNHCI must furnish each patient with the following:

(i) A separate bed of proper size and height for the convenience of the patient.

(ii) A clean, comfortable mattress.

(iii) Bedding appropriate to the weather and climate.

(iv) Functional furniture appropriate to the patient's needs and individual closet space with clothes racks and shelves accessible to the patient.

(3) HCFA may permit variances in requirements specified in paragraphs (b)(1)(i) and (ii) of this section relating to rooms on an individual basis when the RNHCI adequately demonstrates in writing that the variances meet the following:

(i) Are in accordance with the special needs of the patients.

(ii) Will not adversely affect patients' health and safety.

§ 403.744 Condition of participation: Life safety from fire.

(a) General. An RNHCI must meet the following conditions:

(1) Except as provided in paragraph (b) of this section, the RNHCI must meet the new or existing health care occupancies provisions of the 1997 edition of the Life Safety Code of the National Fire Protection Association (NFPA 101), which is incorporated by
reference. Incorporation by reference of NFPA 101, the Life Safety Code, 1997 edition, was approved by the Director of the Federal Register in accordance with 5 U.S.C. 552(a) and 1 CFR part 51.\footnote{1} (See §403.70).

(2) The RNHCI must have written fire control plans that contain provisions for prompt reporting of fires; extinguishing fires; protection of patients, staff, and the public; evacuation; and cooperation with fire fighting authorities.

(3) The RNHCI must maintain written evidence of regular inspection and approval by State or local fire control agencies.

(b) Exceptions. (1) If application of the Life Safety Code required under paragraph (a)(1) of this section would result in unreasonable hardship upon the RNHCI, HCFA may waive specific provisions of the Life Safety Code, but only if the waiver does not adversely affect the health and safety of patients.

(2) If HCFA finds that the fire and safety code imposed by State law adequately protects patients in the institution, the provisions of the Life Safety Code required in paragraph (a)(1) of this section do not apply in that State.

§ 403.746 Condition of participation: Utilization review.

The RNHCI must have in effect a written utilization review plan to assess the necessity of services furnished. The plan must provide that records be maintained of all meetings, decisions, and actions by the utilization review committee.

(a) Standard: Utilization review plan. The utilization review plan must contain written procedures for evaluating the following:

(1) Admissions.

(2) Duration of care.

(3) Continuing care of an extended duration.

(4) Items and services furnished.

(b) Standard: Utilization review committee. The committee is responsible for evaluating each admission and ensuring that the admission is necessary and appropriate. The utilization review plan must be carried out by the utilization review committee, consisting of the governing body, administrator or other individual responsible for the overall administration of the RNHCI, the supervisor of nursing staff, and other staff as appropriate.

§ 403.750 Estimate of expenditures and adjustments.

(a) Estimates. HCFA estimates the level of expenditures for services provided under this subpart before the start of each FFY beginning with FFY 2000.

(b) Adjustments to payments. When the level of estimated expenditures is projected to exceed the FFY trigger level as described in paragraph (d) of this section, for the year of the projection, payments to RNHCIs will be reduced by a proportional percentage to prevent estimated expenditures from exceeding the trigger level. In addition to reducing payments proportionally, HCFA may impose alternative adjustments.

(c) Notification of adjustments. HCFA notifies participating RNHCIs before the start of the FFY of the type and level of expenditure reductions to be made and when these adjustments will apply.

(d) Calculation of trigger level. The trigger level for FFY 1998 is $20,000,000. For subsequent FFYs, the trigger level is the unadjusted trigger level increased or decreased by the carry forward as described in §403.754(b). The unadjusted trigger level is the base year amount (the unadjusted trigger level dollar amount for the prior FFY) increased by the average consumer price index (the single numerical value published monthly by the Bureau of Labor Statistics that presents the relationship in United States urban areas for the current cost of goods and services compared to a base year, to represent the change in spending power) for the 12-month period ending on July 31 preceding the beginning of the FFY.
§ 403.752 Payment provisions.

(a) Payment to RNHCIs. Payment for services may be made to an RNHCI that meets the conditions for coverage described in §403.720 and the conditions of participation described in §§403.730 through 403.746. Payment is made in accordance with §413.40 of this chapter to an RNHCI meeting these conditions. 

(b) Review of estimates and adjustments. There is no administrative or judicial review of the level of estimated expenditures or the adjustments in payments described in §§403.750(a) and (b).

(c) Effect on beneficiary liability. When payments are reduced in accordance with §403.750(b), the RNHCI may bill the beneficiary the amount of the Medicare reduction attributable to his or her covered services.

(d) Notification of beneficiary liability. (1) The RNHCI must notify the beneficiary in writing at the time of admission of any proposed or current proportional Medicare adjustment. A beneficiary currently receiving care in the RNHCI must be notified in writing at least 30 days before the Medicare reduction is to take effect. The notification must inform the beneficiary that the RNHCI can bill him or her for the proportional Medicare adjustment.

(2) The RNHCI must, at time of billing, provide the beneficiary with his or her liability for payment, based on a calculation of the Medicare reduction pertaining to the beneficiary's covered services permitted by §403.750(b).

§ 403.754 Monitoring expenditure level.

(a) Tracking expenditures. Starting in FFY 1999 HCFA begins monitoring Medicare payments to RNHCIs.

(b) Carry forward. The difference between the trigger level and Medicare expenditures for a FFY results in a carry forward that either increases or decreases the unadjusted trigger level described in §403.750(d). In no case may the carry forward exceed $50,000,000 for an FFY.

§ 403.756 Sunset provision.

(a) Effective date. Beginning with FFY 2002, if the level of estimated expenditures for all RNHCIs exceeds the trigger level for 3 consecutive FFYs, HCFA will not accept as the basis for payment of any claim any election executed on or after January 1 of the following calendar year.

(b) Notice of activation. A notice in the Federal Register will be published at least 60 days before January 1 of the calendar year that the sunset provision becomes effective.

(c) Effects of sunset provision. Only those beneficiaries who have a valid election in effect before January 1 of the year in which the sunset provision becomes effective will be able to claim Medicare payment for care in an RNHCI, and only for RNCHI services furnished during that election.
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Subpart B—Medical Services Coverage Decisions That Relate to Health Care Technology

AUTHORITY: Secs. 1102, 1862 and 1871 of the Social Security Act as amended (42 U.S.C. 1302, 1395y, and 1395hh).

SOURCE: 60 FR 48423, Sept. 19, 1995, unless otherwise noted.

§ 405.201 Scope of subpart and definitions.

(a) Scope. This subpart establishes that—
(1) HCFA uses the FDA categorization of a device as a factor in making Medicare coverage decisions; and
(2) HCFA may consider for Medicare coverage certain devices with an FDA-approved investigational device exemption (IDE) that have been categorized as non-experimental/investigational (Category B).

(b) Definitions. As used in this subpart—
Class I refers to devices for which the general controls of the Food, Drug, and Cosmetic Act, such as adherence to good manufacturing practice regulations, are sufficient to provide a reasonable assurance of safety and effectiveness.
Class II refers to devices that, in addition to general controls, require special controls, such as performance standards or postmarket surveillance, to provide a reasonable assurance of safety and effectiveness.
Class III refers to devices that cannot be classified into Class I or Class II because insufficient information exists to
§ 405.203 FDA categorization of investigational devices.

(a) The FDA assigns a device with an FDA-approved IDE to one of two categories:

(1) Experimental/Investigational (Category A) Devices.

(2) Non-Experimental/Investigational (Category B) Devices.

(b) The FDA notifies HCFA, when it notifies the sponsor, that the device is categorized by FDA as experimental/investigational (Category A) or non-experimental/investigational (Category B).

(c) HCFA uses the categorization of the device as a factor in making Medicare coverage decisions.

§ 405.205 Coverage of a non-experimental/investigational (Category B) device.

(a) For any device that meets the requirements of the exception at §411.15(o) of this chapter, the following procedures apply:

(1) The FDA notifies HCFA, when it notifies the sponsor, that the device is categorized by FDA as non-experimental/investigational (Category B).

(2) HCFA uses the categorization of the device as a factor in making Medicare coverage decisions.

(b) If the FDA becomes aware that a categorized device no longer meets the requirements of the exception at §411.15(o) of this chapter, the FDA notifies the sponsor and HCFA and the procedures described in paragraph (a)(2) of this section apply.

§ 405.207 Services related to a non-covered device.

(a) When payment is not made. Medicare payment is not made for medical and hospital services that are related to the use of a device that is not covered because HCFA determines the device is not "reasonable" and "necessary" under section 1862(a)(1)(A) of the Act or because it is excluded from coverage for other reasons. These services include all services furnished in preparation for the use of a noncovered device, and services furnished as necessary after-care that are incidental to recovery from the use of the device or from receiving related noncovered services.

(b) When payment is made. Medicare payment may be made for services, ordinarily covered by Medicare, to treat a condition or complication that arises...
§ 405.209 Payment for a non-experimental/investigational (Category B) device.

Payment under Medicare for a non-experimental/investigational (Category B) device is based on, and may not exceed, the amount that would have been paid for a currently used device serving the same medical purpose that has been approved or cleared for marketing by the FDA.

§ 405.211 Procedures for Medicare contractors in making coverage decisions for a non-experimental/investigational (Category B) device.

(a) General rule. In their review of claims for payment, Medicare contractors are bound by the statute, regulations, and all HCFA administrative issuances, including all national coverage decisions.

(b) Potentially covered non-experimental/investigational (Category B) devices. Medicare contractors may approve coverage for any device with an FDA-approved IDE categorized as a non-experimental/investigational (Category B) device if all other coverage requirements are met.

(c) Other considerations. Medicare contractors must consider whether any restrictions concerning site of service, indications for use, or any other list of conditions for coverage have been placed on the device’s use.

§ 405.213 Re-evaluation of a device categorization.

(a) General rules. (1) Any sponsor that does not agree with an FDA decision that categorizes its device as experimental/investigational (Category A) may request re-evaluation of the categorization decision.

(2) A sponsor may request review by HCFA only after the requirements of paragraph (b) of this section are met.

(3) No reviews other than those described in paragraphs (b) and (c) of this section are available to the sponsor.

(b) Request to FDA. A sponsor that does not agree with the FDA’s categorization of its device may submit a written request to the FDA at any time requesting re-evaluation of its original categorization decision, together with any information and rationale that it believes support recategorization. The FDA notifies both HCFA and the sponsor of its decision.

(c) Request to HCFA. If the FDA does not agree to recategorize the device, the sponsor may seek review from HCFA. A device sponsor must submit its request in writing to HCFA. HCFA obtains copies of relevant portions of the application, the original categorization decision, and supplementary materials. HCFA reviews all material submitted by the sponsor and the FDA’s recommendation. HCFA reviews only information in the FDA record to determine whether to change the categorization of the device. HCFA issues a written decision and notifies the sponsor of the IDE and the FDA.

§ 405.215 Confidential commercial and trade secret information.

To the extent that HCFA relies on confidential commercial or trade secret information in any judicial proceeding, HCFA will maintain confidentiality of the information in accordance with Federal law.

Subpart C—Suspension of Payment, Recovery of Overpayments, and Repayment of Scholarships and Loans

Authority: Secs. 1102, 1815, 1833, 1942, 1866, 1870, 1871, 1879, and 1892 of the Social Security Act (42 U.S.C. 1302, 1395g, 1395l, 1395u, 1395cc, 1395gg, 1395hh, 1395pp, and 1395ccc) and 31 U.S.C. 3711.

§ 405.301 Scope of subpart.

This subpart sets forth the policies and procedures for handling of incorrect payments and recovery of overpayments.

[54 FR 41733, Oct. 11, 1989]

§ 405.350 Individual's liability for payments made to providers and other persons for items and services furnished the individual.

Any payment made under title XVIII of the Act to any provider of services or other person with respect to any item or service furnished an individual shall be regarded as a payment to the individual, and adjustment shall be made pursuant to §§405.352 through 405.358 where:

(a) More than the correct amount is paid to a provider of services or other person and the Secretary determines that:

(1) Within a reasonable period of time, the excess over the correct amount cannot be recouped from the provider of services or other person, or

(2) The provider of services or other person was without fault with respect to the payment of such excess over the correct amount, or

(b) A payment has been made under the provisions described in section 1814(e) of the Act, to a provider of services for items and services furnished the individual.

(c) For purposes of paragraph (a)(2) of this section, a provider of services or other person shall, in the absence of evidence to the contrary, be deemed to be without fault if the determination of the carrier, the intermediary, or the Health Care Financing Administration that more than the correct amount was paid was made subsequent to the third year following the year in which notice was sent to such individual that such amount had been paid.


§ 405.351 Incorrect payments for which the individual is not liable.

Where an incorrect payment has been made to a provider of services or other person, the individual is liable only to the extent that he has benefited from such payment.

§ 405.352 Adjustment of title XVIII incorrect payments.

Where an individual is liable for an incorrect payment (i.e., a payment made under §405.350(a) or §405.350(b)) adjustment is made (to the extent of such liability) by:

(a) Decreasing any payment under title II of the Act, or under the Railroad Retirement Act of 1937, to which the individual is entitled; or

(b) In the event of the individual's death before adjustment is completed, by decreasing any payment under title II of the Act, or under the Railroad Retirement Act of 1937 payable to the estate of the individual or to any other person, that are based on the individual's earnings record (or compensation).


§ 405.353 Certification of amount that will be adjusted against individual title II or railroad retirement benefits.

As soon as practicable after any adjustment is determined to be necessary, the Secretary, for purposes of this subpart, shall certify the amount of the overpayment or payment (see §405.350) with respect to which the adjustment is to be made. If the adjustment is to be made by decreasing subsequent payments under the Railroad Retirement Act of 1937 payable to the estate of the individual or to any other person, that are based on the individual's earnings record (or compensation).

§ 405.355 Waiver of adjustment or recovery.

(a) The provisions of § 405.352 may not be applied and there may be no adjustment or recovery of an incorrect payment (i.e., a payment made under § 405.350(a) or § 405.350(b)) in any case where such incorrect payment has been made with respect to an individual who is without fault, or where such adjustment or recovery would be made by decreasing payments to which another person who is without fault is entitled as provided in section 1870(b) of the Act where such adjustment or recovery would defeat the purpose of title II or title XVIII of the Act or would be against equity and good conscience.

(b) Adjustment or recovery of an incorrect payment (or only such part of an incorrect payment as may be determined to be inconsistent with the purposes of Title XVIII of the Act) against an individual who is without fault shall be deemed to be against equity and good conscience if the determination that such payment was incorrect was made subsequent to the third year following the year in which notice of such payment was sent to such individual. (See 20 CFR 404.509 and 404.512.)

§ 405.356 Principles applied in waiver of adjustment or recovery.

The principles applied in determining waiver of adjustment or recovery (§ 405.355) are the applicable principles of § 405.358 and 20 CFR 404.507-404.509, 404.510a, and 404.512.

§ 405.357 Notice of right to waiver consideration.

Whenever an initial determination is made that more than the correct amount of payment has been made, notice of the provisions of section 1870(c) of the Act regarding waiver of adjustment or recovery shall be sent to the overpaid individual and to any other individual against whom adjustment or recovery of the overpayment is to be effected (see § 405.358).

§ 405.358 When waiver of adjustment or recovery may be applied.

Section 1870(c) of the Act provides that there shall be no adjustment or recovery in any case where an incorrect payment under title XVIII (hospital and supplementary medical insurance benefits) has been made (including a payment under section 1814(e) of the Act with respect to an individual:

(a) Who is without fault, and

(b) Adjustment or recovery would either:

(1) Defeat the purposes of title II or title XVIII of the Act, or

(2) Be against equity and good conscience.

§ 405.359 Liability of certifying or disbursing officer.

No certifying or disbursing officer shall be held liable for any amount certified or paid by him to any provider of services or other person:

(a) Where the adjustment or recovery of such amount is waived (see § 405.355), or

(b) Where adjustment (see § 405.352) or recovery is not completed prior to the death of all persons against whose benefits such adjustment is authorized.

§ 405.370 Definitions.

For purposes of this subpart, the following definitions apply:

Offset. The recovery by Medicare of a non-Medicare debt by reducing present or future Medicare payments and applying the amount withheld to the indebtedness. (Examples are Public Health Service debts or Medicaid debts recovered by HCFA).

Recoupment. The recovery by Medicare of any outstanding Medicare debt by reducing present or future Medicare payments and applying the amount withheld to the indebtedness.
or carrier from a provider or supplier of an approved Medicare payment amount before a determination of the amount of the overpayment exists.

[61 FR 63745, Dec. 2, 1996]

§ 405.371 Suspension, offset, and recoupment of Medicare payments to providers and suppliers of services.

(a) General. Medicare payments to providers and suppliers, as authorized under this subchapter (excluding payments to beneficiaries), may be—

(1) Suspended, in whole or in part, by HCFA, an intermediary, or a carrier if HCFA, the intermediary, or the carrier possesses reliable information that an overpayment or fraud or willful misrepresentation exists or that the payments to be made may not be correct, although additional evidence may be needed for a determination; or

(2) Offset or recouped, in whole or in part, by an intermediary or a carrier if the intermediary, carrier, or HCFA has determined that the provider or supplier to whom payments are to be made has been overpaid.

(b) Steps necessary for suspension of payment, offset, and recoupment. Except as provided in paragraph (c) of this section, HCFA, the intermediary, or carrier suspends payments only after it has complied with the procedural requirements set forth at § 405.372. The intermediary or carrier offsets or recoups payments only after it has complied with the procedural requirements set forth at § 405.373.

(c) Suspension of payment in the case of unfilled cost reports. If a provider has failed to timely file an acceptable cost report, payment to the provider is immediately suspended until a cost report is filed and determined by the intermediary to be acceptable. In the case of an unfilled cost report, the provisions of § 405.372 do not apply. (See § 405.372(a)(2) concerning failure to furnish other information.)

[61 FR 63746, Dec. 2, 1996]

§ 405.372 Proceeding for suspension of payment.

(a) Notice of intention to suspend—(1) General rule. Except as provided in paragraphs (a)(2) through (a)(4) of this section, if the intermediary, carrier, or HCFA has determined that a suspension of payments under § 405.371(a)(1) should be put into effect, the intermediary or carrier must notify the provider or supplier of the intention to suspend payments, in whole or in part, and the reasons for making the suspension.

(2) Failure to furnish information. The notice requirement of paragraph (a)(1) of this section does not apply if the intermediary or carrier suspends payments to a provider or supplier in accordance with section 1815(a) or section 1833(e) of the Act, respectively, because the provider or supplier has failed to submit information requested by the intermediary or carrier that is needed to determine the amounts due the provider or supplier. (See § 405.371(c) concerning failure to file timely acceptable cost reports.)

(3) Harm to Trust Funds. A suspension of payment may be imposed without prior notice if HCFA, the intermediary, or carrier determines that the Medicare Trust Funds would be harmed by giving prior notice. HCFA may base its determination on an intermediary’s or carrier’s belief that giving prior notice would hinder the possibility of recovering the money.

(4) Fraud or misrepresentation. If the intended suspension of payment involves suspected fraud or misrepresentation, HCFA determines whether to impose the suspension and if prior notice is appropriate. HCFA directs the intermediary or carrier as to the timing and content of the notification to the provider or supplier. HCFA is the real party in interest and is responsible for the decision. HCFA may base its decision on information from the intermediary, carrier, law enforcement agencies, or other sources. HCFA determines whether the information is reliable.

(b) Rebuttal—(1) If prior notice is required. If prior notice is required under paragraph (a) of this section, the intermediary or carrier must give the provider or supplier an opportunity for rebuttal in accordance with § 405.374. If a rebuttal statement is received within the specified time period, the suspension of payment goes into effect on the date stated in the notice, and the procedures and provisions set forth in
§ 405.375 apply. If by the end of the period specified in the notice no statement has been received, the suspension goes into effect automatically, and the procedures set forth in paragraph (c) of this section are followed.

(2) If prior notice is not required. If, under the provisions of paragraphs (a)(2) through (a)(4) of this section, a suspension of payment is put into effect without prior notice to the provider or supplier, the intermediary or carrier must, once the suspension is in effect, give the provider or supplier an opportunity to submit a rebuttal statement as to why the suspension should be removed.

(c) Subsequent action. If a suspension of payment is put into effect, the intermediary, carrier, or HCFA takes timely action after the suspension to obtain the additional evidence it may need to make a determination as to whether an overpayment exists or the payments may be made. The intermediary, carrier, or HCFA makes all reasonable efforts to expedite the determination. As soon as the determination is made, the intermediary or carrier informs the provider or supplier and, if appropriate, the suspension is rescinded or any existing recoupment or offset is adjusted to take into account the determination.

(d) Duration of suspension of payment—(1) General rule. Except as provided in paragraphs (d)(2) and (d)(3) of this section, a suspension of payment is limited to 180 days, starting with the date the suspension begins.

(2) 180-day extension. (i) An intermediary, a carrier, or, in cases of fraud and misrepresentation, OIG or a law enforcement agency, may request a one-time only extension of the suspension period for up to 180 additional days if it is unable to complete its examination of the information or investigation, as appropriate, within the 180-day time limit. The request must be submitted in writing to HCFA.

(ii) Upon receipt of a request for an extension, HCFA notifies the provider or supplier of the requested extension. HCFA then either extends the suspension of payment for up to an additional 180 days or determines that the suspended payments are to be released to the provider or supplier.

(3) Exceptions to the time limits. (i) The time limits specified in paragraphs (d)(1) and (d)(2) of this section do not apply if the case has been referred to, and is being considered by, the OIG for administrative action (for example, civil money penalties).

(ii) HCFA may grant an extension in addition to the extension provided under paragraph (d)(2) of this section if the Department of Justice submits a written request to HCFA that the suspension of payment be continued based on the ongoing investigation and anticipated filing of criminal and/or civil actions. At a minimum, the request must include the following:

(A) Identification of the entity under suspension.

(B) The amount of time needed for continued suspension in order to implement the criminal and/or civil proceedings.

(C) A statement of why and/or how criminal and/or civil actions may be affected if the requested extension is not granted.

(e) Disposition of suspended payments. Payments suspended under the authority of § 405.371(b) are first applied to reduce or eliminate any overpayments determined by the intermediary, carrier, or HCFA, including any interest assessed under the provisions of § 405.378, and then applied to reduce any other obligation to HCFA or to HHS. In the absence of a legal requirement that the excess be paid to another entity, the excess is released to the provider or supplier.

[61 FR 63746, Dec. 2, 1996]
§ 405.374 Opportunity for rebuttal.

(a) General rule. If prior notice of the suspension of payment, offset, or recoupment is given under §405.372 or §405.373, the intermediary or carrier must give the provider or supplier an opportunity, before the suspension, offset, or recoupment takes effect, to submit any statement (to include any pertinent information) as to why it should not be put into effect on the date specified in the notice. Except as provided in paragraph (b) of this section, the provider or supplier has at least 15 days following the date of notification to submit the statement.

(b) Exception. The intermediary or carrier may for cause—

(1) Impose a shorter period for rebuttal; or

(2) Extend the time within which the statement must be submitted.

[61 FR 63747, Dec. 2, 1996]

§ 405.375 Time limits for, and notification of, administrative determination after receipt of rebuttal statement.

(a) Submission and disposition of evidence. If the provider or supplier submits a statement, under §405.374, as to why a suspension of payment, offset, or recoupment should not be put into effect, or, under §405.372(b)(2), why a suspension should be terminated, HCFA, the intermediary, or carrier must, within 15 days, from the date the statement is received, consider the statement (including any pertinent evidence submitted), together with any other material bearing upon the case, and determine whether the facts justify the suspension, offset, or recoupment or, if already initiated, justify the termination of the suspension, offset, or recoupment. Suspension, offset, or recoupment is not delayed beyond the date stated in the notice in order to review the statement.

(b) Notification of determination. The intermediary or carrier must send written notice of the determination made under paragraph (a) of this section to the provider or supplier. The notice must—

(1) In the case of offset or recoupment, contain rationale for the determination; and

(2) In the case of suspension of payment, contain specific findings on the conditions upon which the suspension is initiated, continued, or removed and an explanatory statement of the determination.

[61 FR 63747, Dec. 2, 1996]

§ 405.376 Suspension and termination of collection action and compromise of claims for overpayment.

(a) Basis and purpose. This section contains requirements and procedures for the compromise of, or suspension or termination of collection action on, claims for overpayments against a provider or a supplier under the Medicare
program. It is adopted under the authority of the Federal Claims Collection Act (31 U.S.C. 3711). Collection and compromise of claims against Medicare beneficiaries are explained at 20 CFR 404.515.

(b) Definitions. As used in this section, debtor means a provider of services or a physician or other supplier of services that has been overpaid under title XVIII of the Social Security Act. It includes an individual, partnership, corporation, estate, trust, or other legal entity.

(c) Basic conditions. A claim for recovery of Medicare overpayments against a debtor may be compromised, or collection action on it may be suspended or terminated, by the Health Care Financing Administration (HCFA) if:

1. The claim does not exceed $100,000, or such higher amount as the Attorney General may from time to time prescribe, exclusive of interest; and
2. There is no indication of fraud, the filing of a false claim, or misrepresentation on the part of the debtor or any director, partner, manager, or other party having an interest in the claim.

(d) Basis for compromise. A claim may be compromised for one or more of the following reasons:

1. The debtor, or the estate of a deceased debtor, does not have the present or prospective ability to pay the full amount within a reasonable time;
2. The debtor refuses to pay the claim in full and the United States is unable to collect the full amount within a reasonable time by legal proceedings;
3. There is real doubt the United States can prove its case in court; or
4. The cost of collecting the claim does not justify enforced collection of the full amount.

(e) Basis for termination of collection action. Collection action may be terminated for one or more of the following reasons:

1. The United States cannot enforce collection of any significant sum;
2. The debtor cannot be located, there is no security to be liquidated, the statute of limitations has run, and the prospects of collecting by offset are too remote to justify retention of the claim;
3. The cost of further collection action is likely to exceed any recovery;
4. It is determined the claim is without merit; or
5. Evidence to substantiate the claim is no longer available.

(f) Basis for suspension of collection action. Collection action may be suspended for either of the following reasons if future collection action is justified based on potential productivity, including foreseeable ability to pay, and size of claim:

1. The debtor cannot be located;
2. The debtor is unable to make payments on the claim or to fulfill an acceptable compromise.

(g) Factors considered. In determining whether a claim will be compromised, or collection action terminated or suspended, HCFA will consider the following factors:

1. Age and health of the debtor, present and potential income, inheritance prospects, possible concealment or fraudulent transfer of assets, and the availability of assets which may be reached by enforced collection proceedings, for compromise under paragraph (d)(1) of this section, termination under paragraph (e)(1) of this section, and suspension under paragraph (f)(2) of this section;
2. Applicable exemptions available to a debtor and uncertainty concerning the price of the property in a forced sale, for compromise under paragraph (d)(2) of this section and termination under paragraph (e)(1) of this section; and
3. The probability of proving the claim in court, the probability of full or partial recovery, the availability of necessary evidence, and related pragmatic considerations, for compromise under paragraph (d)(3) of this section.

(h) Amount of compromise. The amount accepted in compromise will be reasonable in relation to the amount that can be recovered by enforced collection proceedings. Consideration shall be given to the following:

1. The exemptions available to the debtor under State or Federal law;
2. The time necessary to collect the overpayment;
§ 405.377 Withholding Medicare payments to recover Medicaid overpayments.

(a) Basis and purpose. This section implements section 4185 of the Act, which provides for withholding Medicare payments to certain Medicaid providers that have not arranged to repay Medicaid overpayments as determined by the Medicaid State agency or have failed to provide information necessary to determine the amount (if any) of overpayments.

(b) When withholding may be used. HCFA may withhold Medicare payment to offset Medicaid overpayments that a Medicaid agency has been unable to collect if—

(1) The Medicaid agency has followed the procedure specified in §447.31 of this chapter; and

(2) The institution or person is one described in paragraph (c) of this section and either—

(i) Has not made arrangements satisfactory to the Medicaid agency to repay the overpayment; or

(ii) Has not provided information to the Medicaid agency necessary to enable the agency to determine the existence or amount of Medicaid overpayment.

(c) Institutions or persons affected. Withholding under paragraph (b) of this section may be made with respect to any of the following entities that has or had in effect an agreement with a Medicaid agency to furnish services under an approved Medicaid State plan:

(1) An institutional provider that has in effect an agreement under section 1866 of the Act. (Part 489 (Provider and Supplier Agreements) implements section 1866 of the Act.)

(2) A physician or supplier that has accepted payment on the basis of an assignment under section 1842(b)(3)(B)(ii) of the Act. (Section 424.55 sets forth the conditions a supplier agrees to in accepting assignment.)

(d) Amount to be withheld. HCFA contacts the appropriate intermediary or carrier to determine the amount of Medicare payment to which the institution or person is entitled.

(1) HCFA contacts the appropriate intermediary or carrier to determine the amount of Medicare payment to which the institution or person is entitled.

(2) HCFA may require the intermediary or carrier to withhold Medicare payments to the institution or person by the lesser of the following amounts:

(i) The amount of the Medicare payments to which the institution or person would otherwise be entitled.

(ii) The total Medicaid overpayment to the institution or person.

(e) Notice of withholding. If HCFA intends to withhold payments under this section, it notifies by certified mail, return receipt requested, the institution or person and the appropriate intermediary or carrier of the intention to withhold Medicare payments and follows the procedure in §405.374. The notice includes—

(1) Identification of the institution or person; and

(2) The amount of Medicare overpayment to be withheld from payments to which the institution or person would otherwise be entitled under Medicare.
§ 405.378 Termination of withholding.

(f) Termination of withholding. HCFA terminates the withholding if—

(1) The Medicaid overpayment is completely recovered;

(2) The institution or person enters into an agreement satisfactory to the Medicaid agency to repay the overpayment; or

(3) The Medicaid agency determines that there is no overpayment based on newly acquired evidence or a subsequent audit.

(g) Disposition of funds withheld.

HCFA releases amounts withheld under this section to the Medicaid agency to be applied against the Medicaid overpayment made by the State agency.

[61 FR 63747, Dec. 2, 1996]

§ 405.378 Interest charges on overpayment and underpayments to providers, suppliers, and other entities.

(a) Basis and purpose. This section, which implements sections 1815(d) and 1833(j) of the common law and Act, and authority granted under the Federal Claims Collection Act, provides for the charging and payment of interest on overpayments and underpayments to Medicare providers, suppliers, HMOs, competitive medical plans (CMPs), and health care prepayment plans (HCPPs).

(b) Basic rules. (1) HCFA will charge interest on overpayments, and pay interest on underpayments, to providers and suppliers of services (including physicians and other practitioners), except as specified in paragraphs (f) and (h) of this section.

(2) Interest will accrue from the date of the final determination as defined in paragraph (c) of this section, and will either be charged on the overpayment balance or paid on the underpayment balance for each 30-day period that payment is delayed. (Periods of less than 30 days will be treated as a full 30-day period, and the 30-day interest charge will be applied to any balance.)

(c) Definition of final determination. (1) For purposes of this section, any of the following constitutes a final determination:

(i) A Notice of Amount of Program Reimbursement (NPR) is issued, as discussed in §§405.1803, 417.576, and 417.810, and either—

(A) A written demand for payment is made; or

(B) A written determination of an underpayment is made by the intermediary after a cost report is filed.

(ii) In cases in which an NPR is not used as a notice of determination (that is, primarily under part E of this chapter, interim cost settlements made for HMOs, CMPs, and HCPPs under §§417.574 and 417.810(e) of this chapter, and initial retroactive adjustment determinations under §413.64(f)(2) of this chapter. In the case of interim cost settlements and initial retroactive adjustment determinations, if the debtor does not dispute the adjustment determination within the timeframe designated in the notice of the determination (generally at least 15 days), a final determination is deemed to have been made. If the provider or supplier does dispute portions of the determination, a final determination is deemed to have been made on those portions when the intermediary issues a new determination in response to the dispute.

(iii) Other examples of cases in which an NPR is not used are carrier reasonable charge determinations under subpart E of this part, interim cost settlements made for HMOs, CMPs, and HCPPs under §§417.574 and 417.810(e) of this chapter, and initial retroactive adjustment determinations under §413.64(f)(2) of this chapter. In the case of interim cost settlements and initial retroactive adjustment determinations, if the debtor does not dispute the adjustment determination within the timeframe designated in the notice of the determination (generally at least 15 days), a final determination is deemed to have been made. If the provider or supplier does dispute portions of the determination, a final determination is deemed to have been made on those portions when the intermediary issues a new determination in response to the dispute.

(iv) The due date of a timely-filed cost report that indicates an amount is due HCFA, and is not accompanied by payment in full. (If an additional overpayment or underpayment is determined by the carrier or intermediary, a final determination on the additional amount is made in accordance with paragraphs (c)(1)(i), (c)(1)(ii), or (c)(1)(iii), of this section.)

(v) With respect to a cost report that is not filed on time, the day following the date the cost report was due (plus a single extension of time not to exceed 30 days if granted for good cause), until the time as a cost report is filed. (When the cost report is subsequently filed, there is an additional determination as
specified in paragraphs (c)(1)(i), (ii), (iii), or (iv) of this section.)

(2) Except as required by any subsequent administrative or judicial reversal, interest accrues from the date of final determination as specified in this subsection.

(d) Rate of interest. (1) The interest rate on overpayments and underpayments is the higher of—

(i) The rate as fixed by the Secretary of the Treasury after taking into consideration private consumer rates of interest prevailing on the date of final determination as defined in paragraph (c) of this section (this rate is published quarterly in the Federal Register by the Department under 45 CFR 30.13(a)); or

(ii) The current value of funds rate (this rate is published annually in the Federal Register by the Secretary of the Treasury, subject to quarterly revisions).

(2) [Reserved]

(e) Accrual of interest. (1) If a cost report is filed that does not indicate an amount is due HCFA but the intermediary makes a final determination that an overpayment exists, or if a carrier makes a final determination that an overpayment to a physician or supplier exists, interest will accrue beginning with the date of such final determination. Interest will continue to accrue during periods of administrative and judicial appeal and until final disposition of the claim.

(2)(i) If a cost report is filed and indicates that an amount is due HCFA, interest on the amount due will accrue from the due date of the cost report unless—

(A) Full payment on the amount due accompanies the cost report; or

(B) The provider and the intermediary agree in advance to liquidate the overpayment through a reduction in interim payments over the next 30-day period.

(ii) If the intermediary determines an additional overpayment during the cost settlement process, interest will accrue from the date of each determination.

(iii) The interest rate on each of the final determinations of an overpayment will be the rate of interest in effect on the date the determination is made.

(3) In the case of a cost report that is not filed on time, interest also will accrue on a determined overpayment from the day following the due date of the report (plus a single extension of time not to exceed 30 days if granted for good cause, as specified in §413.24(f)) of this chapter, to the time the cost report is filed.

(4) If an intermediary or a carrier makes a final determination that an underpayment exists, interest to the provider or the supplier will accrue from the date of notification of the underpayment.

(f) Waiver of interest charges. (1) When an intermediary or a carrier makes a final determination that an overpayment or underpayment exists, as specified in paragraphs (e)(1), (e)(2)(ii), and (e)(4)—

(i) Interest charges will be waived if the overpayment or underpayment is completely liquidated within 30 days from the date of the final determination.

(ii) HCFA may waive interest charges if it determines that the administrative cost of collecting them exceeds the interest charges.

(2) Interest will not be waived for that period of time during which the cost report was due but remained unfiled for more than 30 days, as specified in paragraph (e)(3) of this section.

(g) Rules applicable to partial payments. If an overpayment is repaid in installments or recouped by withholding from several payments due the provider or supplier of services—

(1) Each payment or recoupment will be applied first to accrued interest and then to the principal; and

(2) After each payment or recoupment, interest will accrue on the remaining unpaid balance.

(h) Exceptions to applicability. (1) The provisions of this section do not apply to the time period for which interest is payable under §413.64(j) of this chapter because the provider seeks judicial review of a decision of the Provider Reimbursement Review Board, or a subsequent reversal, affirmance, or modification of that decision by the Administrator. Prior to that time, until the provider seeks judicial review, interest
accrues at the rate specified in this section on outstanding unpaid balances resulting from final determinations as defined in paragraph (c) of this section.

(2) If an overpayment or an underpayment determination is reversed administratively or judicially, and the reversal is no longer subject to appeal, appropriate adjustments will be made with respect to the overpayment or underpayment and the amount of interest charged.

(i) Nonallowable cost. As specified in §§412.113 and 413.153 of this chapter, interest accrued on overpayments and interest on funds borrowed specifically to repay overpayments are not considered allowable costs, up to the amount of the overpayment, unless the provider had made a prior commitment to borrow funds for other purposes (for example, capital improvements). (See §413.153(a) of this chapter for exceptions based on administrative or judicial reversal.)


REPAYMENT OF SCHOLARSHIPS AND LOANS

§ 405.380 Collection of past-due amounts on scholarship and loan programs.

(a) Basis and purpose. This section implements section 1892 of the Act, which authorizes the Secretary to deduct from Medicare payments for services amounts considered as past-due obligations under the National Health Service Corps Scholarship program, the Physician Shortage Area Scholarship program, and the Health Education Assistance Loan program.

(b) Offsetting against Medicare payment. (1) Medicare carriers and intermediaries offset against Medicare payments in accordance with the signed repayment agreement between the Public Health Service and individuals who have breached their scholarship or loan obligations and who—

(i) Accept Medicare assignment for services;

(ii) Are employed by or affiliated with a provider, HMO, or Competitive Medical Plan (CMP) that receives Medicare payment for services; or

(iii) Are members of a group practice that receives Medicare payment for services.

(2) For purposes of this section, ‘‘provider’’ includes all entities eligible to receive Medicare payment in accordance with an agreement under section 1866 of the Act.

(c) Beginning of offset. (1) The Medicare carrier offsets Medicare payments beginning six months after it notifies the individual or the group practice of the amount to be deducted and the particular individual to whom the deductions are attributable.

(2) The Medicare intermediary offsets payments beginning six months after it notifies the provider, HMO, CMP or group practice of the amount to be deducted and the particular individuals to whom the deductions are attributable. Offset of payments is made in accordance with the terms of the repayment agreement. If the individual ceases to be employed by the provider, HMO, or CMP, or leaves the group practice, no deduction is made.

(d) Refusal to offset against Medicare payment. If the individual refuses to enter into a repayment agreement, or breaches any provision of the agreement, or if Medicare payment is insufficient to maintain the offset collection according to the agreed upon formula, then—

(1) The Department, within 30 days if feasible, informs the Attorney General; and

(2) The Department excludes the individual from Medicare until the entire past due obligation has been repaid, unless the individual is a sole community practitioner or the sole source of essential specialized services in a community and the State requests that the individual not be excluded.

[57 FR 19092, May 4, 1992]

Subpart D—Private Contracts

AUTHORITY: Secs. 1102, 1802, and 1871 of the Social Security Act (42 U.S.C. 1302, 1395a, and 1395hh).

SOURCE: 63 FR 58901, Nov. 2, 1998, unless otherwise noted.
§ 405.400 Definitions.

For purposes of this subpart, the following definitions apply:

Beneficiary means an individual who is enrolled in Part B of Medicare.

Emergency care services means services furnished to an individual for treatment of an "emergency medical condition" as that term is defined in §422.2 of this chapter.

Legal representative means one or more individuals who, as determined by applicable State law, has the legal authority to enter into the contract with the physician or practitioner on behalf of the beneficiary.

Opt-out means the status of meeting the conditions specified in §405.410.

Opt-out period means the 2-year period beginning on the effective date of the affidavit as specified by §405.410(c)(1) or §405.410(c)(2), as applicable.

Participating physician means a "physician" as defined in this section who has signed an agreement to participate in Part B of Medicare.

Physician means a doctor of medicine or a doctor of osteopathy who is currently licensed as that type of doctor in each State in which he or she furnishes services to patients.

Practitioner means a physician assistant, nurse practitioner, clinical nurse specialist, certified registered nurse anesthetist, certified nurse midwife, clinical psychologist, or clinical social worker, who is currently legally authorized to practice in that capacity by each State in which he or she furnishes services to patients or clients.

Private contract means a document that meets the criteria specified in §405.415.

Properly opt-out means to complete, without defect, the requirements for opt-out as specified in §405.410.

Properly terminate opt-out means to complete, without defect, the requirements for terminating opt-out as specified in §405.445.

Urgent care services means services furnished to an individual who requires services to be furnished within 12 hours in order to avoid the likely onset of an emergency medical condition.

§ 405.405 General rules.

(a) A physician or practitioner may enter into one or more private contracts with Medicare beneficiaries for the purpose of furnishing items or services that would otherwise be covered by Medicare, provided the conditions of this subpart are met.

(b) A physician or practitioner who enters into at least one private contract with a Medicare beneficiary under the conditions of this subpart, and who submits one or more affidavits in accordance with this subpart, opts-out of Medicare for a 2-year period unless the opt-out is terminated early according to §405.445. The physician's or practitioner's opt-out may be renewed for subsequent 2-year periods.

(c) Both the private contracts described in paragraph (a) of this section and the physician's or practitioner's opt-out described in paragraph (b) of this section are null and void if the physician or practitioner fails to properly opt-out in accordance with the conditions of this subpart.

(d) Both the private contracts described in paragraph (a) of this section and the physician's or practitioner's opt-out described in paragraph (b) of this section are null and void for the remainder of the opt-out period if the physician or practitioner fails to remain in compliance with the conditions of this subpart during the opt-out period.

(e) Services furnished under private contracts meeting the requirements of this subpart are not covered services under Medicare, and no Medicare payment will be made for such services either directly or indirectly, except as permitted in accordance with §405.435(c).

§ 405.410 Conditions for properly opting-out of Medicare.

The following conditions must be met for a physician or practitioner to properly opt-out of Medicare:

(a) Each private contract between a physician or a practitioner and a Medicare beneficiary that is entered into prior to the submission of the affidavit described in paragraph (b) of this section must meet the specifications of §405.415.
(b) The physician or practitioner must submit an affidavit that meets the specifications of §405.420 to each Medicare carrier with which he or she would file claims absent completion of opt-out.

(c) A nonparticipating physician or a practitioner may opt-out of Medicare at any time in accordance with the following:

(1) The 2-year opt-out period begins the date the affidavit meeting the requirements of §405.420 is signed, provided the affidavit is filed within 10 days after he or she signs his or her first private contract with a Medicare beneficiary.

(2) If the physician or practitioner does not timely file any required affidavit, the 2-year opt-out period begins when the last such affidavit is filed. Any private contract entered into before the last required affidavit is filed becomes effective upon the filing of the last required affidavit and the furnishing of any items or services to a Medicare beneficiary under such contract before the last required affidavit is filed is subject to standard Medicare rules.

(d) A participating physician may properly opt-out of Medicare at the beginning of any calendar quarter, provided that the affidavit described in §405.420 is submitted to the participating physician's Medicare carriers at least 30 days before the beginning of the selected calendar quarter. A private contract entered into before the beginning of the selected calendar quarter becomes effective at the beginning of the selected calendar quarter and the furnishing of any items or services to a Medicare beneficiary under such contract before the beginning of the selected calendar quarter is subject to standard Medicare rules.

§405.415 Requirements of the private contract.

A private contract under this subpart must:

(a) Be in writing and in print sufficiently large to ensure that the beneficiary is able to read the contract.

(b) Clearly state whether the physician or practitioner is excluded from Medicare under sections 1128, 1156, or 1892 or any other section of the Social Security Act.

(c) State that the beneficiary or his or her legal representative accepts full responsibility for payment of the physician's or practitioner's charge for all services furnished by the physician or practitioner.

(d) State that the beneficiary or his or her legal representative understands that Medicare limits do not apply to what the physician or practitioner may charge for items or services furnished by the physician or practitioner.

(e) State that the beneficiary or his or her legal representative agrees not to submit a claim to Medicare or to ask the physician or practitioner to submit a claim to Medicare.

(f) State that the beneficiary or his or her legal representative understands that Medicare payment will not be made for any items or services furnished by the physician or practitioner that would have otherwise been covered by Medicare if there was no private contract and a proper Medicare claim had been submitted.

(g) State that the beneficiary or his or her legal representative agrees not to submit a claim to Medicare or to ask the physician or practitioner to submit a claim to Medicare.

(h) State the expected or known effective date and expected or known expiration date of the opt-out period.

(i) State that the beneficiary or his or her legal representative understands that Medigap plans do not, and that other supplemental plans may elect not to, make payments for items and services not paid for by Medicare.

(j) Be signed by the beneficiary or his or her legal representative and by the physician or practitioner.

(k) Not be entered into by the beneficiary or by the beneficiary's legal representative during a time when the beneficiary requires emergency care services or urgent care services. (However, a physician or practitioner may...
§ 405.420

furnish emergency or urgent care services to a Medicare beneficiary in accordance with § 405.440.

(l) Be provided (a photocopy is permissible) to the beneficiary or to his or her legal representative before items or services are furnished to the beneficiary under the terms of the contract.

(m) Be retained (original signatures of both parties required) by the physician or practitioner for the duration of the opt-out period.

(n) Be made available to HCFA upon request.

(o) Be entered into for each opt-out period.

§ 405.420 Requirements of the opt-out affidavit.

An affidavit under this subpart must:

(a) Be in writing and be signed by the physician or practitioner.

(b) Contain the physician's or practitioner's full name, address, telephone number, national provider identifier (NPI) or billing number, if one has been assigned, uniform provider identification number (UPIN) if one has been assigned, or, if neither an NPI nor a UPIN has been assigned, the physician's or practitioner's tax identification number (TIN).

(c) State that, except for emergency or urgent care services (as specified in § 405.440), during the opt-out period the physician or practitioner will provide services to Medicare beneficiaries only through private contracts that meet the criteria of paragraph § 405.415 for services that, but for their provision under a private contract, would have been Medicare-covered services.

(d) State that the physician or practitioner will not submit a claim to Medicare for any service furnished to a Medicare beneficiary during the opt-out period, nor will the physician or practitioner permit any entity acting on his or her behalf to submit a claim to Medicare for services furnished to a Medicare beneficiary, except as specified in § 405.440.

(e) State that, during the opt-out period, the physician or practitioner understands that he or she may receive no direct or indirect Medicare payment for services that he or she furnishes to Medicare beneficiaries with whom he or she has privately contracted, whether as an individual, an employee of an organization, a partner in a partnership, under a reassignment of benefits, or as payment for a service furnished to a Medicare beneficiary under a Medicare+Choice plan.

(f) State that a physician or practitioner who opts-out of Medicare acknowledges that, during the opt-out period, his or her services are not covered under Medicare and that no Medicare payment may be made to any entity for his or her services, directly or on a capitated basis.

(g) State a promise by the physician or practitioner to the effect that, during the opt-out period, the physician or practitioner agrees to be bound by the terms of both the affidavit and the private contracts that he or she has entered into.

(h) Acknowledge that the physician or practitioner recognizes that the terms of the affidavit apply to all Medicare-covered items and services furnished to Medicare beneficiaries by the physician or practitioner during the opt-out period (except for emergency or urgent care services furnished to the beneficiaries with whom he or she has not previously privately contracted) without regard to any payment arrangements the physician or practitioner may make.

(i) With respect to a physician who has signed a Part B participation agreement, acknowledge that such agreement terminates on the effective date of the affidavit.

(j) Acknowledge that the physician or practitioner understands that a beneficiary who has not entered into a private contract and who requires emergency or urgent care services may not be asked to enter into a private contract with respect to receiving such services and that the rules of § 405.440 apply if the physician furnishes such services.

§ 405.425 Effects of opting-out of Medicare.

If a physician or practitioner opts-out of Medicare in accordance with this subpart for the 2-year period for which the opt-out is effective, the following results obtain:

(a) Except as provided in § 405.440, no payment may be made directly by
§ 405.435 Failure to maintain opt-out.

(a) A physician or practitioner fails to maintain opt-out if—
   (1) He or she is excluded under sections 1128, 1156, or 1892 of the Social Security Act and fails to maintain opt-out under any provision of this chapter.
   (2) He or she is reenrolled under sections 1128, 1156, or 1892 of the Social Security Act and fails to maintain opt-out under any provision of this chapter.
   (3) He or she is not excluded under sections 1128, 1156, or 1892 of the Social Security Act but fails to maintain opt-out under any provision of this chapter.

(b) The death of a beneficiary who has entered into a private contract (or whose legal representative has done so) does not invoke § 424.62 or § 424.64 of this chapter with respect to the physician or practitioner with whom the beneficiary (or legal representative) has privately contracted.

(c) The physician or practitioner who has not been excluded under sections 1128, 1156, or 1892 of the Social Security Act may order, prescribe, or certify the need for Medicare-covered items and services except as provided in § 1001.1901 of this title, and must otherwise comply with the terms of the exclusion in accordance with § 1001.1901 effective with the date of the exclusion.

§ 405.430 Failure to properly opt-out.

(a) A physician or practitioner fails to properly opt-out if—
   (1) Any private contract between the physician or practitioner and a Medicare beneficiary, that was entered into before the affidavit described in § 405.420 was filed, does not meet the specifications of § 405.415; or
   (2) He or she fails to submit the affidavit(s) in accordance with § 405.420.

(b) If a physician or practitioner fails to properly opt-out in accordance with paragraph (a) of this section, the following results obtain:
   (1) The physician's or practitioner's attempt to opt-out of Medicare is nullified, and all of the private contracts between the physician or practitioner and Medicare beneficiaries for the two-year period covered by the attempted opt-out are deemed null and void.
   (2) The physician or practitioner must submit claims to Medicare for all Medicare-covered items and services furnished to Medicare beneficiaries, including the items and services furnished under the nullified contracts. A nonparticipating physician is subject to the limiting charge provisions of § 414.48 of this chapter. A participating physician is subject to the limitations on charges of the participation agreement he or she signed.
   (3) The practitioner may not reassign any claim except as provided in § 424.60 of this chapter.
   (4) The practitioner may neither bill nor collect an amount from the beneficiary except for applicable deductible and coinsurance amounts.
   (5) The physician or practitioner may make another attempt to properly opt-out at any time.

§ 405.435 Failure to maintain opt-out.

(a) A physician or practitioner fails to maintain opt-out under this subpart if, during the opt-out period—
   (1) He or she knowingly and willfully—

Medicare or by any Medicare+Choice plan to the physician or practitioner or to any entity to which the physician or practitioner reassigns his right to receive payment for services.

(b) The physician or practitioner may not furnish any item or service that would otherwise be covered by Medicare (except for emergency or urgent care services) to any Medicare beneficiary except through a private contract that meets the requirements of this subpart.

(c) The physician or practitioner is not subject to the requirement to submit a claim for items or services furnished to a Medicare beneficiary, as specified in § 424.5(a)(6) of this chapter, except as provided in § 405.440.

(d) The physician or practitioner is prohibited from submitting a claim to Medicare for items or services furnished to a Medicare beneficiary except as provided in § 405.440.

(e) In the case of a physician, he or she is not subject to the limiting charge provisions of § 414.48 of this chapter, except for services provided under § 405.440.

(f) The physician or practitioner is not subject to the prohibition-on-reassignment provisions of § 414.80 of this chapter, except for services provided under § 405.440.

(g) In the case of a practitioner, he or she is not prohibited from billing or collecting amounts from beneficiaries as provided in 42 U.S.C. 1395u(b)(18)(B).

(h) The death of a beneficiary who has entered into a private contract (or whose legal representative has done so) does not invoke § 424.62 or § 424.64 of this chapter with respect to the physician or practitioner with whom the beneficiary (or legal representative) has privately contracted.

(i) The physician or practitioner who has not been excluded under sections 1128, 1156, or 1892 of the Social Security Act may order, prescribe, or certify the need for Medicare-covered items and services except as provided in § 1001.1901 of this title, and must otherwise comply with the terms of the exclusion in accordance with § 1001.1901 effective with the date of the exclusion.
§ 405.440

(i) Submits a claim for Medicare payment (except as provided in § 405.440); or

(ii) Receives Medicare payment directly or indirectly for Medicare-covered services furnished to a Medicare beneficiary (except as provided in § 405.440).

(2) He or she fails to enter into private contracts with Medicare beneficiaries for the purpose of furnishing items and services that would otherwise be covered by Medicare, or enters into contracts that fail to meet the specifications of § 405.435; or

(3) He or she fails to comply with the provisions of § 405.440 regarding billing for emergency care services or urgent care services; or

(4) He or she fails to retain a copy of each private contract that he or she has entered into for the duration of the opt-out period for which the contracts are applicable or fails to permit HCFA to inspect them upon request.

(b) If a physician or practitioner fails to maintain opt-out in accordance with paragraph (a) of this section, and fails to demonstrate, within 45 days of a notice from the carrier of a violation of paragraph (a) of this section, that he or she has taken good faith efforts to maintain opt-out (including by refunding amounts in excess of the charge limits to beneficiaries with whom he or she did not sign a private contract), the following results obtain, effective 46 days after the date of the notice, but only for the remainder of the opt-out period:

(1) All of the private contracts between the physician or practitioner and Medicare beneficiaries are deemed null and void.

(2) The physician's or practitioner's opt-out of Medicare is nullified.

(3) The physician or practitioner must submit claims to Medicare for all Medicare-covered items and services furnished to Medicare beneficiaries.

(4) The physician or practitioner or beneficiary will not receive Medicare payment on Medicare claims for the remainder of the opt-out period, except as provided in paragraph (c) of this section.

(5) The physician is subject to the limiting charge provisions of § 414.48 of this chapter.

(6) The practitioner may not reassign any claim except as provided in § 424.80 of this chapter.

(7) The practitioner may neither bill nor collect any amount from the beneficiary except for applicable deductible and coinsurance amounts.

(8) The physician or practitioner may not attempt to once more meet the criteria for properly opting-out until the 2-year opt-out period expires.

(c) Medicare payment may be made for the claims submitted by a beneficiary for the services of an opt-out physician or practitioner when the physician or practitioner did not privately contract with the beneficiary for services that were not emergency care services or urgent care services and that were furnished no later than 15 days after the date of a notice by the carrier that the physician or practitioner has opted-out of Medicare.

§ 405.440 Emergency and urgent care services.

(a) A physician or practitioner who has opted-out of Medicare under this subpart need not enter into a private contract to furnish emergency care services or urgent care services to a Medicare beneficiary. Accordingly, a physician or practitioner will not be determined to have failed to maintain opt-out if he or she furnishes emergency care services or urgent care services to a Medicare beneficiary with whom the physician or practitioner did not previously entered into a private contract, provided the physician or practitioner complies with the billing requirements specified in paragraph (b) of this section.

(b) When a physician or practitioner who has not been excluded under sections 1128, 1156, or 1892 of the Social Security Act furnishes emergency care services or urgent care services to a Medicare beneficiary with whom the physician or practitioner has not previously entered into a private contract, he or she:

(1) Must submit a claim to Medicare in accordance with both 42 CFR part 424 and Medicare instructions (including but not limited to complying with proper coding of emergency or urgent care services furnished by physicians
§ 405.450 Appeals.

(a) A determination by HCFA that a physician or practitioner has failed to properly opt-out, failed to maintain opt-out, failed to timely renew opt-out, failed to privately contract, or failed to properly terminate opt-out is an initial determination for purposes of §405.803.

(b) A determination by HCFA that no payment can be made to a beneficiary for the services of a physician who has opted-out is an initial determination for purposes of §405.803.

§ 405.455 Application to Medicare+Choice contracts.

An organization that has a contract with HCFA to provide one or more Medicare+Choice (M+C) plans to beneficiaries (part 422 of this chapter):

(a) Must acquire and maintain information from Medicare carriers on physicians and practitioners who have opted-out of Medicare.

(b) Must make no payment directly or indirectly for Medicare covered services furnished to a Medicare beneficiary by a physician or practitioner who has opted-out of Medicare.
§ 405.500

(c) May make payment to a physician or practitioner who furnishes emergency or urgent care services to a beneficiary who has not previously entered into a private contract with the physician or practitioner in accordance with § 405.440.

Subpart E—Criteria for Determining Reasonable Charges

AUTHORITY: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh).


§ 405.500 Basis.

Subpart E is based on the provisions of the following sections of the Act: Section 1814(b) provides for Part A payment on the basis of the lesser of a provider’s reasonable costs or customary charges. Section 1832 establishes the scope of benefits provided under the Part B supplementary medical insurance program. Section 1833(a) sets forth the amounts of payment for supplementary medical insurance services on the basis of the lesser of a provider’s reasonable costs or customary charges. Section 1834(a) specifies how payments are made for the purchase or rental of new and used durable medical equipment for Medicare beneficiaries. Section 1834(b) provides for payment for radiologist services on a fee schedule basis. Section 1834(c) provides for payments and standards for screening mammography. Section 1842(b) sets forth the provisions for a carrier to enter into a contract with the Secretary and to make determinations with respect to Part B claims. Section 1842(h) sets forth the requirements for a physician or supplier to voluntarily enter into an agreement with the Secretary to become a participating physician or supplier. Section 1842(i) sets forth the provisions for the payment of Part B claims. Section 1848 establishes a fee schedule for payment of physician services. Section 1861(b) sets forth the inpatient hospital services covered by the Medicare program. Section 1861(s) sets forth the general authority under which HCFA may establish limits on provider costs recognized as reasonable in determining Medicare program payments. Section 1861(aa) sets forth the rural health clinic services and Federally qualified health center services covered by the Medicare program. Section 1863(j) defines the term “covered osteoporosis drug.” Section 1862(a)(14) lists services that are excluded from coverage. Section 1866(a) specifies the terms for provider agreements. Section 1881 authorizes special rules for the coverage of and payment for services furnished to patients with end-stage renal disease. Section 1886 sets forth the requirements for payment to hospitals for inpatient hospital services. Section 1887 sets forth requirements for payment of provider-based physicians and payment under certain percentage arrangements. Section 1889 provides for Medicare and Medigap information by telephone.

[60 FR 63175, Dec. 8, 1995]

§ 405.501 Determination of reasonable charges.

(a) Except as specified in paragraphs (b), (c), and (d) of this section, Medicare pays no more for Part B medical and other health services than the “reasonable charge” for such service. The reasonable charge is determined by the carriers (subject to any deductible and coinsurance amounts as specified in §§ 410.152 and 410.160 of this chapter).

(b) Part B of Medicare pays on the basis of “reasonable cost” (see part 413 of this chapter) for certain institutional services, certain services furnished under arrangements with institutions, and services furnished by entities that elect to be paid on a cost basis (including health maintenance organizations, rural health clinics, Federally qualified health centers and end-stage renal disease facilities).

(c) Carriers will determine the reasonable charge on the basis of the criteria specified in § 405.502, and the customary and prevailing charge screens in effect when the service was furnished. (Also see §§ 415.55 through 415.70 and §§ 415.100 through 415.130 of this chapter, which pertain to the determination of reimbursement for services.
Health Care Financing Administration, HHS

§ 405.502 Criteria for determining reasonable charges.

(a) Criteria. The law allows for flexibility in the determination of reasonable charges to accommodate reimbursement to the various ways in which health services are furnished and charged for. The criteria for determining what charges are reasonable include:

1. The customary charges for similar services generally made by the physician or other person furnishing such services.
2. The prevailing charges in the locality for similar services.
3. In the case of physicians' services, the prevailing charges adjusted to reflect economic changes as provided under § 405.504 of this subpart.
4. In the case of medical services, supplies, and equipment that are reimbursed on a reasonable charge basis (excluding physicians' services), the inflation-indexed charge as determined under § 405.509.
5. [Reserved]
6. In the case of medical services, supplies, and equipment (including equipment servicing) that the Secretary judges do not generally vary significantly in quality from one supplier to another, the lowest charge levels at which such services, supplies, and equipment are widely and consistently available in a locality.
7. Other factors that may be found necessary and appropriate with respect to a category of service to use in judging whether the charge is inherently reasonable. This includes special reasonable charge limits (which may be either upper or lower limits) established by HCFA or a carrier if it determines that the standard rules for calculating reasonable charges set forth in this subpart result in the grossly deficient or excessive charges. The determination of these limits is described in paragraphs (g) and (h) of this section.
8. In the case of laboratory services billed by a physician but performed by an outside laboratory, the payment levels established in accordance with the criteria stated in § 405.515.
9. Except as provided in paragraph (a)(10) of this section, in the case of services of assistants-at-surgery as defined in § 405.580 in teaching and non-teaching settings, charges that are not more than 16 percent of the prevailing charge in the locality, adjusted by the economic index, for the surgical procedure performed by the primary surgeon. Payment is prohibited for the services of an assistant-at-surgery in surgical procedures for which HCFA has determined that assistants-at-surgery on average are used in less than 5 percent of such procedures nationally.
10. In the case of services of assistants at surgery that meet the exception under § 415.190(c)(2) or (c)(3) of this chapter because the physician is performing a unique, necessary, specialized medical service in the total care of a patient during surgery, reasonable charges consistent with prevailing practice in the carrier's service area rather than the special assistant at surgery rate.

(b) Comparable services limitation. The law also specifies that the reasonable charge cannot be higher than the charge applicable for a comparable service under comparable circumstances to the carriers' own policyholders and subscribers.

(c) Application of criteria. In applying these criteria, the carriers are to exercise judgment based on factual data on the charges made by physicians to patients generally and by other persons...
to the public in general and on special factors that may exist in individual cases so that determinations of reasonable charge are realistic and equitable.

(d) Responsibility of Administration and carriers. Determinations by carriers of reasonable charge are not reviewed on a case-by-case basis by the Health Care Financing Administration, although the general procedures and performance of functions by carriers are evaluated. In making determinations, carriers apply the provisions of the law under broad principles issued by the Health Care Financing Administration. These principles are intended to assure overall consistency among carriers in their determinations of reasonable charge. The principles in §§405.502 through 405.507 establish the criteria for making such determinations in accordance with the statutory provisions.

(e) Determination of reasonable charges under the End-Stage Renal Disease (ESRD) Program—(1) General. Reasonable charges for renal-related items and services (furnished in connection with transplantation or dialysis) must be related to costs and allowances that are reasonable when the treatments are furnished in an effective and economical manner.

(2) Nonprovider (independent) dialysis facilities. Reasonable charges for renal-related items and services furnished before August 1, 1983 must be determined related to costs and charges prior to July, 1973, in accordance with the regulations at §405.541. Items and services related to outpatient maintenance dialysis that are furnished after that date are paid for in accordance with §§405.542 and 413.170 of this chapter.

(3) Provider services and (hospital-based) dialysis facilities. Renal-related items and services furnished by providers, or by ESRD facilities based in hospitals, before August 1, 1983 are paid for under the provider reimbursement provisions found generally in part 413 of this chapter. Items and services related to outpatient maintenance dialysis that are furnished after that date are paid for in accordance with §§405.542 and 413.170 of this chapter.

(4) Physicians’ services. Reasonable charges for renal-related physicians’ services must be determined considering charges made for other services involving comparable physicians’ time and skill requirements, in accordance with regulations at §§405.542 and 405.543.

(f) Determining payments for certain physician services furnished in outpatient hospital settings—(1) General rule. If physician services of the type routinely furnished in physicians’ offices are furnished in outpatient hospital settings before January 1, 1992, carriers determine the reasonable charge for those services by applying the limits described in paragraph (f)(5) of this section.

(2) Definition. As used in this paragraph (f), outpatient settings means—

(i) Hospital outpatient departments, including clinics and emergency rooms; and

(ii) Comprehensive outpatient rehabilitation facilities.

(3) Services covered by limits. The carrier establishes a list of services routinely furnished in physicians’ offices in the area. The carrier has the discretion to determine which professional services are routinely furnished in physicians’ offices, based on current medical practice in the area. Listed below are some examples of routine services furnished by office-based physicians.

Examples

Review of recent history, determination of blood pressure, auscultation of heart and lungs, and adjustment of medication.

Brief history and examination, and initiation of diagnostic and treatment programs.

Treatment of an acute respiratory infection.

(4) Services excluded from limits. The limits established under this paragraph do not apply to the following:

(i) Rural health clinic services.

(ii) Surgical services included on the ambulatory surgical center list of procedures published under §416.69(c) of this chapter.
(iii) Services furnished in a hospital emergency room after the sudden onset of a medical condition manifesting itself by acute symptoms of sufficient severity (including severe pain) such that the absence of immediate medical attention could reasonably be expected to result in—
   (A) Placing the patient's health in serious jeopardy;
   (B) Serious impairment to bodily functions; or
   (C) Serious dysfunction of any bodily organ or part.
(iv) Anesthesiology services and diagnostic and therapeutic radiology services.
(v) Federally qualified health center services paid under the rules in part 405 subpart X.

(5) Methodology for developing limits—
   (i) Development of a charge base. The carrier establishes a charge base for each service identified as a routine office-based physician service. The charge base consists of the prevailing charge in the locality for each such service adjusted by the economic index. The carrier uses the prevailing charges that apply to services by non-specialists in office practices in the locality in which the outpatient setting is located.
   (ii) Calculation of the outpatient limits. The carrier calculates the charge limit for each service by multiplying the charge base amount for each service by .60.

(6) Application of limits. The reasonable charge for physician services of the type described in paragraph (f)(3) of this section that are furnished in an outpatient setting is the lowest of the actual charges, the customary charges in accordance with §405.503, the prevailing charges applicable to these services in accordance with §405.504, or the charge limits calculated in paragraph (f)(5)(ii) of this section.

(g) Determination of payment amounts in special circumstances—(1) General. (i) For purposes of this paragraph, a “category of items or services” may consist of a single item or service or any number of items or services.
   (ii) HCFA or a carrier may determine that the standard rules for calculating Part B payment amounts for a category of items or services identified in section 1861(s) of the Act (other than physician services paid under section 1848 of the Act) will result in grossly deficient or excessive amounts.
   (iii) If HCFA or the carrier determines that the standard rules for calculating payment amounts for a category of items or services set forth in this subpart will result in grossly deficient or excessive amounts, HCFA or the carrier may establish special payment limits that are realistic and equitable for a category of items or services.
   (iv) The limit on the payment amount is either an upper limit to correct a grossly excessive payment amount or a lower limit to correct a grossly deficient payment amount.
   (v) The limit is either a specific dollar amount or is based on a special method to be used in determining the payment amount.
   (vi) Except as provided in paragraph (h) of this section, a payment limit for a given year may not vary by more than 15 percent from the payment amount established for the preceding year.
   (vii) Examples of excessive or deficient payment amounts. Examples of the factors that may result in grossly deficient or excessive payment amounts include, but are not limited to, the following:
      (A) The marketplace is not competitive. This includes circumstances in which the marketplace for a category of items or services is not truly competitive because a limited number of suppliers furnish the item or service.
      (B) Medicare and Medicaid are the sole or primary sources of payment for a category of items or services.
      (C) The payment amounts for a category of items or services do not reflect changing technology, increased facility with that technology, or changes in acquisition, production, or supplier costs.
      (D) The payment amounts for a category of items or services in a particular locality are grossly higher or lower than payment amounts in other comparable localities for the category of items or services, taking into account the relative costs of furnishing the category of items or services in the different localities.

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(E) Payment amounts for a category of items or services are grossly higher or lower than acquisition or production costs for the category of items or services.

(F) There have been increases in payment amounts for a category of items or services that cannot be explained by inflation or technology.

(G) The payment amounts for a category of items or services are grossly higher or lower than the payments made for the same category of items or services by other purchasers in the same locality.

(2) Establishing a limit. In establishing a payment limit for a category of items or services, HCFA or a carrier considers the available information that is relevant to the category of items or services and establishes a payment amount that is realistic and equitable. The factors HCFA or a carrier consider in establishing a specific dollar amount or special payment method for a category of items or services may include, but are not limited to, the following:

(i) Price markup. This is the relationship between the retail and wholesale prices or manufacturer’s costs of a category of items or services. If information on a particular category of items or services is not available, HCFA or a carrier may consider the markup on a similar category of items or services and information on general industry pricing trends.

(ii) Differences in charges. HCFA or a carrier may consider the differences in charges for a category of items or services made to non-Medicare and Medicare patients or to institutions and other large volume purchasers.

(iii) Costs. HCFA or a carrier may consider resources (for example, overhead, time, acquisition costs, production costs, and complexity) required to produce a category of items or services.

(iv) Utilization. HCFA or a carrier may impute a reasonable rate of use for a category of items or services and consider unit costs based on efficient utilization.

(v) Payment amounts in other localities. HCFA or a carrier may consider payment amounts for a category of items or services furnished in another locality.

(3) Notification of limits—(i) National limits. HCFA publishes in the Federal Register proposed and final notices announcing a special payment limit described in this paragraph (g) before it adopts the limit. The notices set forth the criteria and circumstances, if any, under which a carrier may grant an exception to a payment limit for a category of items or services.

(ii) Carrier-level limits. A carrier proposing to establish a special payment limit for a category of items or services must inform the affected suppliers and State Medicaid agencies of the factors it considered in determining and in establishing the limit, as described in paragraphs (g)(1) through (g)(3) of this section, and solicit comments. The carrier must evaluate the comments it receives and inform the affected suppliers, State Medicaid agencies, and HCFA of any final limits it establishes. HCFA acknowledges in writing to the carrier that it received the carrier’s notification. After the carrier has received HCFA’s acknowledgement, the limit may be effective for services furnished at least 30 days after the date of the carrier’s notification.

(h) Special payment limit adjustments greater than 15 percent of the payment amount. In addition to applying the general rules under paragraphs (g)(1) through (g)(3) of this section, HCFA applies the following rules in determining and establishing a payment adjustment greater than 15 percent of the payment amount for a category of items or services within a year:

(1) Potential impact of special limit. HCFA considers the potential impact on quality, access, beneficiary liability, assignment rates, and participation of suppliers.

(2) Supplier consultation. Before making a determination that a payment amount for a category of items or services is not inherently reasonable by reason of its grossly excessive or deficient amount, HCFA consults with representatives of the suppliers likely to be affected by the change in the payment amount.

(3) Publication of national limits. If HCFA determines under this paragraph (h) to establish a special payment limit for a category of items or services, it publishes in the Federal Register...
§ 405.503 Determining customary charges.

(a) Customary charge defined. The term “customary charges” will refer to the uniform amount which the individual physician or other person charges in the majority of cases for a specific medical procedure or service. In determining such uniform amount, token charges for charity patients and substandard charges for welfare and other low income patients are to be excluded. The reasonable charge cannot, except as provided in §405.506, be higher than the individual physician’s or other person’s customary charge. The customary charge for different physicians or other persons may, of course, vary. Payment for covered services would be based on the actual charge for the service when, in a given instance, that charge is less than the amount which the carrier would otherwise have found to be within the limits of acceptable charges for the particular service. Moreover, the income of the individual beneficiary is not to be taken into account by the carrier in determining the amount which is considered to be a reasonable charge for a service rendered to him. There is no provision in the law for a carrier to evaluate the reasonableness of charges in light of an individual beneficiary’s economic status.

(b) Variation of charges. If the individual physician or other person varies his charges for a specific medical procedure or service, so that no one amount is charged in the majority of cases, it will be necessary for the carrier to exercise judgment in the establishment of a “customary charge” for such physician or other person. In making this judgment, an important guide, to be utilized when a sufficient volume of data on the physician’s or other person’s charges is available, would be the median or midpoint of his charges, excluding token and substandard charges as well as exceptional charges on the high side. A significant clustering of charges in the vicinity of the median amount might indicate that a point of such clustering should be taken as the physician’s or other
§ 405.504 Determining prevailing charges.

(a) Ranges of charges. (1) In the case of physicians' services furnished beginning January 1, 1987, the prevailing charges for a nonparticipating physician as defined in this paragraph will be no higher than the same level that was set for services furnished during the previous calendar year for a physician who was a participating physician during that year. A nonparticipating physician is a physician who has not entered into an agreement with the Medicare program to accept payment on an assignment-related basis (in accordance with §424.55 of this chapter) for all items and services furnished to individuals enrolled under Part B of Medicare during a given calendar year.

(b) No charge for Part B medical or other health services may be considered to be reasonable if it exceeds the higher of:

(i) The prevailing charge for similar services in the same locality in effect on December 31, 1970, provided such prevailing charge had been found acceptable by HCFA; or

(ii) The prevailing charge that, on the basis of statistical data and methodology acceptable to HCFA, would cover:

(A) 75 percent of the customary charges made for similar services in the same locality in the 12-month period of July 1 through June 30 preceding the fee screen year (January 1 through December 31) in which the service was furnished; or

(B) In the case of services furnished more than 12 months before the beginning of the fee screen year (January 1 through December 31) in which the claim or request for payment is submitted, 75 percent of the customary charges made for similar services in the same locality during the 12 month period of July 1 through June 30 preceding the fee screen year that ends immediately preceding the fee screen year in which the claim or request for payment is submitted.

(3)(i) In the case of physicians' services, furnished before January 1, 1992, each prevailing charge in each locality may not exceed the prevailing charge determined for the FY ending June 30, 1973 (without reference to the adjustments made in accordance with the economic stabilization program then in effect), except on the basis of appropriate economic index data that demonstrate the higher prevailing charge level is justified by:

(A) Changes in general earnings levels of workers that are attributable to factors other than increases in their productivity; and

(B) changes in expenses of the kind incurred by physicians in office practice. The office-expense component and the earnings component of such index shall be given the relative weights
shown in data on self-employed physicians’ gross incomes.

Example. The available data indicate the office-expense and earnings components of the index should be given relative weights of 40 percent and 60 percent, respectively, and it is calculated that the aggregate increase in expenses of practice for a particular July through June period was 112 percent over the expenses of practice for calendar year 1971 and the increase in earnings (less increases in workers’ productivity) was 110 percent over the earnings for calendar year 1971. The allowable increase in any prevailing charge that could be recognized during the next fee screen year would be 110.8 percent ((.40 \times 112) + (.60 \times 110) = 110.8) above the prevailing charge recognized for fiscal year 1973.

(ii)(A) If the increase in the prevailing charge in a locality for a particular physician service resulting from an aggregate increase in customary charges for that service does not exceed the index determined under paragraph (a)(3)(i) of this section, the increase is permitted and any portion of the allowable increase not used is carried forward and is a basis for justifying increases in that prevailing charge in the future. However, if the increase in the prevailing charge exceeds the allowable increase, the increase will be reduced to the allowable amount. Further increases will be justified only to the degree that they do not exceed further rises in the economic index. The prevailing charge for physicians’ services furnished during the 15-month period beginning July 1, 1984 may not exceed the prevailing charge for physicians’ services in effect for the 12-month period beginning July 1, 1983. The increase in prevailing charges for physicians’ services for subsequent fee screen years similarly may not reflect the rise in the economic index that would have otherwise been provided for the period beginning July 1, 1984, and must be treated as having fully provided for the rise in the economic index which would have been otherwise taken into account.

(B) Notwithstanding the provisions of paragraphs (a)(3)(i) and (ii)(A) of this section, the prevailing charge in the case of a physician service in a particular locality determined pursuant to paragraphs (a)(2) and (3)(i) of this section for the fiscal year beginning July 1, 1975, and for any subsequent fee screen years, if lower than the prevailing charge for the fiscal year ending June 30, 1975, by reason of the application of economic index data, must be raised to such prevailing charge which was in effect for the fiscal year ending June 30, 1975. (If the amount paid on any claim processed by a carrier after the original reasonable charge update for the fiscal year beginning July 1, 1975, and prior to the adjustments required by the preceding sentence, was at least $1 less than the amount due pursuant to the preceding sentence, the difference between the amount previously paid and the amount due shall be paid within 6 months after December 31, 1975; however, no payment shall be made on any claim where the difference between the amount previously paid and the amount due shall be paid within 6 months after December 31, 1975; however, no payment shall be made on any claim where the difference between the amount previously paid and the amount due is less than $1.)

(iii) If, for any reason, a prevailing charge for a service in a locality has no precise counterpart in the carrier’s charge data for calendar year 1971 (the data on which the prevailing charge calculations for fiscal year 1973 were based), the limit on the prevailing charge will be estimated, on the basis of data and methodology acceptable to HCFA, to seek to produce the effect intended by the economic index criterion. The allowance or reduction of an increase in a prevailing charge for any individual medical item or service may affect the allowance or reduction of an increase in the prevailing charges for other items or services if, for example, the limit on the prevailing charge is estimated, or if the prevailing charges for more than one item or service are established through the use of a relative value schedule and dollar conversion factors.

(b) Variation in range of prevailing charges. The range of prevailing charges in a locality may be different for physicians or other persons who engage in a specialty practice or service than for others. Existing differentials in the level of charges between different kinds of practice or service could, in some localities, lead to the
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development of more than one range of prevailing charges for application by the carrier in its determinations of reasonable charges. Carrier decisions in this respect should be responsive to the existing patterns of charges by physicians and other persons who render covered services, and should establish differentials in the levels of charges between different kinds of practice or service only where in accord with such patterns.

(c) Re-evaluation and adjustment of prevailing charges. Determinations of prevailing charges by the carrier are to be re-evaluated and adjusted from time to time on the basis of factual information about the charges made by physicians and other persons to the public in general. This information should be obtained from all possible sources including a carrier’s experience with its own programs as well as with the supplementary medical insurance program.

(d) Computation and issuance of the MEI after CY 1992—(1) For update years after CY 1992, the MEI is a physician input price index, in which the annual percent changes for the direct-labor price components are adjusted by an annual percent change in a 10-year moving average index of labor productivity in the nonfarm business sector.

(2) The MEI is constructed, using as a base year, CY 1989 weights and annual percent changes in the economic price proxies as shown in the following chart:

<table>
<thead>
<tr>
<th>MEDICARE ECONOMIC INDEX EXPENDITURE CATEGORIES, WEIGHTS, AND PRICE PROXIES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expense category</td>
</tr>
<tr>
<td>Total</td>
</tr>
<tr>
<td>1. Physician’s Own Time (net income, general earnings)</td>
</tr>
<tr>
<td>a. Wages and Salaries</td>
</tr>
<tr>
<td>b. Fringe Benefits</td>
</tr>
<tr>
<td>2. Physician Practice Expense</td>
</tr>
<tr>
<td>a. Non-physician Employee Compensation</td>
</tr>
<tr>
<td>(1) Wages and Salaries</td>
</tr>
<tr>
<td>(2) Fringe Benefits</td>
</tr>
<tr>
<td>b. Office Expense</td>
</tr>
<tr>
<td>c. Medical Materials and Supplies</td>
</tr>
<tr>
<td>d. Professional Liability Insurance</td>
</tr>
<tr>
<td>e. Medical Equipment</td>
</tr>
<tr>
<td>f. Other Professional Expense</td>
</tr>
<tr>
<td>(1) Professional Car</td>
</tr>
<tr>
<td>(2) Other</td>
</tr>
</tbody>
</table>


² Due to rounding, weights may not sum to 100.0%.

³ All price proxies are for annual percent changes for the 12 months ending June 30th.

⁴ Annual percent change values for Physicians’ Own Time and Non-physician Employee Compensation are net of the change in the 10-year moving average of output per man-hour to exclude changes in non-farm business sector labor productivity.

(3) If there is no methodological change, HCFA publishes a notice in the FEDERAL REGISTER to announce the annual increase in the MEI before the beginning of the update year to which it applies. If there are changes in the base year weights or price proxies, or if there are any other MEI methodological changes, they are published in the FEDERAL REGISTER with an opportunity for public comment.

Determinations of locality.

``Locality'' is the geographical area for which the carrier is to derive the reasonable charges or fee schedule amounts for services or items. Usually, a locality may be a State (including the District of Columbia, a territory, or a Commonwealth), a political or economic subdivision of a State, or a group of States. It should include a cross section of the population with respect to economic and other characteristics. Where people tend to gravitate toward certain population centers to obtain medical care or service, localities may be recognized on a basis constituting medical services areas (interstate or otherwise), comparable in concept to “trade areas.” Localities may differ in population density, economic level, and other major factors affecting charges for services. Carriers therefore shall delineate localities on the basis of their knowledge of local conditions. However, distinctions between localities are not to be so finely made that a locality includes only a very limited geographic area whose population has distinctly similar income characteristics (e.g., a very rich or very poor neighborhood within a city).

Charges higher than customary or prevailing charges or lowest charge levels.

A charge which exceeds the customary charge of the physician or other person who rendered the medical or other health service, or the prevailing charge in the locality, or an applicable lowest charge level may be found to be reasonable, but only where there are unusual circumstances, or medical complications requiring additional time, effort or expense which support an additional charge, and only if it is acceptable medical or medical service practice in the locality to make an extra charge in such cases. The mere fact that the physician’s or other person’s customary charge is higher than prevailing would not justify a determination that it is reasonable.

Illustrations of the application of the criteria for determining reasonable charges.

The following examples illustrate how the general criteria on customary charges and prevailing charges might be applied in determining reasonable charges under the supplementary medical insurance program. Basically, these examples demonstrate that, except where the actual charge is less, reasonable charges will reflect current customary charges of the particular physician or other person within the ranges of the current prevailing charges in the locality for that type and level of service:

- The prevailing charge for a specific medical procedure ranges from $80 to $100 in a certain locality.
- Doctor A’s bill is for $75 although he customarily charges $80 for the procedure.
- Doctor B’s bill is his customary charge of $85.
- Doctor C’s bill is his customary charge of $125.
- Doctor D’s bill is for $100, although he customarily charges $80, and there are no special circumstances in the case.

The reasonable charge for Doctor A would be limited to $75 since under the law the reasonable charge cannot exceed the actual charge, even if it is lower than his customary charge and below the prevailing charges for the locality.

The reasonable charge for Doctor B would be $85, because it is his customary charge and it falls within the range of prevailing charges.

The reasonable charge for Doctor C could not be more than $100, the top of the range of prevailing charges.

The reasonable charge for Doctor D would be $80, because that is his customary charge. Even though his actual charge of $100 falls within the range of prevailing charges, the reasonable charge cannot exceed his customary charge in the absence of special circumstances.
§ 405.509 Determining the inflation-indexed charge.

(a) Definition. For purposes of this section, inflation-indexed charge means the lowest of the fee screens used to determine reasonable charges (as determined in § 405.503 for the customary charge, § 405.504 for the prevailing charge, this section for the inflation-indexed charge, and § 405.511 for the lowest charge level) for services, supplies, and equipment reimbursed on a reasonable charge basis (excluding physicians' services), that is in effect on December 31 of the previous fee screen year, updated by the inflation adjustment factor, as described in paragraph (b) of this section.

(b) Application of inflation adjustment factor to determine inflation-indexed charge. (1) For fee screen years beginning on or after January 1, 1987, the inflation-indexed charge is determined by updating the fee screen used to determine the reasonable charges in effect on December 31 of the previous fee screen year by application of an inflation adjustment factor, that is, the annual change in the level of the consumer price index for all urban consumers, as compiled by the Bureau of Labor Statistics, for the 12-month period ending on June 30 of each year.

(2) For services, supplies, and equipment furnished from October 1, 1985 through December 31, 1986 the inflation adjustment factor is zero.

(c) The inflation-indexed charge does not apply to any services, supplies, or equipment furnished after December 31, 1991, that are covered under or limited by the fee schedule for physicians' services established under section 1848 of the Act and part 415 of this chapter. These services are subject to the Medicare Economic Index described in § 415.30 of this chapter.

(iv) The lowest charge level at which the item or service is widely and consistently available in the locality (see paragraph (c) of this section); or
(v) The inflation-indexed charge, as determined under §405.509, in the case of medical services, supplies, and equipment that are reimbursed on a reasonable charge basis (excluding physicians' services).

(2) In the case of laboratory services, paragraph (a)(1) of this section is applicable to services furnished by physicians in their offices, by independent laboratories (see §405.1310(a)) and to services furnished by a hospital laboratory for individuals who are neither inpatients nor outpatients of a hospital. Allowance of additional charges exceeding the lowest charge level can be approved by the carrier on the basis of unusual circumstances or medical complications in accordance with §405.506.

(b) Public notice of items and services subject to the lowest charge level rule. Before the Secretary determines that lowest charge levels should be established for an item or service, notice of the proposed determination will be published with an opportunity for public comment. The descriptions or specifications of items or services in the notice will be in sufficient detail to permit a determination that items or services conforming to the descriptions will not vary significantly in quality.

(c) Calculating the lowest charge level. The lowest charge level at which an item or service is widely and consistently available in a locality is calculated by the carrier in accordance with instructions from HCFA as follows:

(1) For items or services furnished on or before December 31, 1986.
   (i) A lowest charge level is calculated for each identified item or service in January and July of each year.
   (ii) The lowest charge level for each identified item or service is set at the 25th percentile of the charges (incurred or submitted on claims processed by the carrier) for that item or service, in the locality designated by the carrier for this purpose, during the 3-month period of July 1 through September 30 preceding the fee screen year (January 1 through December 31) for which the item or service was furnished.

(2) For items or services furnished on or after January 1, 1987.
   (i) A lowest charge level is calculated for each identified item or service in January of each year.
   (ii) The lowest charge level for each identified item or service is set at the 25th percentile of the charges (incurred or submitted on claims processed by the carrier) for that item or service, in the locality designated by the carrier for this purpose, during the 3-month period of July 1 through September 30 preceding the fee screen year (January 1 through December 31) for which the item or service was furnished.

(3) Lowest charge levels for laboratory services. In setting lowest charge levels for laboratory services, the carrier will consider only charges made for laboratory services performed by physicians in their offices, by independent laboratories which meet coverage requirements, and for services furnished by a hospital laboratory for individuals who are neither inpatients nor outpatients of a hospital.

(d) Locality. Subject to the approval of the Secretary, the carrier may designate its entire service area as the locality for purposes of this section, or may otherwise modify the localities used for calculating prevailing charges. (The modified locality for an item or service will also be used for calculating the prevailing charge for that item or service.)

(5) Health Care Financing Administration, HHS § 405.512 Carriers' procedural terminology and coding systems.

(a) General. Procedural terminology and coding systems are designed to provide physicians and third party payers with a common language that accurately describes the kinds and levels of services provided and that can serve as a basis for coverage and payment determinations.
§ 405.515 Modification of terminology and/or coding systems.

A carrier that wishes to modify its system of procedural terminology and coding shall submit its request to the Health Care Financing Administration with all pertinent data and information for approval before the revision is implemented. The Health Care Financing Administration will evaluate the proposal in the light of the guidelines specified in paragraph (c) of this section and such other considerations as may be pertinent, and consult with the Assistant Secretary for Health. The Health Care Financing Administration will approve such a revision if it determines that the potential advantages of the proposed new system, outweigh the disadvantages.

(c) Guidelines. The following considerations and guidelines are taken into account in evaluating a carrier's proposal to change its system of procedural terminology and coding:

(1) The rationale for converting to the new terminology and coding;
(2) The estimated short-run and long-run impact on the cost of the health insurance program, other medical care costs, administrative expenses, and the reliability of the estimates;
(3) The degree to which the conversion to the proposed new terminology and coding can be accomplished in a way that permits full implementation of the reasonable charge criteria in accordance with the provisions of this subpart;
(4) The degree to which the proposed new terminology and coding are accepted by physicians in the carrier's area (physician acceptance is assumed only if a majority of the Medicare and non-Medicare bills and claims completed by physicians in the area and submitted to the carrier can reasonably be expected to utilize the proposed new terminology and coding);
(5) The extent to which the proposed new terminology and coding system is used by the carrier in its non-Medicare business;
(6) The clarity with which the proposed system defines its terminology and whether the system lends itself to:
   (i) Accurate determinations of coverage;
   (ii) Proper assessment of the appropriate level of payment; and
   (iii) Meeting the carrier's or Professional Standards Review Organizations' review needs and such other review needs as may be appropriate;
(7) Compatibility of the new terminology and coding system with other systems that the carrier and other carriers may utilize in the administration of the Medicare program—e.g., its compatibility with systems and statistical requirements and with the historical data in the carrier's processing system; and
(8) Compatibility of the proposed system with the carriers methods for determining payment under the fee schedule for physicians' services for services which are identified by a single element of terminology but which may vary in content.


§ 405.515 Reimbursement for clinical laboratory services billed by physicians.

This section implements section 1842(h) of the Social Security Act, which places a limitation on reimbursement for markups on clinical laboratory services billed by physicians. If a physician's bill, or a request for payment for a physician's services, includes a charge for a laboratory test for which payment may be made under this part, the amount payable with respect to the test shall be determined as follows (subject to the coinsurance and deductible provisions at §§ 410.152 and 410.160 of this chapter):

(a) If the bill or request for payment indicates that the test was personally performed or supervised either by the physician who submitted the bill (or for whose services the request for payment was made), or by another physician with whom that physician shares his or her practice, the payment will be based on the physician's reasonable charge for the test (as determined in accordance with § 405.502).

(b) If the bill or request for payment indicates that the test was performed by an outside laboratory, and identifies both the laboratory and the amount the laboratory charged, payment for the test will be based on the lower of—
Health Care Financing Administration, HHS

§ 405.520 Payment for a physician assistant's, nurse practitioner's, and clinical nurse specialists' services and services furnished incident to their professional services.

(a) General rule. A physician assistant's, nurse practitioner's, and clinical nurse specialists' services, and services and supplies furnished incident to their professional services, are paid in accordance with the physician fee schedule. The payment for a physician assistant's services may not exceed the limits at §414.52 of this chapter. The payment for a nurse practitioners' and clinical nurse specialists' services may not exceed the limits at §414.56 of this chapter.

(b) Requirements. Medicare payment is made only if all claims for payment are made on an assignment-related basis in accordance with §424.55 of this chapter, that sets forth, respectively, the conditions for coverage of physician assistants' services, nurse practitioners' services and clinical nurse specialists' services, and services and supplies furnished incident to their professional services.

(c) Civil money penalties. Any person or entity who knowingly and willingly bills a Medicare beneficiary amounts in excess of the appropriate coinsurance
and deductible is subject to a civil money penalty not to exceed $2,000 for each bill or request for payment.

[63 FR 58905, Nov. 2, 1998]

§ 405.534 Limitation on payment for screening mammography services.

(a) Basis and scope. This section implements section 1834(c) of the Act by establishing a limit on payment for screening mammography examinations. There are three categories of billing for screening mammography services. Those categories and the payment limitations on each are set forth in paragraphs (b) through (d) of this section.

(b) Global or complete service billing representing both the professional and technical components of the procedure. If a fee is billed for a global service, the amount of payment subject to the deductible is equal to 80 percent of the least of the following:

(1) The actual charge for the service.

(2) The amount established for the global procedure for a diagnostic bilateral mammogram under the fee schedule for physicians' services set forth at part 414, subpart A.

(3) The payment limit for the procedure. For screening mammography services furnished in CY 1994, the payment limit is $59.63. On January 1 of each subsequent year, the payment limit is updated by the percentage increase in the Medicare Economic Index (MEI) and reflects the relationship between the relative value units for the professional and technical components of a diagnostic bilateral mammogram under the fee schedule for physicians' services.

(c) Professional component billing representing only the physician's interpretation for the procedure. If the professional component of screening mammography services billed separately, the amount of payment for that professional component, subject to the deductible, is equal to 80 percent of the least of the following:

(1) The actual charge for the professional component of the service.

(2) The amount established for the professional component of a diagnostic bilateral mammogram under the fee schedule for physicians' services.

(3) The professional component of the payment limit for screening mammography services described in paragraph (b)(3) of this section.

(d) Technical component billing representing other resources involved in furnishing the procedure. If the technical component of screening mammography services is billed separately, the amount of payment, subject to the deductible, is equal to 80 percent of the least of the following:

(1) The actual charge for the technical component of the service.

(2) The amount established for the technical component of a diagnostic bilateral mammogram under the fee schedule for physicians' services.

(3) The technical component of the payment limit for screening mammography services described in paragraph (b)(3) of this section.


§ 405.535 Special rules for nonparticipating physicians and suppliers furnishing screening mammography services.

If screening mammography services are furnished to a beneficiary by a nonparticipating physician or supplier that does not accept assignment, a limiting charge applies to the charges billed to the beneficiary. The limiting charge is the lesser of the following:

(a) 115 percent of the payment limit set forth in §405.534(b)(3), (c)(3), and (d)(3) (limitations on the global service, professional component, and technical component of screening mammography services, respectively).

(b) The limiting charge for the global service, professional component, and technical component of a diagnostic bilateral mammogram under the fee schedule for physicians' services set forth at §414.48(b) of this chapter.


Subpart F—[Reserved]

Subpart G—Reconsiderations and Appeals Under Medicare Part A

Authority: SECS. 1102, 1155, 1869(b), 1871, 1872, and 1879 of the Social Security Act (42
§ 405.701 Basis, purpose and definitions.

(a) This subpart implements section 1869 of the Social Security Act. Section 1869(a) provides that the Secretary will make determinations about the following matters, and section 1869(b) provides for a hearing for an individual who is dissatisfied with the Secretary's determination as to:

(1) Whether the individual is entitled to hospital insurance (part A) or supplementary medical insurance (part B) under title XVIII of the Act; or

(2) The amount payable under hospital insurance.

(b) This subpart establishes the procedures governing initial determinations, reconsidered determinations, hearings, and final agency review, and the reopening of determinations and decisions that are applicable to matters arising under paragraph (a) of this section.

(c) Subparts J and R of 20 CFR part 404 (dealing with determinations, the administrative review process and representation of parties) are also applicable to matters arising under paragraph (a) of this section, except to the extent that specific provisions are contained in this subpart.

(d) Definitions. As used in subpart G of this part, the term—

Appellant designates the beneficiary, provider or other person or entity that has filed an appeal concerning a particular determination of benefits under Medicare part A. Designation as an appellant does not in itself convey standing to appeal the determination in question.

Common issues of law and fact, with respect to the aggregation of claims by two or more appellants to meet the minimum amount in controversy needed for a hearing, means like or coordinated services or items provided to the same beneficiary by the appellants.


§ 405.702 Notice of initial determination.

After a request for payment under part A of title XVIII of the Act is filed with the intermediary by or on behalf of the individual who received inpatient hospital services, extended care services, or home health services, and the intermediary has ascertained whether the items and services furnished are covered under part A of title XVIII, and where appropriate, ascertained and made payment of amounts due or has ascertained that no payments were due, the individual will be notified in writing of the initial determination in his case. In addition, if the items or services furnished such individual are not covered under part A of title XVIII by reason of § 411.15(g) or § 411.15(k) and payment may not be made for such items or services under § 411.400 only because the requirements of § 411.400(a)(2) are not met, the provider of services which furnished such items or services will be notified in writing of the initial determination in such individual’s case. These notices shall be mailed to the individual and the provider of services at their last known addresses and shall state in detail the basis for the determination. Such written notices shall also inform the individual and the provider of services of their right to reconsideration of the determination if they are dissatisfied with the determination.

[55 FR 11020, Mar. 26, 1990]

§ 405.704 Actions which are initial determinations.

(a) Applications and entitlement of individuals. An initial determination with respect to an individual includes the following—
§ 405.704

(1) A determination with respect to entitlement to hospital insurance or supplementary medical insurance;
(2) A disallowance of an individual’s application for entitlement to hospital or supplementary medical insurance, if the individual fails to submit evidence requested by SSA to support the application. SSA will specify in the initial determination the conditions of entitlement that the applicant failed to establish by not submitting the requested evidence;
(3) A denial of a request for withdrawal of an application for hospital or supplementary medical insurance;
(4) A denial of a request for cancellation of a “request for withdrawal”; and
(5) A determination as to whether an individual, previously determined to be entitled to hospital or supplementary medical insurance, is no longer entitled to such benefits, including a determination based on nonpayment of premiums.

(b) Requests for payment by or on behalf of individuals. An initial determination with respect to an individual includes any determination made on the basis of a request for payment by or on behalf of the individual under part A of Medicare, including a determination with respect to:
(1) The coverage of items and services furnished;
(2) The amount of an applicable deductible;
(3) The application of the coinsurance feature;
(4) The number of days of inpatient hospital benefits utilized during a spell of illness or for purposes of the inpatient psychiatric hospital 190-day lifetime maximum;
(5) The number of days of the 60-day lifetime reserve utilized for inpatient hospital coverage;
(6) The number of days of posthospital extended care benefits utilized;
(7) The number of home health visits utilized;
(8) The physician certification requirement;
(9) The request for payment requirement;
(10) The beginning and ending of a spell of illness, including a determination made under the presumptions established under §409.60(c)(2) of this chapter, as specified in §409.60(c)(4) of this chapter.
(11) The medical necessity of services (See parts 466 and 473 of this chapter for provisions pertaining to initial and reconsidered determinations made by a P.R.O);
(12) When services are excluded from coverage as custodial care (§411.15(g)) or as not reasonable and necessary (§411.15(k)), whether the individual or the provider of services who furnished the services, or both, knew or could reasonably have been expected to know that the services were excluded from coverage (see §411.402);
(13) Any other issues having a present or potential effect on the amount of benefits to be paid under part A of Medicare, including a determination as to whether there has been an overpayment or underpayment of benefits paid under part A, and if so, the amount thereof; and
(14) Whether a waiver of adjustment or recovery under sections 1870(b) and (c) of the Act is appropriate when an overpayment of hospital insurance benefits or supplementary medical insurance benefits (including a payment under section 1814(e) of the Act) has been made with respect to an individual.

(c) Initial determination with respect to a provider of services. An initial determination with respect to a provider of services shall be a determination made on the basis of a request for payment filed by the provider under part A of Medicare on behalf of an individual who was furnished items or services by the provider, but only if the determination involves the following:
(1) A finding by the intermediary that such items or services are not covered by reason of §411.15(g) or §411.15(k); and
(2) A finding by the intermediary that either such individual or such provider of services, or both, knew or could reasonably have been expected to know that such items or services were excluded from coverage under the program.

[55 FR 11020, Mar. 26, 1990]
§ 405.705 Actions which are not initial determinations.

An initial determination under Part A of Medicare does not include determinations relating to:

(a) The reasonable cost of items or services furnished under Part A of Medicare;
(b) Whether an institution or agency meets the conditions for participation in the program;
(c) Whether an individual is qualified for use of the expedited appeals process as provided in §405.718;
(d) An action regarding compromise of a claim arising under the Medicare program, or termination or suspension of collection action on such a claim under the Federal Claims Collection Act of 1966 (31 U.S.C. 3711). See 20 CFR 404.515 for overpayment claims against an individual, §405.376 for overpayment claims against a provider, physician or other supplier, and §408.110 for claims concerning unpaid Medicare premiums;
(e) The transfer or discharge of residents of skilled nursing facilities in accordance with §483.12 of this chapter; or
(f) The preadmission screening and annual resident review processes required by part 483 subparts C and E of this chapter.


§ 405.706 Decisions of utilization review committees.

(a) General rule. A decision of a utilization review committee is a medical determination by a staff committee of the provider or a group similarly composed and does not constitute a determination by the Secretary within the meaning of section 1869 of the Act. The decision of a utilization review committee may be considered by HCFA along with other pertinent medical evidence in determining whether or not an individual has the right to have payment made under Part A of title XVIII.

(b) Applicability under the prospective payment system. HCFA may consider utilization review committee decisions related to inpatient hospital services paid for under the prospective payment system (see part 412 of this chapter) only as those decisions concern:

(1) The appropriateness of admissions resulting in payments under subparts D, E and G of part 412 of this chapter.
(2) The covered days of care involved in determinations of outlier payments under §412.80(a)(1)(i) of this chapter; and
(3) The necessity of professional services furnished in high cost outliers under §412.80(a)(1)(ii) of this chapter.

[48 FR 39831, Sept. 1, 1983]

§ 405.708 Effect of initial determination.

(a) The initial determination under §405.704 (a) or (b) shall be binding upon the individual on whose behalf payment under part A has been requested or, if such individual is deceased, upon the representative of such individual's estate, unless it is reconsidered in accordance with §§405.710 through 405.717 or revised in accordance with §405.750. Such individual (or the representative of such individual's estate if the individual is deceased) shall be the party to such initial determination.

(b) The initial determination under §405.704(c) shall be binding upon the provider of services unless it is reconsidered in accordance with §§405.710 through 405.717 or revised in accordance with §405.750. Such provider of services shall be the party to such initial determination.


§ 405.710 Right to reconsideration.

(a) An individual who is a party to an initial determination, as specified in §405.704 (a) and (b), (or if such individual is deceased, the representative of such individual's estate) and who is dissatisfied with the initial determination may request a reconsideration of such determination in accordance with §405.711 regardless of the amount in controversy.

(b) A provider of services who is a party to an initial determination (as specified in §405.704(c)) and who is dissatisfied with such initial determination may request a reconsideration of such determination in accordance with §405.711, regardless of the amount in controversy.
§ 405.711 Time and place of filing request for reconsideration.

The request for reconsideration shall be made in writing and filed at an office of the SSA or the HCFA or, in the case of a qualified railroad retirement beneficiary (see 20 CFR 404.368) filed at the Railroad Retirement Board, within 60 days after the date of receipt of notice of initial determination, unless such time is extended as provided in § 405.712. A request for reconsideration which is filed with the intermediary which received the request for payment shall be considered to have been filed at the HCFA as of the date it is filed with the intermediary.


§ 405.712 Extension of time to request reconsideration.

If a party to an initial determination desires to file a request for reconsideration after the time for filing such request in accordance with § 405.711 has passed, such party may file a petition with the SSA or the HCFA or, in the case of a qualified railroad retirement beneficiary, with the Railroad Retirement Board, for an extension of time for the filing of such request. Such petition shall be in writing and shall state the reasons why the request for reconsideration was not filed within the required time. For good cause shown, the HCFA may extend the time for filing the request for reconsideration.


§ 405.714 Withdrawal of request for reconsideration.

A request for reconsideration may be withdrawn by the party to the initial determination who filed the request or by his representative provided that the withdrawal is made in writing and filed at an office of the SSA or the HCFA or, in the case of a qualified railroad retirement beneficiary, with the Railroad Retirement Board prior to the date of the mailing of the notice of reconsideration. A withdrawal filed with the intermediary which received the request for payment submitted on behalf of the individual is considered to have been filed with the HCFA as of the date it is filed with the intermediary.


§ 405.715 Reconsidered determination.

(a) In reconsidering an initial determination, the HCFA shall review such initial determination, the evidence and findings upon which such determination was based, and any additional evidence submitted to the SSA or the HCFA or otherwise obtained by the intermediary or the HCFA; and shall make a determination affirming or revising, in whole or in part, such initial determination.

(b) If the request for reconsideration is filed by an individual with respect to an initial determination specified in § 405.704(b)(12), the provider of services who furnished the items or services shall, prior to the making of the reconsidered determination, be made a party thereto. If pursuant to § 405.710(b) a request for reconsideration is filed by a provider of services with respect to an individual determination under § 405.704(c), the individual who was furnished the items or services shall, prior...
§ 405.716 Notice of reconsidered determination.

Written notice of the reconsidered determination shall be mailed by the HCFA to the parties and their representatives at their last known addresses. Such notice shall state the specific reasons for the reconsidered determination and shall advise the parties of their right to a hearing if the amount in controversy is $100 or more, or, if appropriate, advise them of the requirements for use of the expedited appeals process (see § 405.718).

§ 405.717 Effect of a reconsidered determination.

The reconsidered determination is binding upon all parties unless—
(a) A request for a hearing is filed with SSA or HCFA within 60 days after the date of receipt of notice of the reconsidered determination by the parties (for purposes of this section, the date of receipt of notice of the reconsidered determination is presumed to be 5 days after the date of the notice, unless it is shown that the notice was received earlier or later); or
(b) The reconsidered determination is revised in accordance with § 405.750; or
(c) The expedited appeals process is used in accordance with § 405.718.

§ 405.718 Expedited appeals process.

(a) Conditions for use of expedited appeals process (EAP). A party may use the EAP to request court review in place of an administrative law judge (ALJ) hearing or Departmental Appeals Board (DAB) review if the following conditions are met:
(1) HCFA has made a reconsideration determination; an ALJ has made a hearing decision; or DAB review has been requested, but a final decision has not been issued;
(2) The filing entity is a party referred to in § 405.718(d);
(3) The party has filed a request for an ALJ hearing in accordance with § 405.722, or DAB review in accordance with 20 CFR 404.968;
(4) The amount remaining in controversy is $1,000 or more.
(5) If there is more than one party to the reconsideration determination or hearing decision, each party concurs, in writing, with the request for the EAP.
(b) Content of the request for EAP. The request for the EAP:
(1) Alleges that there are no material issues of fact in dispute; and
(2) Asserts that the only factor precluding a decision favorable to the party is a statutory provision that is unconstitutional or a regulation, national coverage decision under section 1862(a)(1) of the Act, or HCFA Ruling that is invalid.
(c) Place and time for requesting an EAP—(1) Place for filing request. The person must file a written request—
(i) At an office of SSA or HCFA; or
(ii) If the person is in the Philippines, at the Veterans Administration Regional Office or with an ALJ; or
(iii) If the person is a qualified railroad retirement beneficiary, at an office of the Railroad Retirement Board.
(2) Time of filing request. The party may file a request for the EAP—
(i) If the party has requested a hearing, at any time prior to receipt of the notice of the ALJ’s decision;
(ii) Within 60 days after the date of receipt of notice of the ALJ’s decision or dismissal, unless the time is extended in accordance with the standards set out in 20 CFR 404.925(c). For purposes of this section, the date of receipt of the notice is presumed to be 5 days after the date of the notice, unless it is shown that the notice was received later; or
(iii) If the party has requested DAB review, at any time prior to receipt of notice of the Board’s decision.
(d) Parties to the EAP. The parties to the EAP are the persons who were parties to the reconsideration determination and, if appropriate, to the hearing.
(e) Determination on request for EAP.
(1) For EAP requests initiated at the ALJ level, an ALJ determines whether all conditions of paragraphs (a) and (b) of this section are met.
§ 405.720  Hearing; right to hearing.

A person has a right to a hearing regarding any initial determination made under § 405.704 if:
(a) Such initial determination has been reconsidered by the HCFA;
(b) Such person was a party to the reconsidered determination;
(c) Such person or his representative has filed a written request for a hearing in accordance with the procedure described in § 405.722; and
(d) The amount in controversy is $100 or more.


§ 405.722  Time and place of filing request for a hearing.

The request for a hearing shall be made in writing and filed at an office of the SSA or the HCFA or with an ALJ, or, in the case of a qualified railroad retirement beneficiary, at an office of the Railroad Retirement Board. Such request must be filed within 60 days after the date of receipt of notice of the reconsidered determination by such individual, except where the time is extended as provided in 20 CFR 404.933(c). For purposes of this section, the date of receipt of notice of the reconsidered determination shall be presumed to be 5 days after the date of such notice, unless there is a reasonable showing to the contrary.


§ 405.724  Departmental Appeals Board (DAB) review.

Regulations beginning at 20 CFR 404.967 regarding SSA Appeals Council Review are also applicable to DAB review of matters addressed by this subpart.


§ 405.730  Court review.

(a) To the extent authorized by sections 1869, 1876(c)(5)(B), and 1879(d) of the Act, a party to a Departmental Appeals Board (DAB) decision or an ALJ decision if the DAB does not review the ALJ decision, may obtain a court review if the amount remaining in controversy is $1,000 or more. A party may obtain court review by filing a civil action in a district court of the United States in accordance with the provisions of section 205(g) of the Act. The filing procedure is set forth at 20 CFR 422.210.

(b) A party to a reconsidered determination or an ALJ hearing decision may obtain a court review if the amount in controversy is $1,000 or more, and he or she requests and meets
§ 405.732 Review of national coverage decisions (NCDs).

(a) General. (1) HCFA makes NCDs either granting, limiting, or excluding Medicare coverage for a specific medical service, procedure or device. NCDs are made under section 1862(a)(1) of the Act or other applicable provisions of the Act. An NCD is binding on all Medicare carriers, fiscal intermediaries, PROs, HMOs, CMPs, and HCPPs when published in HCFA program manuals or the FEDERAL REGISTER.

(2) Under section 1869(b)(3) of the Act, only NCDs made under section 1862(a)(1) of the Act are subject to the conditions of paragraphs (b) through (d) of this section.

(b) Review by ALJ. (1) An ALJ may not disregard, set aside, or otherwise review an NCD.

(2) An ALJ may review the facts of a particular case to determine whether an NCD applies to a specific claim for benefits and, if so, whether the NCD has been applied correctly to the claim.

(c) Review by Court. (1) A court’s review of an NCD is limited to whether the record is incomplete or otherwise lacks adequate information to support the validity of the decision, unless the case has been remanded to the Secretary to supplement the record regarding the NCD. The court may not invalidate an NCD except upon review of the supplemented record.

(2) A Federal court may not hold unlawful or set aside an NCD because it was not issued in accordance with the notice and comment procedures of the Administrative Procedure Act (5 U.S.C. 553) or section 1871(b) of the Act.

(d) Remands—(1) Secretary’s action. When a court remands an NCD matter to the Secretary because the record in support of the NCD is incomplete or otherwise lacks adequate information, the Secretary remands the case to HCFA in order to supplement the record.

(2) Remand to HCFA. HCFA supplements the record with new or updated evidence, including additional information from other sources, and may issue a revised NCD.

(3) Final Actions. (i) The proceedings to supplement the record are expedited.

(ii) When HCFA does not issue a revised NCD, it returns the supplemented record to the court for review.

(iii) When HCFA issues a revised NCD, it forwards the case to an ALJ who issues a new decision applying the revised NCD to the facts of the claim(s) under consideration. The ALJ’s decision is subject to DAB review and, ultimately, judicial review.


§ 405.740 Principles for determining the amount in controversy.

(a) Individual appellants. For the purpose of determining whether an individual appellant meets the minimum amount in controversy needed for a hearing ($100), the following rules apply:

(1) The amount in controversy is computed as the actual amount charged the individual for the items and services in question, less any amount for which payment has been made by the intermediary and less any deductible and coinsurance amounts applicable in the particular case.

(2) A single beneficiary may aggregate claims from two or more providers to meet the $100 hearing threshold and a single provider may aggregate claims for services provided to one or more beneficiaries to meet the $100 hearing threshold.

(3) In either of the circumstances specified in paragraph (a)(2) of this section, two or more claims may be aggregated by an individual appellant only if the claims have previously been reconsidered and a request for hearing has been made within 60 days after receipt of the reconsideration determination(s).

(4) When requesting a hearing, the appellant must specify in his or her appeal request the specific claims to be aggregated.

(b) Two or more appellants. As specified below, under section 1869(b)(2) of the Act, two or more appellants may aggregate their claims together to meet the minimum amount in controversy needed for a hearing ($100).
§ 405.745 Amount in controversy

The right to aggregate under this statutory provision applies to claims for items and services furnished on or after January 1, 1987.

(1) The aggregate amount in controversy is computed as the actual amount charged the individual(s) for the items and services in question, less any amount for which payment has been made by the intermediary and less any deductible and coinsurance amounts applicable in the particular case.

(2) In determining the amount in controversy, two or more appellants may aggregate their claims together under the following circumstances:
   (i) Two or more beneficiaries may combine claims representing services from the same or different provider(s) if the claims involve common issues of law and fact;
   (ii) Two or more providers may combine their claims if the claims involve the delivery of similar or related services to the same beneficiary; or
   (iii) Two or more providers may combine their claims if the claims involve common issues of law and fact with respect to services furnished to two or more beneficiaries.

(f) Notwithstanding the provisions of paragraphs (a)(1) and (b)(1) of this section, when payment is made for certain excluded services under §411.400 of this chapter or the liability of the beneficiary for those services is limited under §411.402 of this chapter, the amount in controversy is computed as the amount that would have been charged the beneficiary for the items or services in question, less any deductible and coinsurance amounts applicable in the particular case, had such expenses not been paid pursuant to §411.400 of this chapter or had such liability not been limited pursuant to §411.402 of this chapter.

(g) Under this subpart, an appellant may not combine part A and part B claims together to meet the requisite amounts in controversy for a hearing. HMO, CMP and HCPP appellants under part 417 of this chapter may combine part A and part B claims together to meet the requisite amounts in controversy for a hearing.

[59 FR 12181, Mar. 16, 1994]

§ 405.747 Dismissal of request for hearing; amount in controversy less than $100.

The ALJ shall, without holding a hearing, dismiss the request for hearing if the request for hearing plainly shows that less than $100 is in controversy. If a hearing is held and the ALJ finds that the amount in controversy is less than $100, the ALJ shall dismiss the request for hearing and will not rule on the substantive issues involved in the appeal.

§ 405.750 Time period for reopening initial, revised, or reconsidered determinations and decisions or revised decisions of an ALJ or the Departmental Appeals Board (DAB); binding effect of determination and decisions.

(a) Reopenings concerning applications and entitlement. A determination, or decision, or revised determination or decision made by the SSA concerning any matter under §405.704(a), may be reopened and revised under 20 CFR 404.988 (Conditions for reopening).

(b) Reopenings concerning a request for payment. An initial, revised, or reconsidered determination of HCFA, or a decision or revised decision of an ALJ or of the DAB, with respect to an individual’s right concerning a request for payment under Medicare Part A, which is otherwise binding under 20 CFR 404.955 or 404.961 and §§405.708 or 405.717 of this subpart may be reopened:

1. Within 12 months from the date of the notice of the initial or reconsidered determination to the party to such determination;
2. After such 12-month period, but within 4 years after the date of the notice of the initial determination to the individual, upon establishment of good cause for reopening such determination or decision (see 20 CFR 404.988(b) and 404.989); or
3. At any time, when:
   i. Such initial, revised, or reconsidered determination or such decision or revised decision is unfavorable, in whole or in part, to the party thereto, but only for the purpose of correcting clerical error or error on the face of the evidence on which such determination or decision was based; or
   ii. Such initial, revised, or reconsidered determination or such decision or revised decision was procured by fraud or similar fault of the beneficiary or some other person.

§ 405.753 Appeal of a categorization of a device.

(a) HCFA’s acceptance of the FDA categorization of a device as an experimental/investigational (Category A) device under §405.203 is a national coverage decision under section 1862(a)(1) of the Act.

(b) HCFA’s acceptance of the FDA categorization of a device as an experimental/investigational (Category A) device under §405.203 is an aspect of an initial determination that, under section 1862 of the Act, payment may not be made.

(c) In accordance with section 1869(b)(3)(A) of the Act, HCFA’s acceptance of the FDA categorization of a device as an experimental/investigational (Category A) device under §405.203 may not be reviewed by an administrative law judge.

[60 FR 48424, Sept. 19, 1995]

Subpart H—Appeals Under the Medicare Part B Program

AUTHORITY: Secs. 1102, 1842(b)(3)(C), 1869(b), and 1871 of the Social Security Act (42 U.S.C. 1302, 1395u(b)(3)(C), 1395f(b), and 1395hh).


§ 405.801 Part B appeals—general description.

(a) The Medicare carrier makes an initial determination when a request for payment for Part B benefits is submitted. If an individual beneficiary is dissatisfied with the initial determination, he or she may request, and the carrier will perform, a review of the claim. Following the carrier’s review determination, the beneficiary may obtain a carrier hearing if the amount remaining in controversy is at least $100. The beneficiary is also entitled to a carrier hearing without the benefit of a review determination when the initial request for payment is not being acted upon with reasonable promptness (as defined in §405.802). Following the carrier’s hearing, the beneficiary may obtain a carrier hearing if the amount remaining in controversy is at least $1,000. The beneficiary is also entitled to a carrier hearing without the benefit of a review determination when the initial request for payment is not being acted upon with reasonable promptness (as defined in §405.802). Following the carrier’s hearing, the beneficiary may obtain a carrier hearing if the amount remaining in controversy is at least $500. If the beneficiary is dissatisfied with the decision of the ALJ, he or she may request the Departmental Appeals Board (DAB) to review the case. Following the action of the DAB, the beneficiary may file suit in Federal district court if the amount remaining in controversy is at least $1,000.
§ 405.802 Definitions.

As used in subpart H of this part, the term—

After receipt of the notice means 5 days after the date of the notice, unless it is shown that the notice was received earlier or later.

Appellant designates the beneficiary, assignee or other person or entity that has filed an appeal concerning a particular determination of benefits under Medicare part B. Designation as an appellant does not in itself convey standing to appeal the determination in question.

Assignee means a physician or supplier who furnishes services to a beneficiary under Medicare part B and who has accepted a valid assignment executed by the beneficiary.

Assignment means the transfer by the assignor of his or her claim for payment to the assignee in return for the latter’s promise not to charge more for his or her services than the carrier finds to be the reasonable charge or other approved amount.

Assignor means a beneficiary under Medicare part B whose physician or supplier has taken assignment of a claim.

Carrier means an organization which has entered into a contract with the Secretary pursuant to section 1842 of the Act and which is authorized to make determinations with respect to part B of title XVIII of the Act. For purposes of this subpart, the term carrier also refers to an intermediary that has entered into a contract with the Secretary under section 1816 of the Act and is authorized to make determinations with respect to part B provider services, as specified in §421.5(c) of this chapter.

Common issues of law and fact, with respect to the aggregation of claims by two or more appellants to meet the minimum amount in controversy needed for an ALJ hearing, occurs when the claims sought to be aggregated are denied or reduced for similar reasons and arise from a similar fact pattern material to the reason the claims are denied.

Delivery of similar or related services, with respect to the aggregation of claims by two or more physician/supplier appellants to meet the minimum amount in controversy needed for an ALJ hearing, means like or coordinated services or items provided to the same beneficiary by the appellants.

Representative means an individual meeting the conditions described in §§ 405.870 through 405.871.

With reasonable promptness means within a period of 60 consecutive days after the receipt by the carrier of a request for payment.

§ 405.803 Initial determination.

(a) Carriers make initial determinations regarding claims for benefits under Medicare Part B.

(b) An initial determination for purposes of this subpart includes determinations such as the following:

1. Whether services furnished are covered.

2. Whether the deductible has been met.

3. Whether the receipted bill or other evidence of payment is acceptable.

4. Whether the charges for services furnished are reasonable.
(5) If the services furnished to a beneficiary by a physician or a supplier pursuant to an assignment under §424.55 of this chapter are not covered because they are determined to be not reasonable and necessary under §411.15(k) of this chapter, whether the beneficiary, physician or supplier, or a physician who meets the requirements of §411.408, knew or could reasonably have been expected to know at the time the services were furnished that the services were not covered.

(c) The following are not initial determinations for purposes of this subpart:

(1) Any issue or factor for which SSA or HCFA has sole responsibility, for example, whether an independent laboratory meets the conditions for coverage of services; whether a Medicare overpayment claim should be compromised, or collection action terminated or suspended.

(2) Any issue or factor which relates to hospital insurance benefits under Medicare Part A.


§ 405.804 Notice of initial determination.

After a carrier has made an initial determination on a request for payment written notice of this determination shall be mailed to each party to the determination at his last known address. The notice of the determination shall inform each party to the determination of his right to have such determination reviewed.

§ 405.805 Parties to the initial determination.

The parties to the initial determination (see §405.803) may be any party described in §405.802.

[64 FR 52670, Sept. 30, 1999]

§ 405.806 Effect of Initial Determination.

The initial determination is binding upon all parties to the claim for benefits unless the determination is—

(a) Reviewed in accordance with §§405.810 through 405.812; or

(b) Revised as a result of a reopening in accordance with §405.841.


§ 405.807 Request for review of initial determination.

(a) General. A party to an initial determination by a carrier, that is dissatisfied with the initial determination and wants to appeal the matter, may request that the carrier review the determination. The request for review by the party to an initial determination must clearly indicate that he or she is dissatisfied with the initial determination and wants to appeal the matter. The request for review does not constitute a waiver of the party's right to a hearing (under §405.815) after the review.

(b) Place and method of filing a request. A request by a party for a carrier to review the initial determination may be made in one of the following ways:

(1) In writing and filed at an office of the carrier, SSA, or HCFA.

(2) By telephone to the telephone number designated by the carrier as the appropriate number for the receipt of requests for review.

(c) Time of filing request. (1) The carrier must provide a period of 6 months after the date of the notice of the initial determination within which the party to the initial determination may request a review.

(2) The carrier may, upon request by the party, extend the period for requesting the review of the initial determination.

[64 FR 52670, Sept. 30, 1999]

§ 405.808 Parties to the review.

The parties to the review (as provided for in §405.807(a)) shall be the persons who were parties to the carrier's initial determination as described in §405.805, and any other party whose rights with respect to the particular claim being reviewed may be affected by such review.


§ 405.809 Opportunity to submit evidence.

The parties to the review (as provided for in §405.807(a)) shall have a
§ 405.810 Review determination.

Subject to the provisions of §§ 405.807 through 405.809, the carrier shall review the claim in dispute and, upon the basis of the evidence of record, shall make a separate determination affirming or revising inwhole or in part the findings and determination in question.


§ 405.811 Notice of review determination.

Written notice of the review determination is mailed to a party at his or her last known address. The review determination states the basis of the determination and advises the party of his or her right to a carrier hearing when the amount in controversy is $100 or more as determined in accordance with § 405.817. The notice states the place and manner of requesting a carrier hearing as well as the time limit under which a hearing must be requested (see § 405.821).


§ 405.812 Effect of review determination.

The review determination is binding upon all parties to the review unless a carrier hearing decision is issued pursuant to a request for hearing made in accordance with § 405.821 or is revised as a result of reopening in accordance with § 405.841.


§ 405.815 Amount in controversy for carrier hearing, ALJ hearing and judicial review.

Any party designated in § 405.822 is entitled to a carrier hearing after a review determination has been made by the carrier if the amount remaining in controversy is $100 or more and the party meets the requirements of § 405.821 of this subpart. To be entitled to a hearing before an ALJ following the carrier hearing, the amount remaining in controversy must be $500 or more, and for judicial review following the ALJ hearing and Departmental Appeals Board Review, the amount remaining in controversy must be $1000 or more.


§ 405.817 Principles for determining amount in controversy.

(a) Individual appellants. For the purpose of determining whether an individual appellant meets the minimum amount in controversy needed for a carrier hearing ($100) or ALJ hearing ($500), the following rules apply:

(1) The amount in controversy is computed as the actual amount charged the individual for the items and services in question, less any amount for which payment has been made by the carrier and less any deductible and coinsurance amounts applicable in the particular case.

(2) A single beneficiary may aggregate claims from two or more physicians/suppliers to meet the $100 or $500 thresholds. A single physician/supplier may aggregate claims from two or more beneficiaries to meet the $100 or $500 threshold levels of appeal.

(3) In either of the circumstances specified in paragraph (a)(2) of this section, two or more claims may be aggregated by an individual appellant to meet the amount in controversy for a carrier hearing only if the claims have previously been reviewed and a request for hearing has been made within six months after the date of the review determination(s).

(4) In either of the circumstances specified in paragraph (a)(2) of this section, two or more claims may be aggregated by an individual appellant to meet the amount in controversy for an ALJ hearing only if the claims have previously been decided by a carrier hearing officer and a request for an ALJ hearing has been made within 60 days after receipt of the carrier hearing officer decision(s).

(5) When requesting a carrier hearing or an ALJ hearing, the appellant must specify in his or her appeal request the specific claims to be aggregated.

(b) Two or more appellants. As specified in this paragraph, under section...
Health Care Financing Administration, HHS § 405.821

§ 405.821 Request for carrier hearing.

(a) A request for a carrier hearing is any clear expression in writing by a claimant asking for a hearing to adjudicate a claim when not acted upon with reasonable promptness or by a party to a review determination who states, in effect, that he or she is dissatisfied with the carrier's review determination and wants further opportunity to appeal the matter to the carrier.

(b) The hearing request must be filed at an office of the carrier or at an office of SSA or HCFA.

(c) Except when a carrier hearing is held because the carrier did not act upon a claim with reasonable promptness, a party to a review determination may request a carrier hearing within six months after the date of the notice of the review determination. The carrier may, upon request by the
§ 405.822 Parties to a carrier hearing.

The parties to a hearing shall be the persons who were parties to the carrier's review determination (§ 405.808) which is in question. Any other person may be made a party if that person's rights with respect to supplementary medical insurance benefits may be prejudiced by the decision.

§ 405.823 Carrier hearing officer.

Any hearing provided for in this subpart shall be conducted by a hearing officer designated by the appropriate official of the carrier.

§ 405.824 Disqualification of carrier hearing officer.

A hearing officer shall not conduct a hearing in any case in which he is prejudiced or partial with respect to any party, or if he has any interest in the matter before him. Notice of any objection with respect to the hearing officer who will conduct the hearing shall be made by the objecting party at his earliest opportunity. The hearing officer shall consider such objection and shall, at his discretion, withdraw. If the hearing officer withdraws, the appropriate official of the carrier shall designate another hearing officer to conduct the hearing. If the hearing officer does not withdraw, the objecting party may present his objections to the carrier for consideration at any time prior to the issuance of a decision. The carrier shall review the request and take appropriate action. The fact that a hearing officer is an employee of the carrier may not serve as prima facie cause for disqualification.
§ 405.832 Dismissal of request for carrier hearing.

(a) By application of party. With the approval of the hearing officer, a request for a hearing may be withdrawn or dismissed at any time prior to the mailing of notice of the decision on the application of the party or parties filing the request for such hearing. A party may request a dismissal by filing a written notice of such request with the carrier, the hearing officer or orally stating such request at the hearing. The dismissal of a request for hearing shall be binding unless vacated (see paragraph (d) of this section).

(b) Dismissal by abandonment of party. A hearing officer may dismiss a request for hearing upon abandonment by the party or parties who filed the request. A party shall be deemed to have abandoned a request for hearing, other than where personal appearance is waived in accordance with § 405.831, if neither the party nor his representative appears at the time and place fixed for the hearing and within 10 days after the mailing of a notice to him by the hearing officer to show cause, such party does not show good and sufficient cause for such failure to appear and failure to notify the hearing officer prior to the time fixed for hearing that he cannot appear.

(c) Dismissal for cause. The hearing officer may, on his own motion, dismiss a hearing request, either entirely or as to any stated issue, under either of the following circumstances:

(1) Where the party requesting a hearing is not a proper party under § 405.822 or does not otherwise have a right to a hearing under section 1842(b)(3)(C) of the Act; or
§ 405.833 Record of carrier hearing.

A complete record of the proceedings at the carrier hearing is made. The testimony is transcribed and copies of other documentary evidence are reproduced in any case when directed by the hearing officer, the carrier, or HCFA. The record will also be transcribed and reproduced at the request of any party to the hearing provided the requesting party bears the cost.


§ 405.834 Carrier hearing officer's decision.

(a) As soon as practicable after the close of a carrier hearing, the carrier hearing officer issues a decision in the case based upon the evidence presented at the hearing or otherwise included in the hearing record. The decision is issued as a written notice to the parties and contains—

(1) Findings of fact,

(2) A statement of reasons, and

(3) Notification to the parties of their right to an ALJ hearing when the amount remaining in controversy is at least $500.

(b) A copy of the decision is mailed to the parties to the hearing at their last known addresses.


§ 405.835 Effect of carrier hearing officer's decision.

The carrier hearing officer's decision is binding upon all parties to the hearing unless—

(a) A request for an ALJ hearing is filed in accordance with §405.855, or

(b) The decision is revised in accordance with §405.841.


§ 405.836 Authority of the carrier hearing officer.

The carrier hearing officer, in adjudicating Medicare Part B claims, complies with all of the provisions of, and regulations issued under, title XVIII of the Act, as well as with HCFA Rulings, national coverage decisions, and other policy statements, instructions, and guides issued by HCFA.

§ 405.842 Notice of reopening and revision.

(a) Notice. When any determination or decision is reopened as provided in § 405.841, notice of such reopening shall be mailed to the parties to such determination or decision at their last known addresses. A notice of revision following a reopening of a decision, shall be mailed to the parties and shall state the basis for the revised determination or decision.

(b) Effect of revised determination. The revision of a determination (see § 405.841) shall be binding upon all parties thereto unless a party files a written request for a hearing with respect to a revised determination when the amount in controversy is $100 or more.


§ 405.850 Change of ruling or legal precedent.

Change of a legal interpretation or administrative ruling upon which a determination or decision was made shall not be considered as good and sufficient reason for reopening the determination or decision.

§ 405.853 Expedited appeals process.

(a) Conditions for use of expedited appeals process (EAP). A party may use the EAP set forth in § 405.718 of this chapter to request court review in place of the ALJ hearing or Departmental Appeals Board (DAB) review if the following conditions are met:

1. The carrier hearing officer has made a decision; an ALJ has made a hearing decision; or DAB review has been requested, but a final decision has not been issued.

2. The filing entity is a party referred to in § 405.718(d) of this chapter.

3. The party has filed a request for an ALJ hearing in accordance with § 405.855, or DAB review in accordance with 20 CFR 404.968.

4. The amount remaining in controversy is $1,000 or more.

5. If there is more than one party to the hearing decision, each party concurs, in writing, with the request for an EAP.

(b) Content of the request for EAP. The request for an EAP:

1. Alleges that there are no material issues of fact in dispute; and

2. Asserts that the only factor precluding a decision favorable to the party is a statutory provision that is unconstitutional or a regulation, national coverage decision under section 1862(a)(1) of the Act, or HCFA Ruling that is invalid.


§ 405.855 ALJ hearing.

(a) Right to Hearing. A party to the carrier hearing has a right to a hearing before an ALJ if:

1. The party files a written request for an ALJ hearing within 60 days after receipt of the notice of the carrier hearing decision; and

2. The amount remaining in controversy is $500 or more.

(b) Place of filing hearing request. The request for an ALJ hearing must be made in writing and filed with the carrier that issued the decision, a Social Security office, or, in the case of a qualified railroad retirement beneficiary, an office of the Railroad Retirement Board.

(c) Effect of ALJ hearing decision. (1) An ALJ’s decision is binding on all parties to the hearing unless—

i. The DAB reviews the ALJ decision;

ii. The DAB does not review the ALJ decision, and the party requests judicial review;

iii. The decision is revised by the DAB or an ALJ in accordance with the provisions of § 405.750 of this chapter; or

iv. The expedited appeals process is used.


§ 405.856 Departmental Appeals Board (DAB) review.

Regulations beginning at 20 CFR 404.967 regarding SSA Appeals Council Review are applicable to DAB review of matters addressed by this subpart.


§ 405.857 Court review.

(a) General rule. To the extent authorized by sections 1869, 1876(c)(5)(B), and 1879(d) of the Act, a party to a DAB decision, or an ALJ decision if the DAB does not review the ALJ’s decision,
may obtain a court review if the amount remaining in controversy is $1,000 or more. A party may obtain court review by filing a civil action in a district court of the United States in accordance with the provisions of section 205(g) of the Act. The filing procedure is set forth in 20 CFR 422.210.

(b) Prohibition against court review of certain Part B regulations or instructions. Under section 1869(b)(4) of the Act, a court may not review a regulation or instruction that relates to a method of payment under Part B if the regulation was promulgated, or the instruction issued, before January 1, 1981. [62 FR 25854, May 12, 1997]

§ 405.860 Review of national coverage decisions (NCDs).

(a) General. (1) HCFA makes NCDs either granting, limiting, or excluding Medicare coverage for a specific medical service, procedure or device. NCDs are made under section 1862(a)(1) of the Act or other applicable provisions of the Act. An NCD is binding on all Medicare carriers, fiscal intermediaries, PROs, HMOs, CMPs, and HCPPs when published in HCFA program manuals or the FEDERAL REGISTER.

(2) Under section 1869(b)(3) of the Act, only NCDs made under section 1862(a)(1) of the Act are subject to the conditions of paragraphs (b) through (d) of this section.

(b) Review by ALJ. (1) An ALJ may not disregard, set aside, or otherwise review an NCD.

(2) An ALJ may review the facts of a particular case to determine whether an NCD applies to a specific claim for benefits and, if so, whether the NCD has been applied correctly to the claim.

(c) Review by Court. (1) A court’s review of an NCD is limited to whether the record is incomplete or otherwise lacks adequate information to support the validity of the decision, unless the case has been remanded to the Secretary to supplement the record regarding the NCD. The court may not invalidate an NCD except upon review of the supplemented record.

(2) A Federal court may not hold unlawful or set aside an NCD because it was not issued in accordance with the notice and comment procedures of the Administrative Procedure Act (5 U.S.C. 553) or section 1871(b) of the Act.

(d) Remands—(1) Secretary’s action. When a court remands an NCD matter to the Secretary because the record in support of the NCD is incomplete or otherwise lacks adequate information, the Secretary remands the case to HCFA in order to supplement the record.

(2) Remand to HCFA. HCFA supplements the record with new or updated evidence, including additional information from other sources, and may issue a revised NCD.

(3) Final Actions. (i) The proceedings to supplement the record, are expedited.

(ii) When HCFA does not issue a revised NCD, it returns the supplemented record to the court for review.

(iii) When HCFA issues a revised NCD, it forwards the case to an ALJ who issues a new decision applying the revised NCD to the facts of the claim(s) under consideration. The ALJ’s decision is subject to DAB review and, ultimately, judicial review. [62 FR 25854, May 12, 1997]

§ 405.870 Appointment of representative.

A party to an initial determination, informal review or hearing as provided in §§ 405.803 through 405.934, may appoint as his representative in any such proceeding any person qualified under §§ 405.803 through 405.934, may appoint as his representative in any such proceeding any person qualified under §§ 405.871. Where the representative is an attorney, in the absence of information to the contrary, his representation that he has such authority shall be accepted as evidence of the attorney’s authority to represent a party.

§ 405.871 Qualifications of representatives.

Any individual may be appointed to act as representative in accordance with §405.870, unless he is disqualified or suspended from acting as a representative in proceedings before the SSA or the HCFA or unless otherwise prohibited by law. [39 FR 12098, Apr. 3, 1974. Redesignated at 42 FR 52926, Sept. 30, 1977, as amended at 62 FR 25854, May 12, 1997]
§ 405.872 Authority of representatives.

A representative, appointed and qualified as provided in §§ 405.870 and 405.871, may make or give, on behalf of the party he represents, any request or notice relative to any proceeding before the carrier including review and hearing. A representative shall be entitled to present evidence and allegations as to facts and law in any proceeding affecting the party he represents and to obtain information with respect to the claim of such party to the same extent as such party. Notice to any party in any action, determination, or request to any party for the production of evidence, shall be sent to the representative of such party.

§ 405.874 Appeals of carrier decisions that supplier standards are not met.

(a) An entity serving as a National Supplier Clearinghouse must act promptly to determine if any entity submitting a request for a billing number as a Medicare supplier of part B items meets the standards set forth in part 424. Effective July 1, 1993, the National Supplier Clearinghouse must accept, reject or request additional information within 15 days of the receipt of an enrollment application.

(b) If the National Supplier Clearinghouse disallows an entity’s request for a billing number or revokes, with the concurrence of HCFA, an entity’s billing number, the National Supplier Clearinghouse notifies the entity by certified mail. Revocation is effective 15 days after the National Supplier Clearinghouse mails notice of its determination. The carrier disallows payment for items furnished by the supplier beginning with that effective date. The notice must inform the entity of the reason for the rejection or revocation, its right to appeal, the date by which it must file that appeal (90 days after the postmark of the notice) and the address to which the appeals must be sent in writing.

(c) A fair hearing officer not involved in the original determination to disallow an entity’s request for a billing number, or to revoke an entity’s billing number, must schedule a hearing to be held within one week of receipt of an appeal, or later at the request of the entity. Both the entity and carrier may offer evidence. The hearing officer issues notice of his/her decision within 2 weeks of the hearing. The notice is sent by certified letter to HCFA, the carrier, and the appealing entity. This notice must include information about the supplier’s further right to appeal, the carrier’s right to appeal, the date by which the appeal must be filed (90 days after the postmark of the notice) and the address to which the appeals must be sent in writing. Either the carrier or entity may appeal the hearing officer’s decision to HCFA.

(d) A HCFA official, designated by the Administrator of HCFA, must make an appeal decision based on the evidence presented to the fair hearing officer and his or her decision. The HCFA official requests any additional information he or she deems necessary from either the carrier or the entity within two weeks of receipt by the HCFA of the appeal. Notice of the HCFA official’s decision contains information about any further appeals the entity and carrier may have.

(e) A billing number is not issued, or remains revoked, and payment is not made, for items or services furnished by any entity which a carrier determines does not qualify for a billing number, until the carrier (upon re-application of the entity), a fair hearing officer, or a HCFA official designated to hear such appeals, determines that the entity qualifies for a billing number. Any claims for items or services furnished after revocation of the supplier’s billing number and submitted by the entity during the appeals period are held and not processed, i.e., are neither approved, denied or developed, until all administrative appeals have been exhausted. If an entity is determined not to have qualified for a billing number in one period but to
have qualified in another, the carrier pays for claims for items sold or rented to beneficiaries during the period the entity qualified as a supplier. If there is evidence of an overpayment, see subpart C of part 405 of this Chapter.

(f) A billing number may be reinstated after revocation when an entity completes a corrective action plan, to which HCFA has agreed, and provided sufficient assurance of its intent to comply fully with the supplier standards.

[57 FR 27305, June 18, 1992]

§ 405.877 Appeal of a categorization of a device.

(a) HCFA's acceptance of the FDA categorization of a device as an experimental/investigational (Category A) device under § 405.203 is a national coverage decision under section 1862(a)(1) of the Act.

(b) HCFA's acceptance of the FDA categorization of a device as an experimental/investigational (Category A) device under § 405.203 is an aspect of an initial determination that, under section 1862 of the Act, payment may not be made.

(c) In accordance with section 1869(b)(3)(A) of the Act, HCFA's acceptance of the FDA categorization of a device as an experimental/investigational (Category A) device under § 405.203 may not be reviewed by an administrative law judge.

[60 FR 48424, Sept. 19, 1995]

Subparts I–Q—[Reserved]

Subpart R—Provider Reimbursement Determinations and Appeals

Authority: Secs. 205, 1102, 1814(b), 1815(a), 1833, 1833(v), 1871, 1872, 1878, and 1886 of the Social Security Act (42 U.S.C. 405, 1302, 1395f(b), 1395f(a), 1395l, 1395x(v), 1395hh, 1395i, 1395oo, and 1395ww).


§ 405.1801 Introduction.

(a) Definitions. As used in this subpart:

Administrator means the Administrator or Deputy Administrator of HCFA.

Administrator’s review means that review provided for in section 1878(f) of the Act (42 U.S.C. 1395oo(f)) and § 405.1875.

Board means the Provider Reimbursement Review Board established in accordance with section 1878 of the Act (42 U.S.C. 1395oo) and § 405.1845.

Board hearing means that hearing provided for in section 1878(a) of the Act (42 U.S.C. 1395oo(a)), and § 405.1835.

Date of filing and date of submission of materials mean the day of the mailing (as evidenced by the postmark) or hand-delivery of materials, unless otherwise defined in this subpart.

Date of receipt means the date on the return receipt of “return receipt requested” mail, unless otherwise defined in this subpart.

Intermediary determination means the following:

(1) With respect to a provider of services that has filed a cost report under §§ 413.20 and 413.24(f) of this chapter, the term means a determination of the total amount of payment due the provider, pursuant to § 405.1803 following the close of the provider’s cost reporting period, for items and services furnished to beneficiaries for which reimbursement may be made on a reasonable cost basis under Medicare for the period covered by the cost report.

(2) With respect to a hospital that receives payments for inpatient hospital services under the prospective payment system (part 412 of this chapter), the term means a determination of the total amount of payment due the hospital, pursuant to § 405.1803 following the close of the hospital’s cost reporting period, under that system for the period covered by the determination.

(3) For purposes of appeal to the Provider Reimbursement Review Board, the term is synonymous with the phrases “intermediary’s final determination” and “final determination of the Secretary”, as those phrases are used in section 1878(a) of the Act.

(4) For purposes of § 405.376 concerning claims collection activities, the term does not include an action by HCFA with respect to a compromise of
§ 405.1803 Intermediary determination and notice of amount of program reimbursement.

(a) General requirement. Upon receipt of a provider's cost report, or amended cost report where permitted or required, the intermediary must, within a reasonable period of time (see § 405.1835(b)), furnish the provider and other parties as appropriate (see § 405.1805) a written notice reflecting the intermediary's determination of the total amount of reimbursement due the provider for the cost reporting period covered by the notice. The intermediary must include in the notice the maximum extent possible, the amounts determined to be due, and the amounts received by the hospital during the cost reporting period covered by the notice.

(b) Requirements for intermediary notices. The intermediary must include in each notice appropriate references to law, regulations, HCFA Rulings, or
§ 405.1804

Program instructions to explain why the intermediary's determination of the amount of program reimbursement for the period differs from the amount the provider claimed. The notice must also inform the provider of its right to an intermediary or Board hearing (see §§ 405.1809, 405.1811, 405.1815, 405.1835, and 405.1843) and that the provider must request the hearing within 180 days after the date of the notice.

(c) Use of notice as basis for recoupment of overpayments. The intermediary's determination contained in its notice is the basis for making the retroactive adjustment (required by § 413.64(f) of this chapter) to any program payments made to the provider during the period to which the determination applies, including recoupment under § 405.373 from ongoing payments to the provider of any overpayments to the provider identified in the determination. Recoupment is made notwithstanding any request for hearing on the determination the provider may make under § 405.1811 or § 405.1835.


§ 405.1807

Effect of intermediary determination.

The determination shall be final and binding on the party or parties to such determination unless:

(a) an intermediary hearing is requested in accordance with § 405.1811 and an intermediary hearing decision rendered in accordance with § 405.1831; or

(b) the intermediary determination is revised in accordance with § 405.1885; or

(c) a Board hearing is requested in accordance with § 405.1835 and a hearing decision rendered pursuant thereto.

§ 405.1809

Intermediary hearing procedures.

(a) Hearings. Each intermediary must establish and maintain written procedures for intermediary hearings, in accordance with the regulations in this subpart, for resolving issues that may arise between the intermediary and a provider concerning the amount of reasonable cost reimbursement, or prospective payment due the provider (except as provided in § 405.1804) under the Medicare program. The procedures must provide for a hearing on the intermediary determination contained in the notice of program reimbursement (§ 405.1803), if the provider files a timely request for a hearing.

(b) Amount in controversy. In order for an intermediary to grant a hearing, the following dates and amounts in controversy apply:

(1) For cost reporting periods ending prior to June 30, 1973, the amount of program reimbursement in controversy must be at least $1000.

(2) For cost reporting periods ending on or after June 30, 1973, the amount of program reimbursement in controversy must be at least $1000 but less than $10,000.

§ 405.1811 Right to intermediary hearing; time, place, form, and content of request for intermediary hearing.

(a) A provider that has been furnished a notice of amount of program reimbursement may request an intermediary hearing if it is dissatisfied with the intermediary's determination contained in the notice and the amount in controversy requirement described in § 405.1809 is met. The request must be in writing and be filed with the intermediary within 180 calendar days after the date of the notice. (See § 405.1835(c)). No other individual, entity, or party has the right to an intermediary hearing.

(b) The request must (1) identify the aspect(s) of the determination with which the provider is dissatisfied, and (2) explain why the provider believes the determination on these matters is incorrect, and (3) be submitted with any documentary evidence the provider considers necessary to support its position.

(c) Following the timely filing of the request for hearing, the provider may identify in writing, prior to the onset of the hearing proceedings, additional aspects of the determination with which it is dissatisfied and furnish any documentary evidence in support thereof. If such additional aspects are submitted, the hearing officer may postpone the hearing to allow for his examination of such additional aspects.


§ 405.1813 Failure to timely request an intermediary hearing.

If a provider requests an intermediary hearing on an intermediary's determination after the time limit prescribed in § 405.1811, the designated intermediary hearing officer or panel of hearing officers will dismiss the request and furnish the provider a written notice that explains the time limitation, except that for good cause shown, the time limit prescribed in § 405.1811 may be extended. However, an extension may not be granted if the extension request is filed more than 3 years after the date of the original notice of the intermediary determination.

[48 FR 39835, Sept. 1, 1983]

§ 405.1815 Parties to the intermediary hearing.

The parties to the intermediary hearing shall be the parties to the intermediary determination and any other entity determined by the intermediary to be a related organization of such provider. Said parties shall be given reasonable notice of the time, date, and place of such hearing. Neither the intermediary nor the Health Care Financing Administration are parties (see § 405.1819).

§ 405.1817 Hearing officer or panel of hearing officers authorized to conduct intermediary hearing; disqualification of officers.

The intermediary hearing provided for in § 405.1809 shall be conducted by a hearing officer or panel of hearing officers designated by the intermediary. Such hearing officer or officers shall be persons knowledgeable in the field of health care reimbursement. The hearing officer or officers shall not have had any direct responsibility for the program reimbursement determination with respect to which a request for hearing is filed; no hearing officer (or officers) shall conduct a hearing in a case in which he is prejudiced or partial with respect to any party, or where he has any interest in the matter pending for determination before him. Notice of any objection which a party may have with respect to a hearing officer shall be presented in writing to such officer by the objecting party at the party's earliest opportunity. The hearing officer shall consider the objection and shall, at his discretion, either proceed in the conduct of the hearing or withdraw. If the hearing officer does not withdraw, the objecting party may, after the hearing, present his objections to an executive official of the intermediary, who shall rule promptly on the objection.

§ 405.1819 Conduct of intermediary hearing.

The hearing shall be open to all parties thereto (see § 405.1815) and to representatives of the intermediary and of
§ 405.1821 Prehearing discovery and other proceedings prior to the intermediary hearing.

(a) Prehearing discovery shall be permitted upon timely request of any party. To be timely, a request for discovery and inspection shall be made before the beginning of the hearing. A reasonable time for inspection and reproduction of documents shall be provided by order of the hearing officer(s).

(b) If, in the discretion of the hearing officer(s), the purpose of defining the issues more clearly would be served, the hearing officer(s) may schedule a prehearing conference. For this purpose, a single member of a panel of hearing officers, when such is the case, may be appointed to act for the panel with respect to prehearing activities.

§ 405.1823 Evidence at intermediary hearing.

Evidence may be received at the intermediary hearing even though inadmissible under the rules of evidence applicable to court procedure. The hearing officer(s) shall give the parties opportunity for submission and consideration of facts and arguments, and during the course of the hearing, should in ruling upon admissibility of evidence, exclude irrelevant, immaterial, or unduly repetitious evidence. The hearing officer(s) shall render a final ruling on the admissibility of evidence.

§ 405.1825 Witnesses at intermediary hearing.

The hearing officer(s) may examine the witnesses and shall allow the parties and their representatives to do so. Parties to the proceedings may also cross-examine witnesses.

§ 405.1827 Record of intermediary hearing.

A complete recordation of the proceedings at the intermediary hearing shall be made and transcribed in all cases. It shall be made available to any party upon request. The record will not be closed until a decision (see § 405.1831) has been issued.

§ 405.1829 Authority of hearing officer(s) at intermediary hearing.

(a) The hearing officer(s) in exercising his authority must comply with all the provisions of title XVIII of the Act and regulations issued thereunder, as well as with HCFA Rulings issued under the authority of the Administrator of the Health Care Financing Administration (see 42 CFR 401.108), and with the general instructions issued by the Health Care Financing Administration in accordance with the Secretary's agreement with the intermediary.

(b) The determination of a fiscal intermediary that no payment may be made under title XVIII of the Act for any expense incurred for items and services furnished to an individual because such items and services are excluded from coverage pursuant to section 1862 of the Act, 42 U.S.C. 1395y (see subpart C of this part), shall not be reviewed by the hearing officer(s). Such determination shall be reviewed only in accordance with the applicable provisions of subparts G and H of this part.

§ 405.1831 Intermediary hearing decision and notice.

The hearing officer(s) shall, on a timely basis, render a decision in writing based on the evidence in the record; such decision shall constitute the final determination of the intermediary. In such decision, he will cite applicable law, regulations, HCFA Rulings, and general instructions of the Health Care Financing Administration, as well as
findings on all the matters in issue at the hearing. A copy of the decision will be mailed to all parties to the hearing at their last known addresses.

§ 405.1833 Effect of intermediary hearing decision.

The intermediary hearing decision provided for in § 405.1831 shall be final and binding upon all parties to the hearing unless such intermediary determination is revised in accordance with § 405.1885.

§ 405.1835 Right to Board hearing.

(a) Criteria. The provider (but no other individual, entity, or party) has a right to a hearing before the Board about any matter designated in § 405.1801(a)(1), if:

(1) An intermediary determination has been made with respect to the provider; and

(2) The provider has filed a written request for a hearing before the Board under the provisions described in § 405.1841(a)(1); and

(3) The amount in controversy (as determined in § 405.1839(a)) is $10,000 or more.

(b) Prospective payment exceptions. Except with respect to matters for which administrative or judicial review is not permitted as specified in § 405.1804, hospitals that are paid under the prospective payment system are entitled to hearings before the Board under this section if they otherwise meet the criteria described in paragraph (a) of this section.

(c) Right to hearing based on late intermediary determination about reasonable cost. Notwithstanding the provisions of paragraph (a)(1) of this section, the provider also has a right to a hearing before the Board if an intermediary’s determination concerning the amount of reasonable cost reimbursement due a provider is not rendered within 12 months after receipt by the intermediary of a provider’s perfected cost report or amended cost report (as permitted or as required to furnish sufficient data for purposes of making such determination—see § 405.1803(a)) provided such delay was not occasioned by the fault of the provider.

§ 405.1837 Group appeal.

(a) Criteria for group appeals. Subject to paragraph (b) of this section, a group of providers may bring an appeal before the Board but only if—

(1) Each provider in the group is identified as one which would, upon the filing of a request for a hearing before the Board, but without regard to the $10,000 amount in controversy requirement, be entitled to a hearing under § 405.1835;

(2) The matters at issue involve a common question of fact or of interpretation of law, regulations or HCFA Rulings; and

(3) The amount in controversy is, in the aggregate, $50,000 or more.

(b) Providers under common ownership or control. Effective April 20, 1983, any appeal filed by providers that are under common ownership or control must be brought by the providers as a group appeal in accordance with the provisions of paragraph (a) of this section with respect to any matters involving an issue common to the providers and for which the amount in controversy is, in the aggregate, $50,000 or more (see § 405.1841(a)(2)). A single provider involved in a group appeal that also wishes to appeal issues that are not common to the other providers in the group must file a separate hearing request (see § 405.1841(a)(1)) and must separately meet the requirements in § 405.1811 or § 405.1835, as applicable.

§ 405.1839 Amount in controversy.

(a) Single appeals. The $1,000 amount in controversy required under § 405.1809 for an intermediary hearing and the $10,000 amount in controversy required under § 405.1835 for a Board hearing is, as applicable to the matters for which the provider has requested a hearing, the combined total of the amounts computed as follows:

(1) Providers under prospective payment. For providers that are paid under the prospective payment system, by deducting—

(i) The total of the payment due the provider on other than a reasonable cost basis under the prospective payment system from the total amount
that would be payable after a recomputation that takes into account any exclusion, exception, adjustment, or additional payment denied the provider under part 412 of this chapter, as applicable;

(ii) The total of the payment due the provider on a reasonable cost basis under the prospective payment system from the total reimbursable costs claimed by the provider; and

(iii) The adjusted total reimbursable costs due the provider on a reasonable cost basis under other than the prospective payment system from the total reimbursable costs claimed in the aggregate by the providers.

(2) Providers not under prospective payment. For providers that are not paid under the prospective payment system, by deducting the adjusted total reimbursable program costs due the providers (in the aggregate) on a reasonable cost basis from the total reimbursable costs claimed in the aggregate by the providers.

[49 FR 323, Jan. 3, 1984]

§ 405.1841 Time, place, form, and content of request for Board hearing.

(a) General requirements. (1) The request for a Board hearing must be filed in writing with the Board within 180 days of the date the notice of the intermediary’s determination was mailed to the provider or, where notice of the determination was not timely rendered, within 180 days after the expiration of the period specified in § 405.1835(c). Such request for Board hearing must identify the aspects of the determination with which the provider is dissatisfied, explain why the provider believes the determination is incorrect in such particulars, and be accompanied by any documenting evidence the provider considers necessary to support its position. Prior to the commencement of the hearing proceedings, the provider may identify in writing additional aspects of the intermediary’s determination with which it is dissatisfied and furnish any documentary evidence in support thereof.

(2) Effective April 20, 1983, any request for a Board hearing by providers that are under common ownership or control (see § 413.17 of this chapter) must be brought by the providers as a group appeal (see § 405.1837(b)) with respect to any matters at issue involving a question of fact or of interpretation of law, regulations, or HCFA Rulings common to the providers and for which the amount in controversy is $50,000 or more in the aggregate. If a group appeal is filed, the provider seeking the appeal must be separately identified in the request for hearing, which must be prepared and filed consistently with the requirements of paragraph (a)(1) of this section.
(b) Extension of time limit for good cause. A request for a Board hearing filed after the time limit prescribed in paragraph (a) of this section shall be dismissed by the Board, except that for good cause shown, the time limit may be extended. However, no such extension shall be granted by the Board if such request is filed more than 3 years after the date the notice of the intermediary's determination is mailed to the provider.


§ 405.1842 Expediting Board proceedings.

(a) Basis and purpose. This section implements section 1878(f)(1) of the Social Security Act, as amended by section 955 of Public Law 96-499 (42 U.S.C. 1395oo(f)(1)). The amendment provides an opportunity for providers to obtain expedited administrative review when the Board determines that it does not have the authority to decide a question of law, regulation, or HCFA Ruling relevant to the case (see § 405.1867).

(b) Basic rule. (1) Except as provided in paragraph (b)(4) of this section, a provider may submit a written request to the Board, with supporting documentation, to determine whether the Board has the authority to decide a question of law, regulations, or HCFA Rulings relevant to and controlling upon an issue to be reviewed by the Board. The Board is required to make an expedited review determination in writing, either denying or granting the request, within 30 days after the date of receipt of the request, as defined in paragraph (1) of this section. The Board may also issue a determination on its own motion that it lacks the authority to decide a question of law, regulations, or HCFA Rulings relevant to and controlling upon an issue to be reviewed by the Board. The provider is required to make an expedited review determination in writing, either denying or granting the request, within 30 days after the date of receipt of the request, as defined in paragraph (1) of this section. The provider may also issue a determination on its own motion that it lacks authority to decide a question of law, regulations, or HCFA Rulings.

(2) The Board must determine that the provider (including each provider in a group appeal) is entitled to a hearing under section 1878(a) of the Act before making the determination described in paragraph (b)(1) of this section. Thus, the provider must file (or have already filed) a written request for a Board hearing that meets the requirements in § 405.1841. The information and documentation required with respect to the filing of a request for a hearing is used by the Board to determine jurisdiction under section 1878(a) of the Act.

(3) A provider's request for an expedited review determination cannot be considered to be filed with the Board, nor can the 30-day time period during which the Board is required to make an expedited review determination begin, until such time as the Board accepts jurisdiction of the case.

(4) Proceedings conducted by the Board under an authority other than section 1878(a) of the Act and §§ 405.1835 through 405.1873 of this subpart are not hearings for purposes of this section and are not subject to the expedited Board proceedings set forth in this section. For example, proceedings concerning reimbursement for capital expenditures conducted under section 1122(f) of the Act and §§ 405.1891 through 405.1873 of this subpart are not hearings for purposes of this section. (Section 1122(f) specifically bars any administrative or judicial review.)

(c) "Own motion" review. If the Board is considering issuing a determination on its own motion that it lacks the authority to decide a question of law, regulations, or HCFA Rulings, it will notify the provider and intermediary of its proposed determination and allow them a reasonable period of time to file evidence or arguments either to support or oppose the proposed determination.

(d) Provider requests. (1) If a provider seeks an expedited Board proceeding, it must—(i) File its appropriately documented request in writing with the Board; and

(ii) Send a copy of the request and documentation simultaneously to the intermediary.

(2) The request to the Board for an expedited review determination must—(i) Identify the issues and the controlling law, regulation, or HCFA Ruling for which the Board is to make a determination;

(ii) Allege and demonstrate that there are no factual issues in dispute;

(iii)含an an explanation of why the provider believes the Board cannot decide the legal issue or issues that are in dispute; and

(iv) Include all other information or details that support the request.
(3) If the information in the provider request is insufficient for the Board to determine whether it has the authority to decide an issue, the Board will request more information from the provider. Such a request will affect the 30-day time limit as provided in paragraph (i) of this section. If the provider does not send more information or sends inadequate information, the Board will determine that it has the authority to decide the issue and will begin the regular procedure for a hearing.

(e) Intermediary participation. (1) After receiving a copy of the provider's request for an expedited review determination, the intermediary may send comments to the Board on the provider's request and supporting documentation. The intermediary will send a copy of its comments to the provider simultaneously.

(2) If the intermediary's comments raise questions about the provider's request for expedited review, the Board may request additional information from the provider as provided in paragraph (d)(3) of this section.

(f) Criteria for a Board determination. The Board will review all documentation forwarded by the provider and the intermediary relevant to the request for a Board determination concerning the Board's authority to decide an issue. In its review, the Board will consider—

(1) The controlling facts in the case;
(2) The applicability of law, regulations, or HCFA Rulings;
(3) Whether there are factual issues for the Board to resolve; and
(4) Whether there are legal issues within the authority of the Board to decide.

(g) Board determination. (1) Within 30 days after the date of receipt (as defined in paragraph (i) of this section) of a provider's request and all necessary documentation the Board will issue a determination concerning its authority to decide the question of law, regulations, or HCFA Rulings relevant to the issues identified by the provider in its request.

(2) If there are factual or legal issues in dispute on an issue within the authority of the Board to decide, the Board will not make an expedited review determination on the particular issue but will proceed with a hearing. The Board has the authority to decide when two or more issues are sufficiently related to preclude separation for purposes of an expedited review determination on one or more of them and a hearing on the other or others.

(3) The Board will promptly notify the provider in writing of its determination and will send a copy of the determination to the intermediary.

(4) The Board's determination concerning its authority or its lack of a determination is not subject to the Secretary's review under §405.1875.

(h) Effect of a Board decision. (1) The Board's determination, issued on its own motion or at the request of a provider, that it lacks authority to decide a question of law, regulations or HCFA Rulings is a final decision permitting a provider to seek judicial review with respect to the matter or matters in controversy contained in the determination, within 60 days of the date of the Board's determination.

(2) After the Board has determined that it does not have the authority to decide an issue, the provider will not be granted a hearing on the same issue.

(3) If the Board fails to issue an expedited review determination within 30 days of the date of receipt of a complete request (as determined under paragraph (i) of this section), the provider may, within 60 days from the end of that period, seek judicial review of the matters for which it requested the Board's determination.

(4) If the Board fails to make an expedited review determination within the required 30 days, it will begin regular hearing procedures as though it has the authority to decide the issue.

(5) If the provider seeks judicial review because the Board fails to make a determination as provided in paragraph (g)(1) of this section, it should notify the Board at the time it files for judicial review. The Board will not hold a hearing, even if one has been scheduled, on the matter or matters for which the provider is seeking judicial review.

(6) The Board's determination does not affect the right of the provider to a Board hearing for issues for which the
provider did not request expedited review, or for which the Board determines it does have the authority to decide, or for which the Board did not make a determination and the provider did not request judicial review.

(i) Date of receipt. For purposes of this section, the date of receipt of the provider's request is the later of—

(1) The actual date of receipt by the Board of the information required under paragraph (d)(2) of this section, or of additional information requested by the Board under paragraph (d)(3) of this section, whichever the Board receives later; or

(2) The date indicated on the Board's written notification to the provider that the Board has accepted jurisdiction of the case.

(j) Examples. Below are examples showing when a provider may expect to receive an expedited review determination, in relation to various circumstances affecting its request for the determination.

(1) The provider requests a hearing and expedited review at or about the same time. If all information is complete, the Board could send notification that it has accepted jurisdiction of the case and the expedited review determination simultaneously.

(2) The provider requests both a hearing and an expedited review determination, and supplies complete information. The Board accepts jurisdiction but, for example, because of the complexity of the case, the Board makes its expedited review determination within 30 days after it has accepted jurisdiction.

(3) The provider requests both a hearing and an expedited review determination, but the request for a hearing does not contain enough information for the Board to determine jurisdiction. The Board would request more information to determine jurisdiction and would make its expedited review determination within 30 days after it has accepted jurisdiction.

(4) The provider requests both a hearing and an expedited review determination, but does not send enough information for the Board to make an expedited review determination. Assuming the Board accepts jurisdiction, the Board would request more information about the request for expedited review and make its determination within 30 days after it receives the additional information.

(5) The provider requests an expedited review determination after the Board has accepted jurisdiction. The Board would make its determination within 30 days after receipt of an appropriately documented request for an expedited review determination.

§ 405.1843 Parties to Board hearing.

(a) The parties to the Board hearing shall be the provider, the intermediary (including the Health Care Financing Administration when acting directly as intermediary) that rendered the determination being appealed (see §405.1833), and any other entity found by the intermediary to be a related organization of such provider.

(b) Except as provided in paragraph (a), neither the Secretary nor the Health Care Financing Administration may be made a party to the hearing. However, the Board may call as a witness any employee or officer of the Department of Health and Human Services having personal knowledge of the facts and the issues in controversy in a hearing pending before the Board and may call as a consultant to the Board in connection with any such hearing any individual designated by the Secretary for such purpose. (See §405.1863.)

§ 405.1845 Composition of Board.

(a) The Board will consist of five members appointed by the Secretary. All shall be knowledgeable in the field of cost reimbursement. At least one shall be a certified public accountant. Two Board members shall be representative of providers of services.

(b) The term of office for Board members shall be 3 years, except that initial appointments may be for such shorter terms as the Secretary may designate to permit staggered terms of office. No member shall serve more than two consecutive 3-year terms of office. The Secretary shall have the authority to terminate a Board member's term of office for good cause.
§ 405.1847 Disqualification of Board members.

No Board member shall join in the conduct of a hearing in a case in which he is prejudiced or partial with respect to any party or in which he has any interest in the matter pending for decision before him. Notice of any objection which a party may have with respect to a Board member shall be presented in writing to such Board member by the objecting party at its earliest opportunity. The Board member shall consider the objection and shall, in his discretion, either proceed to join in the conduct of the hearing or withdraw. If he does not withdraw, the objecting party may petition the Board, presenting its objection and reasons therefor, and be entitled to a ruling thereon before the hearing can proceed.

§ 405.1849 Establishment of time and place of hearing by the Board.

The Board shall fix the time and place for the hearing and shall mail written notice thereof to the parties at their last known addresses, not less than 30 days prior to the scheduled time. Either on its own motion or for good cause shown by a party, the Board may, as appropriate, reschedule, adjourn, postpone, or reopen the hearing, provided that reasonable written notice is given to the parties.

§ 405.1851 Conduct of Board hearing.

The Board hearing shall be open to the parties, to representatives of the Health Care Financing Administration, and to such other persons as the Board deems necessary and proper. The Board shall inquire fully into all of the matters at issue and shall receive into evidence the testimony of witnesses and any documents which are relevant and material to such matters. If the Board believes that there is relevant and material evidence available which has not been presented at the hearing, it may at any time prior to the mailing of notice of the decision, reconvene the hearing for the receipt of such evidence. The order in which the evidence and the allegations shall be presented and the conduct of the hearing shall be at the discretion of the Board.

§ 405.1853 Prehearing discovery and other proceedings prior to the Board hearing.

(a) Upon notification that a request for Board hearing has been filed, the intermediary shall forthwith review the materials submitted by the provider in accordance with § 405.1841. Simultaneously, the intermediary shall review the information which formed the basis for its determination of the amount of program reimbursement. Based on the findings of such review, the intermediary shall expeditiously attempt to join with the provider in written stipulations setting forth the issues that said review has resolved and designating the issues that remain for Board resolution. Having obtained such stipulations and being satisfied that no further agreements can be negotiated, the intermediary shall ensure that all available documentary evidence in support of each party's position is part of the record. Such evidence will ordinarily include a position paper from the provider, a position paper from the intermediary, and any documents which support the issues addressed in the stipulations. These materials, in addition to all relevant documents which formed the basis for its determination of the amount of program reimbursement, shall be forwarded to the Board within 60 days after the date of the provider's request for Board review.
Health Care Financing Administration, HHS § 405.1867

(b) Prehearing discovery shall be permitted upon timely request of a party. To be timely, a request for discovery and inspection shall be made before the beginning of the hearing. A reasonable time for inspection and reproduction of documents shall be provided by order of the Board. The Board’s order on all discovery matters shall be final.

(c) If, in the discretion of the Board, the purpose of defining the issues more clearly would be served, the Board may schedule a prehearing conference. For this purpose, a single member of the Board may be appointed to act for the Board with respect to prehearing activities.

§ 405.1855 Evidence at Board hearing.

Evidence may be received at the Board hearing even though inadmissible under the rules of evidence applicable to court procedure. The Board shall give the parties opportunity for submission and consideration of facts and arguments and during the course of the hearing should, in ruling upon admissibility of evidence, exclude irrelevant, immaterial, or unduly repetitious evidence. The Board shall render a final ruling on the admissibility of evidence.

§ 405.1857 Subpoenas.

When reasonably necessary for the full presentation of a case, the Board may, either upon its own motion or upon the request of a party, issue subpoenas for the attendance and testimony of witnesses and for the production of books, records, correspondence, papers, or other documents which are relevant and material to any matter in issue at the hearing. Parties who desire the issuance of a subpoena shall, not less than 10 days prior to the time fixed for the hearing, file with the Board a written request therefor, designating the witnesses or documents to be produced, and describing the address, or location thereof with sufficient particularity to permit such witnesses or documents to be found. The request for a subpoena shall state the pertinent facts which the party expects to establish by such witnesses or documents and whether such facts could be established by other evidence without the use of a subpoena. Subpoenas, as provided for above, shall be issued in the name of the Board, and the Health Care Financing Administration shall assume the cost of the issuance and the fees and mileage of any witness so subpoenaed, as provided in section 205(d) of the Act, 42 U.S.C. 405(d).

§ 405.1859 Witnesses.

Witnesses at the hearing shall testify under oath or affirmation, unless excused by the Board for cause. The Board may examine the witnesses and shall allow the parties or their representatives to do so. Parties to the proceeding may also cross-examine witnesses.

§ 405.1861 Oral argument and written allegations.

The parties, upon their request, shall be allowed a reasonable time for the presentation of oral argument or for the filing of briefs or other written statements of allegations as to facts or law. Copies of any brief or other written statement shall be filed in sufficient number that they may be made available to all parties and to the Health Care Financing Administration.

§ 405.1863 Administrative policy at issue.

Where a party to the Board hearing puts into issue an administrative policy which is interpretative of the law or regulations, the Board will promptly notify to the Health Care Financing Administration.

§ 405.1865 Record of Board hearing.

A complete record of the proceedings at the hearing shall be made and transcribed in all cases. It shall be made available to the parties upon request. The record will not be closed until a decision has been issued.

§ 405.1867 Sources of Board’s authority.

In exercising its authority to conduct the hearings described herein, the Board must comply with all the provisions of title XVIII of the Act and regulations issued thereunder, as well as HCFA Rulings issued under the authority of the Administrator of the Health Care Financing Administration (see §401.108 of this subchapter). The Board
§ 405.1869 Scope of Board's decision-making authority.

The Board shall have the power to affirm, modify, or reverse a determination of an intermediary with respect to a cost report and to make any other modifications on matters covered by such report (including modifications adverse to the provider or other parties) even though such matters were not considered in the intermediary's determination. The opinion of the majority of those Board members deciding the case will constitute the Board's decision.

§ 405.1871 Board hearing decision and notice.

(a) The Board shall, as soon as practicable after the conclusion of its hearing, render a written decision based upon the record made at such hearing, the record established in support of the determination of the intermediary (see § 405.1803), and such other evidence as may be obtained or received by the Board. Such Board decision shall be supported by substantial evidence when the record of the Board hearing is viewed as a whole and shall cite applicable law, regulations, and HCFA Rulings. A copy of the decision shall be mailed to all parties to the hearing at their last known addresses and, at the same time, to the Administrator and HCFA.

(b) The decision of the Board provided for in paragraph (a) of this section shall be final and binding upon all parties to the hearing before the Board unless it is reviewed by the Secretary in accordance with § 405.1875, or revised in accordance with § 405.1885.

§ 405.1875 Administrator's review.

(a) General rule. (1) Except for a Board determination under § 405.1842 that it lacks the authority to decide an issue, the Administrator, at his or her discretion, may review any final decision of the Board, including a decision under § 405.1873 about the Board's jurisdiction to grant a hearing. The Administrator may exercise this discretion on his or her own motion, in response to a request from a party to a Board hearing or in response to a request from HCFA.

(b) Request for review. A party or HCFA requesting the Administrator to review a Board decision must file a written request with the Administrator within 15 days of the receipt of the Board decision.

(c) Criteria for deciding whether to review. In deciding whether to review a Board decision, either on his or her own motion or in response to a request from a party to the hearing or HCFA,
the Administrator will normally consider whether it appears that:

(1) The Board made an erroneous interpretation of law, regulation or HCFA Ruling;
(2) The Board’s decision is not supported by substantial evidence; or
(3) The case presents a significant policy issue having a basis in law and regulations, and review is likely to lead to the issuance of a HCFA Ruling or other directive needed to clarify a statutory or regulatory provision;
(4) The Board has incorrectly assumed or denied jurisdiction or extended its authority to a degree not provided for by statute, regulation or HCFA Ruling; and
(5) The decision of the Board requires clarification, amplification, or an alternative legal basis for the decision.

(d) Decision to review.

(1) Whether or not a party or HCFA has requested review, the Administrator will promptly notify the parties and HCFA whether he or she has decided to review a decision of the Board and, if so, will indicate the particular issues he or she will consider.

(2) The Administrator may decline to review a case or any issue in a case even if a party has filed a written request for review under paragraph (b) of this section.

(e) Written submissions.

(1) Within 15 days of receipt of a notice that the Administrator has decided to review a Board decision, a party or HCFA may submit to the Administrator, in writing:

(i) Proposed findings and conclusions;

(ii) Supporting views or exceptions to the Board decision;

(iii) Supporting reasons for the exceptions and proposed findings; and

(iv) A rebuttal of the other party’s request for review or other submissions already filed with the Administrator.

(2) These submissions shall be limited to issues the Administrator has decided to review and confined to the record of the Board hearing.

(3) A party or HCFA, within 15 days of receipt of a notice that the Administrator has decided to review a decision, may also request that the decision be remanded and state reasons for doing so. Reasons for a request to remand may include new, substantial evidence concerning—

(i) Issues presented to the Board; and

(ii) New issues that have arisen since the case was presented to the Board.

(4) A copy of any written submission made under this paragraph shall be sent simultaneously to each other party to the Board hearing and to HCFA, if HCFA has previously—

(i) Requested that the Administrator review a Board decision or filed a written submission in response to a party’s request for review;

(ii) Responded to a party’s request for review; or

(iii) Submitted material after the Administrator has announced that he or she will review a Board decision.

(f) Ex parte communications prohibited.

All communications from any of the parties or HCFA about a Board decision being reviewed by the Administrator must be in writing and must contain a certification that copies have been served on the parties and HCFA, as appropriate. The Administrator will not consider any communication that does not meet these requirements or is not submitted within the required time limits.

(g) Administrator’s decision.

(1) If the Administrator has notified the parties and HCFA that he or she has decided to review a Board decision, the Administrator will affirm, reverse, modify or remand the case.

(2) The Administrator will make this decision within 60 days after the provider received notification of the Board decision and will promptly mail a copy of the decision to each party and to HCFA.

(3) Any decision other than to remand will be confined to—

(i) The record of the Board, as forwarded by the Board;

(ii) Any materials submitted under paragraphs (b) or (e) of this section; and

(iii) Generally known facts that are not subject to reasonable dispute.

(4) The Administrator may rely on prior decisions of the Board, the Administrator and the courts, and other applicable law, whether or not cited by the parties and HCFA.
§ 405.1877 Judicial review.

(a) General rule. Section 1878(f) of the Act permits a provider to obtain judicial review of a final decision of the Board, or of a reversal, affirmation, or modification by the Administrator of a Board decision, by filing a civil action pursuant to the Federal Rules of Civil Procedure within 60 days of the date on which the provider received notice of—

(1) A final decision by the Board; or

(2) Any reversal, affirmation, or modification by the Administrator.

The Board's decision is not final if the Administrator reverses, affirms, or modifies the decision within 60 days of the date on which the provider received notice of the decision.

(b) Administrator declines to review a Board decision. If the Administrator declines to review a Board decision, the provider must file its appeal within 60 days of receipt of the decision of the Board.

(c) Administrator does not act after reviewing a Board decision. If the Administrator notifies the parties that he or she has decided to review a Board decision and then does not make a decision within the 60 days allotted for his or her review, this subsequent inaction constitutes an affirmance allowing a provider an additional 60 days in which to file for judicial review, beginning with the date the Administrator's time expires for taking action under §405.1875(g)(2).

(d) Matters not subject to judicial review. Certain matters affecting payments to hospital under the prospective payment system are not subject to judicial review, as provided in section 1886(d)(7) of the Act and §405.1804.

(e) Group appeals. Any action under this section by providers that are under common ownership or control (see §413.17 of this chapter) must be brought by the providers as a group with respect to any matter involving an issue common to the providers.

(f) Venue for appeals. An action for judicial review must be brought in the District Court of the United States for the judicial district in which the provider is located (or, effective April 20, 1983, in an action brought jointly by several providers, the judicial district in which the greatest number of such providers are located) or in the District Court for the District of Columbia. Effective April 20, 1983, any action for judicial review by providers under common ownership or control (§413.17 of this chapter), must be brought by such providers as a group with respect to any matter involving an issue common to the providers.

(g) Service of process. Process must be served as described under 45 CFR part 4.

§ 405.1877 Remand.

1. A remand to the Board by the Administrator vacates the Board's decision.

2. The Administrator may direct the Board to take further action with respect to the development of additional facts or new issues, or to consider the applicability of laws or regulations other than those considered by the Board. The following are not acceptable bases for remand—

(i) Presentation of evidence existing at the time of the Board hearing that was known or reasonably could have been known;

(ii) Introduction of a favorable court case that was either not available in print at the time of the Board hearing or was decided after the Board hearing;

(iii) Change of a party's representation before the Board;

(iv) Presentation of an alternative legal basis concerning an issue in dispute; or

(v) Attempted retraction of a waiver of a right made before or at the Board hearing.

3. After remand, the Board will take the action requested in the remand action and issue a new decision.

4. The new decision will be final unless the Administrator reverses, affirms, modifies, or again remands the decision in accordance with the provisions of the section.

[48 FR 45773, Oct. 7, 1983]
person it appoints to act as its representative at the proceedings, conducted in accordance with §§405.1819 and 405.1851.

§ 405.1883 Authority of representative.
A representative appointed by a provider or other party may accept or give on behalf of the provider or other party any request or notice relative to any proceeding before a hearing officer or the Board. A representative shall be entitled to present evidence and allegations as to facts and law in any proceeding affecting the party he represents and to obtain information with respect to a request for an intermediary hearing or a Board hearing made in accordance with §§405.1811, 405.1815, or 405.1837 to the same extent as the party he represents. Notice to a provider or other party of any action, determination, or decision, or a request for the production of evidence by a hearing officer or the Board sent to the representative of the provider or other party shall have the same force and effect as if it had been sent to the provider or other party.

§ 405.1885 Reopening a determination or decision.
(a) A determination of an intermediary, a decision by a hearing officer or panel of hearing officers, a decision by the Board, or a decision of the Secretary may be reopened with respect to findings on matters at issue in such determination or decision, by such intermediary officer or panel of hearing officers, Board, or Secretary, as the case may be, either on motion of such intermediary officer or panel of hearing officers, Board, or Secretary, or on the motion of the provider affected by such determination or decision to revise any matter in issue at any such proceedings. Any such request to reopen must be made within 3 years of the date of the notice of the intermediary or Board hearing decision, or where there has been no such decision, any such request to reopen must be made within 3 years of the date of notice of the intermediary determination. No such determination or decision may be reopened after such 3-year period except as provided in paragraphs (d) and (e) of this section.

(b) A determination or a hearing decision rendered by the intermediary shall be reopened and revised by the intermediary if, within the aforementioned 3-year period, the Health Care Financing Administration notifies the intermediary that such determination or decision is inconsistent with the applicable law, regulations, or general instructions issued by the Health Care Financing Administration in accordance with the Secretary's agreement with the intermediary.

(c) Jurisdiction for reopening a determination or decision rests exclusively with that administrative body that rendered the last determination or decision.

(d) Notwithstanding the provisions of paragraph (a) of this section, an intermediary determination or hearing decision, a decision of the Board, or a decision of the Secretary shall be reopened and revised at any time if it is established that such determination or decision was procured by fraud or similar fault of any party to the determination or decision.

(e) Paragraphs (a) and (b) of this section apply to determinations on cost reporting periods ending on or after December 31, 1971. (See §405.1801(c).) However, the 3-year period described shall also apply to determinations with respect to cost reporting periods ending prior to December 31, 1971, but only if the reopening action was undertaken after May 27, 1972 (the effective date of regulations which, prior to the publication of this subpart R, governed the reopening of such determinations).

§ 405.1887 Notice of reopening.
(a) All parties to any reopening described above shall be given written notice of the reopening. When such reopening results in any revision in the prior decision notice of said revision or revisions will be mailed to the parties with a complete explanation of the basis for the revision or revisions. Notices of reopenings by the Board shall also be sent to the Secretary.

(b) In any such reopening, the parties to the prior decision shall be allowed a reasonable period of time in which to present any additional evidence or argument in support of their position.
§ 405.1889 Effect of a revision.

Where a revision is made in a determination or decision on the amount of program reimbursement after such determination or decision has been reopened as provided in § 405.1885, such revision shall be considered a separate and distinct determination or decision to which the provisions of §§ 405.1811, 405.1835, 405.1875 and 405.1877 are applicable. (See § 405.1801(c) for applicable effective dates.)

Subparts S–T—[Reserved]

Subpart U—Conditions for Coverage of Suppliers of End-Stage Renal Disease (ESRD) Services

 AUTHORITY: Secs. 1102, 1138, 1861, 1862(a), 1871, 1874, and 1881 of the Social Security Act (42 U.S.C. 1302, 1320b–8, 1395x, 1395y(a), 1395hh, 1395kk, and 1395rr), unless otherwise noted.


§ 405.2100 Scope of subpart.

(a) The regulations in this subpart prescribe the role which End-Stage Renal Disease (ESRD) networks have in the ESRD program, establish the mechanism by which minimal utilization rates are promulgated and applied, under section 1881(b)(1) of the Act, and describe the health and safety requirements that facilities furnishing ESRD care to beneficiaries must meet. These regulations further prescribe the role of ESRD networks in meeting the requirements of section 1881(c) of the Act.

(b) The general objectives of the ESRD program are contained in § 405.2101, and general definitions are contained in § 405.2102. The provisions of §§ 405.2110, 405.2112 and 405.2113 discuss the establishment and activities of ESRD networks, network organizations and membership requirements and restrictions for members of the medical review boards. Sections 405.2120 through 405.2124 discuss general requirements for, and description of, all facilities furnishing ESRD services. Sections 405.2160 through 405.2164 discuss specific requirements for facilities which furnish ESRD dialysis services. Sections 405.2170 and 405.2171 discuss specific requirements for facilities which furnish ESRD transplantation services.

§ 405.2101 Objectives of the end-stage renal disease (ESRD) program.

The objectives of the end-stage renal disease program are:

(a) To assist beneficiaries who have been diagnosed as having end-stage renal disease (ESRD) to receive the care they need;

(b) To encourage proper distribution and effective utilization of ESRD treatment resources while maintaining or improving the quality of care;

(c) To provide the flexibility necessary for the efficient delivery of appropriate care by physicians and facilities; and

(d) To encourage self-dialysis or transplantation for the maximum practical number of patients who are medically, socially, and psychologically suitable candidates for such treatment.

§ 405.2102 Definitions.

As used in this subpart, the following definitions apply:

Agreement. A written document executed between an ESRD facility and another facility in which the other facility agrees to assume responsibility for furnishing specified services to patients and for obtaining reimbursement for those services.

Arrangement. A written document executed between an ESRD facility and another facility in which the other facility agrees to assume responsibility for furnishing specified services to patients and for obtaining reimbursement for those services.

Dialysis. A process by which dissolved substances are removed from a patient's body by diffusion from one fluid compartment to another across a semipermeable membrane. The two types of dialysis that are currently in
common use are hemodialysis and peritoneal dialysis. End-Stage Renal Disease (ESRD). That stage of renal impairment that appears irreversible and permanent, and requires a regular course of dialysis or kidney transplantation to maintain life.

ESRD facility. A facility which is approved to furnish at least one specific ESRD service (see definition of “ESRD service”). Such facilities are:

(a) Renal Transplantation Center. A hospital unit which is approved to furnish directly transplantation and other medical and surgical specialty services required for the care of the ESRD transplant patients, including inpatient dialysis furnished directly or under arrangement. A Renal Transplantation Center may also be a Renal Dialysis Center.

(b) Renal dialysis center. A hospital unit which is approved to furnish the full spectrum of diagnostic, therapeutic, and rehabilitative services required for the care of ESRD dialysis patients (including inpatient dialysis furnished directly or under arrangement). A hospital need not provide renal transplantation to qualify as a renal dialysis center.

(c) Renal dialysis facility. A unit which is approved to furnish dialysis service(s) directly to ESRD patients.

(d) Self-dialysis unit. A unit that is part of an approved renal transplantation center, renal dialysis center, or renal dialysis facility, and furnishes self-dialysis services.

(e) Special purpose renal dialysis facility. A renal dialysis facility which is approved under §405.2164 to furnish dialysis at special locations on a short-term basis to a group of dialysis patients otherwise unable to obtain treatment in the geographical area. The special locations must be either special rehabilitative (including vacation) locations serving ESRD patients temporarily residing there, or locations in need of ESRD facilities under emergency circumstances.

ESRD service. The type of care or services furnished to an ESRD patient. Such types of care are:

(a) Transplantation service. A process by which (1) a kidney is excised from a live or cadaveric donor, (2) that kidney is implanted in an ESRD patient, and (3) supportive care is furnished to the living donor and to the recipient following implantation.

(b) Dialysis service—(1) Inpatient dialysis. Dialysis which, because of medical necessity, is furnished to an ESRD patient on a temporary inpatient basis in a hospital;

(2) Outpatient dialysis. Dialysis furnished on an outpatient basis at a renal dialysis center or facility. Outpatient dialysis includes:

(i) Staff-assisted dialysis. Dialysis performed by the staff of the center or facility.

(ii) Self-dialysis. Dialysis performed, with little or no professional assistance, by an ESRD patient who has completed an appropriate course of training.

(3) Home dialysis. Dialysis performed by an appropriately trained patient at home.

(c) Self-dialysis and home dialysis training. A program that trains ESRD patients to perform self-dialysis or home dialysis with little or no professional assistance, and trains other individuals to assist patients in performing self-dialysis or home dialysis.

Furnishes directly. The ESRD facility provides the service through its own staff and employees, or through individuals who are under direct contract to furnish such services personally for the facility (i.e., not through “agreements” or “arrangements”).

Furnishes on the premises. The ESRD facility furnishes services on its main premises; or on its other premises that are (a) contiguous with or in immediate proximity to the main premises, and under the direction of the same professional staff and governing body as the main premises, or (b) approved on a time-limited basis as a special purpose renal dialysis facility.

Histocompatibility testing. Laboratory test procedures which determine compatibility between a potential organ donor and a potential organ transplant recipient.

Medical care criteria. Predetermined elements against which aspects of the quality of a medical service may be
compared. They are developed by professionals relying on professional expertise and on the professional literature.

Medical care norms. Numerical or statistical measures of usual observed performance. Norms are derived from aggregate information related to the health care provided to a large number of patients over a period of time.

Medical care standards. Professionally developed expressions of the range of acceptable variation from a norm or criterion.

Medical care evaluation study (MCE). Review of health care services, usually performed retrospectively, in which an indepth assessment of the quality and/or utilization of such services is made.

Network, ESRD. All Medicare-approved ESRD facilities in a designated geographic area specified by HCFA.

Network organization. The administrative governing body to the network and liaison to the Federal government.

Organ procurement. The process of acquiring donor kidneys. (See definition of Organ procurement organization in $485.302 of this chapter.)

Qualified personnel. Personnel that meet the requirements specified in this paragraph.

(a) Chief executive officer. A person who:

(1) Holds at least a baccalaureate degree or its equivalent and has at least 1 year of experience in an ESRD unit; or

(2) Is a registered nurse or physician director as defined in this definition; or

(3) As of September 1, 1976, has demonstrated capability by acting for at least 2 years as a chief executive officer in a dialysis unit or transplantation program.

(b) Dietitian. A person who:

(1) Is eligible for registration by the American Dietetic Association under its requirements in effect on June 3, 1976, and has at least 1 year of experience in clinical nutrition; or

(2) Has a baccalaureate or advanced degree with major studies in food and nutrition or dietetics, and has at least 1 year of experience in clinical nutrition.

(c) Medical record practitioner. A person who:

(1) Has graduated from a program for Medical Record Administrators accredited by the Council on Medical Education of the American Medical Association and the American Medical Record Association, and is eligible for certification as a Registered Record Administrator (RRA) by the American Medical Record Association under its requirements in effect on June 3, 1976.

(2) Has graduated from a program for Medical Record Technicians approved jointly by the Council on Medical Education of the American Medical Association and the American Medical Record Association, and is eligible for certification as an Accredited Record Technician (ART) by the American Medical Record Association under its requirements in effect June 3, 1976.

(3) Has successfully completed and received a satisfactory grade in the American Medical Record Association's Correspondence Course for Medical Record Personnel approved by the Accrediting Commission of the National Home Study Council, and is eligible for certification as an Accredited Record Technician by the American Medical Record Association under its requirements in effect June 3, 1976.

(d) Nurse responsible for nursing service. A person who is licensed as a registered nurse by the State in which practicing, and (1) has at least 12 months of experience in clinical nursing, and an additional 6 months of experience in nursing care of the patient with permanent kidney failure or undergoing kidney transplantation, including training in and experience with the dialysis process; or

(2) Has 18 months of experience in nursing care of the patient on maintenance dialysis, or in nursing care of the patient with a kidney transplant, including training in and experience with the dialysis process;

(3) If the nurse responsible for nursing service is in charge of self-care dialysis training, at least 3 months of the total required ESRD experience is in training patients in self-care.

(e) Physician-director. A physician who:

(1) Is board eligible or board certified in internal medicine or pediatrics by a professional board, and has had at least 12 months of experience or training in...
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the care of patients at ESRD facilities; or

(2) During the 5-year period prior to September 1, 1976, served for at least 12 months as director of a dialysis or transplantation program;

(3) In those areas where a physician who meets the definition in paragraph (1) or (2) of this definition is not available to direct a participating dialysis facility, another physician may direct the facility, subject to the approval of the Secretary.

(f) Social worker. A person who is licensed, if applicable, by the State in which practicing, and

(1) Has completed a course of study with specialization in clinical practice at, and holds a masters degree from, a graduate school of social work accredited by the Council on Social Work Education; or

(2) Has served for at least 2 years as a social worker, 1 year of which was in a dialysis unit or transplantation program prior to September 1, 1976, and has established a consultative relationship with a social worker who qualifies under paragraph (f)(1) of this definition.

(g) Transplantation surgeon. A person who:

(1) Is board eligible or board certified in general surgery or urology by a professional board; and

(2) Has at least 12 months training or experience in the performance of renal transplantation and the care of patients with renal transplants.

§ 405.2110 Designation of ESRD networks.

HCFA designated ESRD networks in which the approved ESRD facilities collectively provide the necessary care for ESRD patients.

(a) Effect on patient choice of facility. The designation of networks does not require an ESRD patient to seek care only through the facilities in the designated network where the patient resides, nor does the designation of networks limit patient choice of physicians or facilities, or preclude patient referral by physicians to a facility in another designated network.

(b) Redesignation of networks. HCFA will redesignate networks, as needed, to ensure that the designations are consistent with ESRD program experience, consistent with ESRD program objectives specified in §405.2101, and compatible with efficient program administration.

[51 FR 30361, Aug. 26, 1986]

§ 405.2111 [Reserved]

§ 405.2112 ESRD network organizations.

HCFA will designate an administrative governing body (network organization) for each network. The functions of a network organization include but are not limited to the following:

(a) Developing network goals for placing patients in settings for self-care and transplantation.

(b) Encouraging the use of medically appropriate treatment settings most compatible with patient rehabilitation and the participation of patients, providers of services, and renal disease facilities in vocational rehabilitation programs.

(c) Developing criteria and standards relating to the quality and appropriateness of patient care and, with respect to working with patients, facilities, and providers of services, for encouraging participation in vocational rehabilitation programs.

(d) Evaluating the procedures used by facilities in the network in assessing patients for placement in appropriate treatment modalities.

(e) Making recommendations to member facilities as needed to achieve network goals.

(f) On or before July 1 of each year, submitting to HCFA an annual report that contains the following information:

(1) A statement of the network goals.

(2) The comparative performance of facilities regarding the placement of patients in appropriate settings for—

(i) Self-care;

(ii) Transplants; and

(iii) Vocational rehabilitation programs.

(3) Identification of those facilities that consistently fail to cooperate with...
the goals specified under paragraph (f)(1) of this section or to follow the recommendations of the medical review board.

(4) Identification of facilities and providers that are not providing appropriate medical care.

(5) Recommendations with respect to the need for additional or alternative services in the network including self-dialysis training, transplantation and organ procurement.

(g) Evaluating and resolving patient grievances.

(h) Appointing a network council and a medical review board (each including at least one patient representative) and supporting and coordinating the activities of each.

(i) Conducting on-site reviews of facilities and providers as necessary, as determined by the medical review board or HCFA, using standards of care as specified under paragraph (c) of this section.

(j) Collecting, validating, and analyzing such data as necessary to prepare the reports required under paragraph (f) of this section and the Secretary's report to Congress on the ESRD program and to assure the maintenance of the registry established under section 1881(c)(7) of the Act.

§ 405.2114 [Reserved]

§ 405.2120 Minimum utilization rates: general.

Section 1881(b)(1) of the Social Security Act (42 U.S.C. 1395rr(b)(1)) authorizes the Secretary to limit payment for ESRD care to those facilities that meet the requirements that the Secretary may prescribe, including minimum utilization rates for covered transplantations. The minimum utilization rates, which are explained and specified in §§405.2121 through 405.2130, may be changed from time to time in accordance with program experience. Changes will be published as amendments to these regulations.

§ 405.2121 Basis for determining minimum utilization rates.

In developing minimum utilization rates, the Secretary takes into account the performance of ESRD facilities, the availability of care, the quality of care, and the efficient utilization of equipment and personnel, based on the following evidence:

(a) Information on the geographic distribution of ESRD patients and facilities;

(b) Information on quality of care; and

(c) Information on operational and management efficiency.

§ 405.2122 Types and duration of classification according to utilization rates.

A renal transplantation center that meets all the other conditions for coverage of ESRD services that are classified according to its utilization rate(s) as follows: Unconditional status, conditional status, exception status, or not eligible for reimbursement for that ESRD service. Such classification will be based on previously reported utilization data (see §405.2124, except as specified in paragraph (a) of this section),
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and will be effective until notification of subsequent classification occurs. (See § 405.2123 for reporting requirements; § 405.2124 for method of calculating rates; § 405.2130 for specific standards.)

(a) Initial classification. (1) A renal transplantation center that has not previously participated in the ESRD program will be granted conditional status if it submits a written plan, detailing how it will achieve the utilization rates for conditional status by the end of the second calendar year of its operation under the ESRD program, and the rates required for unconditional status by the end of its fourth calendar year of operation.

(2) The renal transplantation center's performance will be evaluated at the end of the first calendar year to ascertain whether it is properly implementing the plan.

(b) Exception status. (1) A renal transplantation center that does not meet the minimum utilization rate for unconditional or conditional status may be approved by the Secretary for a time limited exception status if:

(i) It meets all other conditions for coverage under this subpart;

(ii) It is unable to meet the minimum utilization rate because it lacks a sufficient number of patients and is located in an area without a sufficient population base to support a center or facility which would meet the rate; and

(iii) Its absence would adversely affect the achievement of ESRD program objectives.

(2) A hospital that furnishes renal transplantation services primarily to pediatric patients and is approved as a renal dialysis center under this subpart, but does not meet the utilization standards prescribed in § 405.2130(a), may be approved by the Secretary for a time limited exception status if:

(i) It meets all other conditions for coverage as a renal transplantation center;

(ii) The surgery is performed under the direct supervision of a qualified transplantation surgeon (§ 405.2102) who is also performing renal transplantation surgery at an approved renal transplantation center that is primarily oriented to adult nephrology;

(iii) It has an agreement, with the other hospital serviced by the surgeon, for sharing limited resources that are needed for kidney transplantation; and

(iv) There are pediatric patients who need the surgery and who cannot obtain it from any other hospital located within a reasonable distance.


§ 405.2123 Reporting of utilization rates for classification.

Each hospital furnishing renal transplantation services must submit an annual report to HCFA on its utilization rates. The report must include both the number of transplants performed during the most recent year of operation and the number performed during each of the preceding 2 calendar years.

[55 FR 23441, June 8, 1990]

§ 405.2124 Calculation of utilization rates for comparison with minimal utilization rate(s) and notification of status.

For purposes of classification the Secretary will use either the utilization rate for the preceding 12 months or the average utilization rate of the preceding 2 calendar years, whichever is higher. The Secretary will inform each ESRD facility and the network coordinating council of the network area in which the ESRD facility is located of the results of this classification.

§ 405.2130 Condition: Minimum utilization rates.

Unless a renal transplantation center is granted an exception under § 405.2122(b), the center must meet the following minimum utilization rate(s) for unconditional or conditional status:

(a) Unconditional status: 15 or more transplants performed annually.

(b) Conditional status: 7 to 14 transplants performed annually.

[55 FR 23441, June 8, 1990]

§ 405.2131 Condition: Provider status: Renal transplantation center or renal dialysis center.

A renal transplantation center or a renal dialysis center (§ 405.2102(e) (1) or
§ 405.2132 [Reserved]

§ 405.2133 Condition: Furnishing data and information for ESRD program administration.

The ESRD facility, laboratory performing histocompatibility testing, and organ procurement organization furnishes data and information in the manner and at the intervals specified by the Secretary, pertaining to its ESRD patient care activities and costs, for inclusion in a national ESRD medical information system and in compilations relevant to program administration, including claims processing and reimbursement. Such information is treated as confidential when it pertains to individual patients and is not disclosed except as authorized by Department regulations on confidentiality and disclosure (see 45 CFR parts 5, 5b, and part 401 of this chapter).

[53 FR 6548, Mar. 1, 1988]

§ 405.2134 Condition: Participation in network activities.

Each facility must participate in network activities and pursue network goals.

[51 FR 30362, Aug. 26, 1986]

§ 405.2135 Condition: Compliance with Federal, State and local laws and regulations.

The ESRD facility is in compliance with applicable Federal, State and local laws and regulations.

(a) Standard: licensure. Where State or applicable local law provides for the licensing of ESRD facilities, the facility is:

(1) Licensed pursuant to such law; or

(2) Approved by the agency of such State or locality responsible for such licensing as meeting the standards established for such licensing.

(b) Standard: licensure or registration of personnel. Each staff member is currently licensed or registered in accordance with applicable law.

(c) Standard: conformity with other laws. The facility is in conformity with applicable laws and regulations pertaining to fire safety, equipment, and other relevant health and safety requirements.

§ 405.2136 Condition: Governing body and management.

The ESRD facility is under the control of an identifiable governing body, or designated person(s) so functioning, with full legal authority and responsibility for the governance and operation of the facility. The governing body adopts and enforces rules and regulations relative to its own governance and to the health care and safety of patients, to the protection of the patients' personal and property rights, and to the general operation of the facility. The governing body receives and acts upon recommendations from the network organization. The governing body appoints a chief executive officer who is responsible for the overall management of the facility.

(a) Standard: disclosure of ownership. The ESRD facility supplies full and complete information to the State survey agency (§ 405.1902(a)) as to the identity of:

(1) Each person who has any direct or indirect ownership interest of 10 per centum or more in the facility, or who is the owner (in whole or in part) of any mortgage, deed of trust, note, or other obligation secured (in whole or in part) by the facility or any of the property or assets of the facility;

(2) Each officer and director of the corporation, if the facility is organized as a corporation; and

(3) Each partner, if the facility is organized as a partnership; and promptly reports to the State survey agency any changes which would affect the current accuracy of the information so required to be supplied.

(b) Standard: Operational objectives. The operational objectives of the ESRD facility, including the services that it provides, are established by the governing body and delineated in writing. The governing body adopts effective administrative rules and regulations that are designed to safeguard the health and safety of patients and to govern the general operations of the facility, in accordance with legal requirements. Such rules and regulations...
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are in writing and dated. The governing body ensures that they are operational, and that they are reviewed at least annually and revised as necessary. If the ESRD facility is engaged in the practice of hemodialyzer reuse, the governing body ensures that there are written policies and procedures with respect to reuse, to assure that recommended standards and conditions are being followed, and requires that patients be informed of the policies and procedures.

(1) The objectives of the facility are formulated in writing and clearly stated in documents appropriate for distribution to patients, facility personnel, and the public.

(2) A description of the services provided by the facility, together with a categorical listing of the types of diagnostic and therapeutic procedures that may be performed, is readily available upon request to all concerned.

(3) Admission criteria that insure equitable access to services are adopted by the facility and are readily available to the public. Access to the self-dialysis unit is available only to patients for whom the facility maintains patient care plans (see § 405.2137).

(4) The operational objectives and administrative rules and regulations of the facility are reviewed at least annually and revised as necessary by the administrative staff, medical director, and other appropriate personnel of the facility, and are adopted when approved by the governing body.

(c) Standard: chief executive officer. The governing body appoints a qualified chief executive officer who, as the ESRD facility’s administrator: Is responsible for the overall management of the facility; enforces the rules and regulations relative to the level of health care and safety of patients, and to the protection of their personal and property rights; and plans, organizes, and directs those responsibilities delegated to him by the governing body. Through meetings and periodic reports, the chief executive officer maintains ongoing liaison among the governing body, medical and nursing personnel, and other professional and supervisory staff of the facility, and acts upon recommendations made by the medical staff and the governing body. In the absence of the chief executive officer, a qualified person is authorized in writing to act on the officer’s behalf.

(1) The governing body delineates in writing the responsibilities of the chief executive officer, and ensures that he/she is sufficiently free from other duties to provide effective direction and management of the operations and fiscal affairs of the facility.

(2) The chief executive officer serves on a full-time or part-time basis, in accordance with the scope of the facility’s operations and administrative needs, and devotes sufficient time to the conduct of such responsibilities.

(3) The responsibilities of the chief executive officer include but are not limited to:

(i) Implementing the policies of the facility and coordinating the provision of services, in accordance with delegations by the governing body.

(ii) Organizing and coordinating the administrative functions of the facility, redelegating duties as authorized, and establishing formal means of accountability for those involved in patient care.

(iii) Authorizing expenditures in accordance with established policies and procedures.

(iv) Familiarizing the staff with the facility’s policies, rules, and regulations, and with applicable Federal, State, and local laws and regulations.

(v) Maintaining and submitting such records and reports, including a chronological record of services provided to patients, as may be required by the facility’s internal committees and governing body, or as required by the Secretary.

(vi) Participating in the development, negotiation, and implementation of agreements or contracts into which the facility may enter, subject to approval by the governing body of such agreements or contracts.

(vii) Participating in the development of the organizational plan and ensuring the development and implementation of an accounting and reporting system, including annual development of a detailed budgetary program, maintenance of fiscal records, and quarterly submission to the governing body of reports of expenses and revenues generated through the facility’s operation.
(viii) Ensuring that the facility employs the number of qualified personnel needed; that all employees have appropriate orientation to the facility and their work responsibilities upon employment; and that they have an opportunity for continuing education and related development activities.

(d) Standard: personnel policies and procedures. The governing body, through the chief executive officer of the ESRD facility, is responsible for maintaining and implementing written personnel policies and procedures that support sound patient care and promote good personnel practices. These policies and procedures ensure that:

1. All members of the facility’s staff are qualified to perform the duties and responsibilities assigned to them and meet such Federal, State, and local professional requirements as may apply.

2. A safe and sanitary environment for patients and personnel exists, and reports of incidents and accidents to patients and personnel are reviewed to identify health and safety hazards. Health supervision of personnel is provided, and they are referred for periodic health examinations and treatments as necessary or as required by Federal, State, and local laws. Procedures are established for routine testing to ensure detection of hepatitis and other infectious diseases.

3. If the services of trainees are utilized in providing ESRD services, such trainees are under the direct supervision of qualified professional personnel.

4. Complete personnel records are maintained on all personnel. These include health status reports, resumes of training and experience, and current job descriptions that reflect the employees’ responsibilities and work assignments.

5. Personnel policies are written and made available to all personnel in the facility. The policies provide for an effective mechanism to handle personnel grievances.

6. All personnel of the facility participate in educational programs on a regular basis. These programs cover initial orientation, and continuing in-service training, including procedures for infection control. Records are maintained showing the content of training sessions and the attendance at such sessions.

7. Personnel manuals are maintained, periodically updated, and made available to all personnel involved in patient care.

(e) Standard: use of outside resources. If the ESRD facility makes arrangements for the provision of a specific service as authorized in this subpart, the responsibilities, functions, objectives, and the terms of each arrangement, including financial provisions and charges, are delineated in a document signed by an authorized representative of the facility and the person or agency providing the service. The chief executive officer when utilizing outside resource, as a consultant, assures that he is apprised of recommendations, plans for implementation, and continuing assessment through dated, signed reports, which are retained by the chief executive officer for follow-up action and evaluation of performance.

(f) Standard: patient care policies. The ESRD facility has written policies, approved by the governing body, concerning the provision of dialysis and other ESRD services to patients. The governing body reviews implementation of policies periodically to ensure that the intent of the policies is carried out. These policies are developed by the physician responsible for supervising and directing the provision of ESRD services, or the facility’s organized medical staff (if there is one), with the advice of (and with provision for review of such policies from time to time, but at least annually, by) a group of professional personnel associated with the facility, including, but not limited to, one or more physicians and one or more registered nurses experienced in rendering ESRD care.

1. The patient care policies cover the following:

   i. Scope of services provided by the facility (either directly or under arrangement).

   ii. Admission and discharge policies (in relation to both in-facility care and home care).

   iii. Medical supervision and physician services.
§ 405.2137 Condition: Patient long-term program and patient care plan.

Each facility maintains for each patient a written long-term program and a written patient care plan to ensure that each patient receives the appropriate modality of care and the appropriate care within that modality. The patient, or where appropriate, parent or legal guardian is involved with the health team in the planning of care. A copy of the current program and plan accompany the patient on interfacility transfer.

(a) Standard: patient long-term program. There is a written long-term program representing the selection of a suitable treatment modality (i.e., dialysis or transplantation) and dialysis setting (e.g., home, self-care) for each patient.

(b) The program is developed by a professional team which includes but is not limited to the physician director of the dialysis facility or center where the patient is currently being treated,
§ 405.2138 Condition: Patients’ rights and responsibilities.

The governing body of the ESRD facility adopts written policies regarding a physician director of a center or facility which offers self-care dialysis training (if not available at the location where the patient is being treated), a transplant surgeon, a qualified nurse responsible for nursing services, a qualified dietitian and a qualified social worker.

(2) The program is formally reviewed and revised in writing as necessary by a team which includes but is not limited to the physician director of the dialysis facility or center where the patient is presently being treated, in addition to the other personnel listed in paragraph (a)(1) of this section at least every 12 months or more often as indicated by the patient’s response to treatment (see §405.2161(b)(1) and §405.2170(a)).

(3) The patient, parent, or legal guardian, as appropriate, is involved in the development of the patient’s long-term program, and due consideration is given to his preferences.

(4) A copy of the patient’s long-term program accompanies the patient on interfacility transfer or is sent within 1 working day.

(b) Standard: patient care plan. There is a written patient care plan for each patient of an ESRD facility (including home dialysis patients under the supervision of the ESRD facility; see §405.2163(e)), based upon the nature of the patient’s illness, the treatment prescribed, and an assessment of the patient’s needs.

(1) The patient care plan is personalized for the individual, reflects the psychological, social, and functional needs of the patient, and indicates the ESRD and other care required as well as the individualized modifications in approach necessary to achieve the long-term and short-term goals.

(2) The plan is developed by a professional team consisting of at least the physician responsible for the patient’s ESRD care, a qualified nurse responsible for nursing services, a qualified social worker, and a qualified dietitian.

(3) The patient, parent, or legal guardian, as appropriate, is involved in the development of the care plan, and due consideration is given to his preferences.

(4) The care plan for patients whose medical condition has not become stabilized is reviewed at least monthly by the professional patient care team described in paragraph (b)(2) of this section. For patients whose condition has become stabilized, the care plan is reviewed every 6 months. The care plan is revised as necessary to insure that it provides for the patients ongoing needs.

(5) If the patient is transferred to another facility, the care plan is sent with the patient or within 1 working day.

(6) For a home-dialysis patient whose care is under the supervision of the ESRD facility, the care plan provides for periodic monitoring of the patient’s home adaptation, including provisions for visits to the home by qualified facility personnel to the extent appropriate. (See §405.2163(e).)

(7) Beginning July 1, 1991, for a home dialysis patient, and beginning January 1, 1994, for any dialysis patient, who uses EPO in the home, the plan must provide for monitoring home use of EPO that includes the following:

(i) Review of diet and fluid intake for indiscretions as indicated by hyperkalemia and elevated blood pressure secondary to volume overload.

(ii) Review of medications to ensure adequate provision of supplemental iron.

(iii) Ongoing evaluations of hematocrit and iron stores.

(iv) A reevaluation of the dialysis prescription taking into account the patient’s increased appetite and red blood cell volume.

(v) A method for physician followup on blood tests and a mechanism (such as a patient log) for keeping the physician informed of the results.

(vi) Training of the patient to identify the signs and symptoms of hypertension and hypotension.

(vii) The decrease or discontinuance of EPO if hypertension is uncontrolled.

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the rights and responsibilities of patients and, through the chief executive officer, is responsible for development of, and adherence to, procedures implementing such policies. These policies and procedures are made available to patients and any guardians, next of kin, sponsoring agency(ies), representative payee(s) (selected pursuant to section 205(j) of the Social Security Act and subpart Q of 20 CFR part 404), and to the public. The staff of the facility is trained and involved in the execution of such policies and procedures. The patients' rights policies and procedures ensure at least the following:

(a) Standard: informed patients. All patients in the facility:
   (1) Are fully informed of these rights and responsibilities, and of all rules and regulations governing patient conduct and responsibilities;
   (2) Are fully informed of services available in the facility and of related charges including any charges for services not covered under title XVIII of the Social Security Act;
   (3) Are fully informed by a physician of their medical condition unless medically contraindicated (as documented in their medical records);
   (4) Are fully informed regarding the facility's reuse of dialysis supplies, including hemodialyzers. If printed materials such as brochures are utilized to describe a facility and its services, they must contain a statement with respect to reuse; and
   (5) Are fully informed regarding their suitability for transplantation and home dialysis.

(b) Standard: participation in planning. All patients treated in the facility:
   (1) Are afforded the opportunity to participate in the planning of their medical treatment and to refuse to participate in experimental research;
   (2) Are transferred or discharged only for medical reasons or for the patient's welfare or that of other patients, or for nonpayment of fees (except as prohibited by title XVIII of the Social Security Act), and are given advance notice to ensure orderly transfer or discharge.

(c) Standard: respect and dignity. All patients are treated with consideration, respect, and full recognition of their individuality and personal needs, including the need for privacy in treatment. Provision is made for translators where a significant number of patients exhibit language barriers.

(d) Standard: confidentiality. All patients are ensured confidential treatment of their personal and medical records, and may approve or refuse release of such records to any individual outside the facility, except in case of their transfer to another health care institution or as required by Federal, State, or local law and the Secretary for proper administration of the program.

(e) Standard: grievance mechanism. All patients are encouraged and assisted to understand and exercise their rights. Grievances and recommended changes in policies and services may be addressed to facility staff, administration, the network organization, and agencies or regulatory bodies with jurisdiction over the facility, through any representative of the patient's choice, without restraint or interference, and without fear of discrimination or reprisal.


§ 405.2139 Condition: Medical records.

The ESRD facility maintains complete medical records on all patients (including self-dialysis patients within the self-dialysis unit and home dialysis patients whose care is under the supervision of the facility) in accordance with accepted professional standards and practices. A member of the facility's staff is designated to serve as supervisor of medical records services, and ensures that all records are properly documented, completed, and preserved. The medical records are completely and accurately documented, readily available, and systematically organized to facilitate the compilation and retrieval of information.

(a) Standard: medical record. Each patient's medical record contains sufficient information to identify the patient clearly, to justify the diagnosis and treatment, and to document the results accurately. All medical records contain the following general categories of information: Documented evidence of assessment of the needs of the patient, whether the patient is

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Condition: Physical environment.

The physical environment in which ESRD services are furnished affords a functional, sanitary, safe, and comfortable setting for patients, staff, and the public.

(a) Standard: building and equipment.

The physical structure in which ESRD services are furnished is constructed, equipped, and maintained to assure the safety of patients, staff, and the public.

(1) Fire extinguishers are conveniently located on each floor of the facility and in areas of special hazard.
Fire regulations and fire management procedures are prominently posted and properly followed.

(2) All electrical and other equipment used in the facility is maintained free of defects which could be a potential hazard to patients or personnel. There is established a planned program of preventive maintenance of equipment used in dialysis and related procedures in the facility.

(3) The areas used by patients are maintained in good repair and kept free of hazards such as those created by damaged or defective parts of the building.

(4) [Reserved]

(5)(i) The ESRD facility must employ the water quality requirements listed in paragraph (a)(5)(ii) of this section developed by the Association for the Advancement of Medical Instrumentation (AAMI) and published in “Hemodialysis Systems,” second edition, which is incorporated by reference.

(ii) Required water quality requirements are those listed in sections 3.2.1, Water Bacteriology; 3.2.2, Maximum Level of Chemical Contaminants; and in Appendix B: Guideline for Monitoring Purity of Water Used for Hemodialysis as B1 through B5.

(iii) Incorporation by reference of the AAMI’s “Hemodialysis Systems,” second edition, 1992, was approved by the Director of the Federal Register in accordance with 5 U.S.C. 552(a) and 1 CFR part 51. If any changes in “Hemodialysis Systems,” second edition, are also to be incorporated by reference, a notice to that effect will be published in the Federal Register.

(b) Standard: Favorable environment for patients. The facility is maintained and equipped to provide a functional sanitary, and comfortable environment with an adequate amount of well-lighted space for the service provided.

(1) There are written policies and procedures in effect for preventing and controlling hepatitis and other infections. These policies include, but are not limited to, appropriate procedures for surveillance and reporting of infections, housekeeping, handling and disposal of waste and contaminants, and sterilization and disinfection, including the sterilization and maintenance of equipment where dialysis supplies are reused; there are written policies and procedures covering the rinsing, cleaning, disinfection, preparation and storage of reused items which conform to requirements for reuse in §405.2150.

(2) Treatment areas are designed and equipped to provide adequate and safe dialysis therapy, as well as privacy and comfort for patients. The space for treating each patient is sufficient to accommodate medically needed emergency equipment and staff and to ensure that such equipment and staff can reach the patient in an emergency. There is sufficient space in units for safe storage of self-dialysis supplies.

(3) There is a nursing/monitoring station from which adequate surveillance of patients receiving dialysis services can be made.

(4) Heating and ventilation systems are capable of maintaining adequate and comfortable temperatures.

(5) Each ESRD facility utilizing a central-batch delivery system provides, either on the premises or through affiliation agreement or arrangement (see §405.2160) sufficient individual delivery systems for the treatment of any patient requiring special dialysis solutions.

(c) Standard contamination prevention. The facility employs appropriate techniques to prevent cross-contamination between the unit and adjacent hospital or public areas including, but not limited to, food service areas, laundry, disposal of solid waste and blood-contaminated equipment, and disposal of contaminants into sewage systems. Waste storage and disposal are carried out in accordance with applicable local laws and accepted public health procedures. The written patient care policies (see §405.2136(f)(1)) specify the functions that are carried out by facility personnel and by the self-dialysis patients.
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with respect to contamination prevention. Where dialysis supplies are reused, records are maintained that can be used to determine whether established procedures covering the rinsing, cleaning, disinfection, preparation and storage of reused items, conform to requirements for reuse in § 405.2150.

(d) Standard: emergency preparedness. Written policies and procedures specifically define the handling of emergencies which may threaten the health or safety of patients. Such emergencies would exist during a fire or natural disaster or during functional failures in equipment. Specific emergency preparedness procedures exist for different kinds of emergencies. These are reviewed and tested at least annually and revised as necessary by, or under the direction of, the chief executive officer. All personnel are knowledgeable and trained in their respective roles in emergency situations.

(1) There is an established written plan for dealing with fire and other emergencies which, when necessary, is developed in cooperation with fire and other expert personnel.

(2) All personnel are trained, as part of their employment orientation, in all aspects of preparedness for any emergency or disaster. The emergency preparedness plan provides for orientation and regular training and periodic drills for all personnel in all procedures so that each person promptly and correctly carries out a specified role in case of an emergency.

(3) There is available at all times on the premises a fully equipped emergency tray, including emergency drugs, medical supplies, and equipment, and staff are trained in its use.

(4) The staff is familiar with the use of all dialysis equipment and procedures to handle medical emergencies.

(5) Patients are trained to handle medical and nonmedical emergencies. Patients must be fully informed regarding what to do, where to go, and whom to contact if a medical or nonmedical emergency occurs.

(See, 1102, 1871, 1901(b), Social Security Act; 42 U.S.C. 1302, 1395hh, 1395rr(b))


§ 405.2150 Condition: Reuse of hemodialyzers and other dialysis supplies.

An ESRD facility that reuses hemodialyzers and other dialysis supplies meets the requirements of this section. Failure to meet any of paragraphs (a) through (c) of this section constitutes grounds for denial of payment for the dialysis treatment affected and termination from participation in the Medicare program.

(a) Standard: Hemodialyzers. If the ESRD facility reuses hemodialyzers, it conforms to the following:

(1) Reuse guidelines. Voluntary guidelines adopted by the AAMI ("Reuse of Hemodialyzers," second edition). Incorporation by reference of the AAMI’s "Reuse of Hemodialyzers," second edition, 1993, was approved by the Director of the Federal Register in accordance with 5 U.S.C. 552(a) and 1 CFR part 51. If any changes in "Reuse of Hemodialyzers," second edition, are also to be incorporated by reference, a notice to that effect will be published in the Federal Register.

(2) Procedure for chemical germicides. To prevent any risk of dialyzer membrane leaks due to the combined action of different chemical germicides,
dialyzers are exposed to only one chemical germicide during the reprocessing procedure. If a dialyzer is exposed to a second germicide, the dialyzer must be discarded.

(3) Surveillance of patient reactions. In order to detect bacteremia and to maintain patient safety when unexplained events occur, the facility—
   (i) Takes appropriate blood cultures at the time of a febrile response in a patient; and
   (ii) If pyrogenic reactions, bacteremia, or unexplained reactions associated with ineffective reprocessing are identified, terminates reuse of hemodialyzers in that setting and does not continue reuse until the entire reprocessing system has been evaluated.

(b) Standard: Transducer filters. To control the spread of hepatitis, transducer filters are changed after each dialysis treatment and are not reused.

(c) Standard: Bloodlines. If the ESRD facility reuses bloodlines, it must—
   (1) Limit the reuse of bloodlines to the same patient;
   (2) Not reuse bloodlines labeled for “single use only”;
   (3) Reuse only bloodlines for which the manufacturer’s protocol for reuse has been accepted by the Food and Drug Administration (FDA) pursuant to the premarket notification (section 510(k)) provision of the Food, Drug, and Cosmetic Act; and
   (4) Follow the FDA-accepted manufacturer’s protocol for reuse of that bloodline.


§ 405.2160 Condition: Affiliation agreement or arrangement.

(a) A renal dialysis facility and a renal dialysis center (see §405.2102(e)(2)) have in effect an affiliation agreement or arrangement with each other, in writing, for the provision of inpatient care and other hospital services.

(b) The affiliation agreement or arrangement provides the basis for effective working relationships under which inpatient hospital care or other hospital services are available promptly to the dialysis facility’s patients when needed. The dialysis facility has in its files documentation from the renal dialysis center to the effect that patients from the dialysis facility will be accepted and treated in emergencies. There are reasonable assurances that:
   (1) Transfer or referral of patients will be effected between the renal dialysis center and the dialysis facility whenever such transfer or referral is determined as medically appropriate by the attending physician, with timely acceptance and admission;
   (2) There will be interchange, within 1 working day, of the patient long-term program and patient care plan, and of medical and other information necessary or useful in the care and treatment of patients transferred or referred between the facilities, or in determining whether such patients can be adequately cared for otherwise than in either of such facilities; and
   (3) Security and accountability for patients’ personal effects are assured.

§ 405.2161 Condition: Director of a renal dialysis facility or renal dialysis center.

Treatment is under the general supervision of a Director who is a physician. The physician-director need not devote full time as Director but is responsible for planning, organizing, conducting, and directing the professional ESRD services and must devote sufficient time to carrying out these responsibilities. The director may also serve as the Chief Executive Officer of the facility.

(a) Standard: qualifications. The director of a dialysis facility is a qualified physician-director. (See §405.2102.)

(b) Standard: responsibilities. The responsibilities of the physician-director include but are not limited to the following:
   (1) Participating in the selection of a suitable treatment modality, i.e., transplantation or dialysis, and dialysis setting, for all patients;
   (2) Assuring adequate training of nurses and technicians in dialysis techniques;
   (3) Assuring adequate monitoring of the patient and the dialysis process, including, for self-dialysis patients, assuring periodic assessment of patient performance of dialysis tasks;
   (4) Assuring the development and availability of a patient care policy
§ 405.2162 Condition: Staff of a renal dialysis facility or renal dialysis center.

Properly trained personnel are present in adequate numbers to meet the needs of the patients, including those arising from medical and nonmedical emergencies.

(a) Standard: Registered nurse. The dialysis facility employs at least one full time qualified nurse responsible for nursing service. (See §405.2102.)

(b) Standard: On-duty personnel. Whenever patients are undergoing dialysis:

(1) One currently licensed health professional (e.g., physician, registered nurse, or licensed practical nurse) experienced in rendering ESRD care is on duty to oversee ESRD patient care;

(2) An adequate number of personnel are present so that the patient/staff ratio is appropriate to the level of dialysis care being given and meets the needs of patients; and

(3) An adequate number of personnel are readily available to meet medical and nonmedical needs.

(c) Standard: Self-care dialysis training personnel. If the facility offers self-care dialysis training, a qualified nurse is in charge of such training (see §405.2102.)

§ 405.2163 Condition: Minimal service requirements for a renal dialysis facility or renal dialysis center.

The facility must provide dialysis services, as well as adequate laboratory, social, and dietetic services to meet the needs of the ESRD patient.

(a) Standard: Outpatient dialysis services—(1) Staff-assisted dialysis services. The facility must provide all necessary institutional dialysis services and staff required in performing the dialysis.

(2) Self-dialysis services. If the facility offers self-dialysis services, it must provide all medically necessary supplies and equipment and any other service specified in the facility's patient care policies.

(b) Standard: Laboratory services. The dialysis facility makes available laboratory services (other than the specialty of tissue pathology and histocompatibility testing), to meet the needs of the ESRD patient. All laboratory services must be performed by an appropriately certified laboratory in accordance with part 493 of this chapter. If the renal dialysis facility furnishes its own laboratory services, it must meet the applicable requirements established for certification of laboratories found in part 493 of this chapter. If the facility does not provide laboratory services, it must make arrangements to obtain these services from a laboratory certified in the appropriate specialties and subspecialties of service in accordance with the requirements of part 493 of this chapter.

(c) Standard: Social services. Social services are provided to patients and their families and are directed at supporting and maximizing the social functioning and adjustment of the patient. Social services are furnished by a qualified social worker (§405.2102) who has an employment or contractual relationship with the facility. The qualified social worker is responsible for conducting psychosocial evaluations, participating in team review of patient progress and recommending changes in treatment based on the patient's current psychosocial needs, providing casework and groupwork services to patients and their families in dealing with the special problems associated with ESRD.
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with ESRD, and identifying community social agencies and other resources and assisting patients and families to utilize them.

(d) Standard: Dietetic services. Each patient is evaluated as to his nutritional needs by the attending physician and by a qualified dietician (§405.2102) who has an employment or contractual relationship with the facility. The dietician, in consultation with the attending physician, is responsible for assessing the nutritional and dietetic needs of each patient, recommending therapeutic diets, counseling patients and their families on prescribed diets, and monitoring adherence and response to diets.

(e) Standard: Self-dialysis support services. The renal dialysis facility or center furnishing self-dialysis training upon completion of the patient's training, furnishes (either directly, under agreement or by arrangement with another ESRD facility) the following services:

(1) Surveillance of the patient's home adaptation, including provisions for visits to the home or the facility;

(2) Consultation for the patient with a qualified social worker and a qualified dietician;

(3) A recordkeeping system which assures continuity of care;

(4) Installation and maintenance of equipment;

(5) Testing and appropriate treatment of the water; and

(6) Ordering of supplies on an ongoing basis.

(f) Standard: Participation in recipient registry. The dialysis facility or center participates in a patient registry program with an OPO designated or redesignated under part 486, subpart G of this chapter, for patients who are awaiting cadaveric donor transplantation.

(g) Use of EPO at home: Patient selection. The dialysis facility, or the physician responsible for all dialysis-related services furnished to the patient, must make a comprehensive assessment that includes the following:

(1) Pre-selection monitoring. The patient's hematocrit (or hemoglobin), serum iron, transferrin saturation, serum ferritin, and blood pressure must be measured.

(2) Conditions the patient must meet. The assessment must find that the patient meets the following conditions:

(i) On or after July 1, 1991, is a home dialysis patient or, on or after January 1, 1994, is a dialysis patient;

(ii) Has a hematocrit (or comparable hemoglobin level) that is as follows:

(A) For a patient who is initiating EPO treatment, no higher than 30 percent unless there is medical documentation showing the need for EPO despite a hematocrit (or comparable hemoglobin level) higher than 30 percent. (Patients with severe angina, severe pulmonary distress, or severe hypertension may require EPO to prevent adverse symptoms even if they have higher hematocrit or hemoglobin levels.)

(B) A renal dialysis facility that establishes the plan of care and monitors the progress of the home EPO therapy.

(3) Conditions the patient or the patient's caregiver must meet. The assessment must find that the patient or a caregiver who assists the patient in performing self-dialysis meets the following conditions:

(i) Is trained by the facility to inject EPO and is capable of carrying out the procedure.

(ii) Is capable of reading and understanding the drug labeling.

(iii) Is trained in, and capable of observing, aseptic techniques.

(4) Care and storage of drug. The assessment must find that EPO can be stored in the patient's residence under refrigeration and that the patient is aware of the potential hazard of a child's having access to the drug and syringes.

(h) Use of EPO at home: Responsibilities of the physician or the dialysis facility. The patient's physician or dialysis facility must—

(1) Develop a protocol that follows the drug label instructions;
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(2) Make the protocol available to the patient to ensure safe and effective home use of EPO; and

(3) Through the amounts prescribed, ensure that the drug “on hand” at any time does not exceed a 2-month supply.


§ 405.2164 Conditions for coverage of special purpose renal dialysis facilities.

(a) A special purpose renal dialysis facility must comply with all conditions for coverage for renal dialysis facilities specified in §§ 405.2130 through 405.2164, with the exception of §§ 405.2134, and 405.2137 that relate to participation in the network activities and patient long-term programs.

(b) A special purpose renal dialysis facility must consult with a patient’s physician to assure that care provided in the special purpose dialysis facility is consistent with the patient’s long-term program and patient care plan required under § 405.2137.

(c) The period of approval for a special purpose renal dialysis facility may not exceed 8 calendar months in any calendar year.

(d) A special purpose renal dialysis facility may provide services only to those patients who would otherwise be unable to obtain treatments in the geographical areas served by the facility.


§ 405.2170 Condition: Director of a renal transplantation center.

The renal transplantation center is under the general supervision of a qualified transplantation surgeon (§ 405.2102) or a qualified physician-director (§ 405.2102), who need not serve full time. This physician is responsible for planning, organizing, conducting, and directing the renal transplantation center and devotes sufficient time to carry out these responsibilities, which include but are not limited to the following:

(a) Participating in the selection of a suitable treatment modality for each patient.

(b) Assuring adequate training, of nurses in the care of transplant patients.

(c) Assuring that tissue typing and organ procurement services are available either directly or under arrangement.

(d) Assuring that transplantation surgery is performed under the direct supervision of a qualified transplantation surgeon.

(c) Standard: Dietetic services. Each patient is evaluated as to his nutritional needs by the attending physician and a qualified dietician (§ 405.2102) who has an employment or contractual relationship with the facility. The dietician, in consultation with the attending physician, is responsible for assessing the nutritional and dietetic needs of each patient, recommending therapeutic diets, counseling patients and their families on prescribed diets, and monitoring adherence and response to diets.

(d) Standard: Laboratory services: (1) The renal transplantation center makes available, directly or under arrangements, laboratory services to meet the needs of ESRD patients. Laboratory services are performed in a laboratory facility certified in accordance with part 493 of this chapter.

(2) Laboratory services for crossmatching of recipient serum and donor lymphocytes for pre-formed antibodies by an acceptable technique are available on a 24-hour emergency basis.

(e) Standard: Organ procurement. A renal transplantation center using the services of an organ procurement organization designated or redesignated under part 485, subpart D of this chapter to obtain donor organs has a written agreement covering these services. The renal transplantation center agrees to notify HCFA in writing within 30 days of the termination of the agreement.

§ 405.2180 Termination of Medicare coverage.

(a) Except as provided in § 405.2181, failure of a supplier of ESRD services to meet one or more of the conditions for coverage set forth in this subpart U will result in termination of Medicare coverage of the services furnished by that supplier.

(b) If termination of coverage is based solely on a supplier’s failure to participate in network activities and pursue network goals, as required by § 405.2134, coverage may be reinstated when HCFA determines that the supplier is making reasonable and appropriate efforts to meet that condition.

(c) If termination of coverage is based on failure to meet any of the other conditions specified in this subpart, coverage will not be reinstated until HCFA finds that the reason for termination has been removed and there is reasonable assurance that it will not recur.

§ 405.2181 Alternative sanctions.

(a) Basis for application of alternative sanctions. HCFA may, as an alternative to termination of Medicare coverage, impose one of the sanctions specified in paragraph (b) of this section if HCFA finds that—

(1) The supplier fails to participate in the activities and pursue the goals of the ESRD network that is designated to encompass its geographic area; and

(2) This failure does not jeopardize patient health and safety.

(b) Alternative sanctions. The alternative sanctions that HCFA may apply in the circumstances specified in paragraph (a) of this section include the following:

(1) Denial of payment for services furnished to patients first accepted for care after the effective date of sanction as specified in the sanction notice.

(2) Reduction of payments, for all ESRD services furnished by the supplier, by 20 percent for each 30-day period after the effective date of sanction.

(3) Withholding of all payments, without interest, for all ESRD services furnished by the supplier to Medicare beneficiaries.

(c) Duration of sanction. An alternative sanction remains in effect until HCFA finds that the supplier is in substantial compliance with the requirements to cooperate in the network plans and goals, or terminates coverage of
§ 405.2182 Notice of sanction and appeal rights: Termination of coverage.

(a) Notice of sanction. HCFA gives the supplier and the general public notice of sanction and of the effective date of the sanction. The effective date of the sanction is at least 30 days after the date of the notice.

(b) Appeal rights. Termination of Medicare coverage of a supplier's ESRD services because the supplier no longer meets the conditions for coverage of its services is an initial determination appealable under part 498 of this chapter.

[53 FR 36277, Sept. 19, 1988]

§ 405.2184 Notice of appeal rights: Alternative sanctions.

If HCFA proposes to apply a sanction specified in §405.2181(b), the following rules apply:

(a) HCFA gives the facility notice of the proposed sanction and 15 days in which to request a hearing.

(b) If the facility requests a hearing, HCFA provides an informal hearing by a HCFA official who was not involved in making the appealed decision.

(c) During the informal hearing, the facility—

(1) May be represented by counsel;

(2) Has access to the information on which the allegation was based; and

(3) May present, orally or in writing, evidence and documentation to refute the finding of failure to participate in network activities and pursue network goals.

(d) If the written decision of the informal hearing supports application of the alternative sanction, HCFA provides the facility and the public, at least 30 days before the effective date of the sanction, with a written notice that specifies the effective date and the reasons for the sanction.

[53 FR 36277, Sept. 19, 1988]

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year for items and services covered under Part B of title XVIII; and
(2) The expenses incurred for the first 3 pints of blood or 3 units of packed red blood cells furnished to a beneficiary during any calendar year. (See §§ 410.160 and 410.161 of this chapter for greater detail.)

Federally qualified health center (FQHC) means an entity that has entered into an agreement with HCFA to meet Medicare program requirements under §§ 405.2434 and—
(1) Is receiving a grant under section 329, 330, or 340 of the Public Health Service Act, or is receiving funding from such a grant under a contract with the recipient of such a grant and meets the requirements to receive a grant under section 329, 330 or 340 of the Public Health Service Act;
(2) Based on the recommendation of the PHS, is determined by HCFA to meet the requirements for receiving such a grant;
(3) Was treated by HCFA, for purposes of part B, as a comprehensive federally funded health center (FFHC) as of January 1, 1990; or
(4) Is an outpatient health program or facility operated by a tribe or tribal organizations under the Indian Self-Determination Act or by an Urban Indian organization receiving funds under title V of the Indian Health Care Improvement Act.

HCFA stands for Health Care Financing Administration.

Intermittent nursing care means a medically predictable need for nursing care from time to time, but usually not less frequently than once every 60 days.

Nurse-midwife means a registered professional nurse who meets the following requirements:
(1) Is currently licensed to practice in the State as a registered professional nurse.
(2) Is legally authorized under State law or regulations to practice as a nurse-midwife.
(3) Except as provided in paragraph (b)(10)(iv) of this section, has completed a program of study and clinical experience for nurse-midwives, as specified by the State.
(4) If the State does not specify a program of study and clinical experience that nurse-midwives must complete to practice in that State, meets one of the following conditions:
   (i) Is currently certified as a nurse-midwife by the American College of Nurse-Midwives.
   (ii) Has satisfactorily completed a formal education program (of at least one academic year) that, upon completion, qualifies the nurse to take the certification examination offered by the American College of Nurse-Midwives.
   (iii) Has successfully completed a formal educational program for preparing registered nurses to furnish gynecological and obstetrical care to women during pregnancy, delivery, and the postpartum period, and care to normal newborns, and was practicing as a nurse-midwife for a total of 12 months during any 18-month period from August 8, 1976 to July 16, 1982.

Nurse practitioner and physician assistant means individuals who meet the applicable education, training experience and other requirements of § 491.2 of this chapter.

Part-time nursing care means nursing care that is required on less than a full-time basis, that is, less than 8 hours a day or 40 hours a week.

Physician means the following:
(1) A doctor of medicine or osteopathy legally authorized to practice medicine and surgery by the State in which the function is performed.
(2) Within limitations as to the specific services furnished, a doctor of dentistry or dental or oral surgery, a doctor of optometry, a doctor of podiatry or surgical chiropody or a chiropractor. (See section 1861(r) of the Act for specific limitations.)
(3) A resident (including residents as defined in § 415.152 of this chapter who meet the requirements in § 415.206(b) of this chapter for payment under the physician fee schedule).

Reporting period means a period of 12 consecutive months specified by the intermediary as the period for which a clinic or center must report its costs and utilization. The first and last reporting periods may be less than 12 months.

Rural health clinic means a facility that:
§ 405.2402  Basic requirements.

(a) Certification by the State survey agency. The rural health clinic must be certified in accordance with part 491 of this chapter.

(b) Acceptance of the clinic as qualified to furnish rural health clinic services. If the Secretary, after reviewing the survey agency recommendation and other evidence relating to the qualifications of the rural health clinic, determines that it meets the requirements of this subpart and of part 491 of this chapter, he will send the clinic:

  (1) Written notice of the determination; and
  (2) Two copies of the agreement to be filed as required by section 1861(aa)(1) of the Act.

(c) Filing of agreement by the rural health clinic. If the rural health clinic wishes to participate in the program, it must:

  (1) Have both copies of the agreement signed by an authorized representative; and
  (2) File them with the Secretary.

(d) Acceptance by the Secretary. If the Secretary accepts the agreement filed by the rural health clinic, he will return to the clinic one copy of the agreement, with a notice of acceptance specifying the effective date.

(e) Duration of agreement. The agreement shall be for a term of one year and may be renewed annually by mutual consent of the Secretary and the rural health clinic.

(f) Appeal rights. If the Secretary does not certify a rural health clinic, or refuses to enter into or renew an agreement, the facility is entitled to a hearing in accordance with part 498 of this chapter.

§ 405.2403  Content and terms of the agreement with the Secretary.

(a) Under the agreement, the rural health clinic agrees to the following:

  (1) Maintaining compliance with conditions. The clinic agrees to maintain compliance with the conditions set forth in part 491 of this chapter and to report promptly to HCFA any failure to do so.

  (2) Charges to beneficiaries. The clinic agrees not to charge the beneficiary or any other person for items and services for which the beneficiary is entitled to have payment made under the provisions of this part (or for which the beneficiary would have been entitled if the rural health clinic had filed a request for payment in accordance with § 410.165 of this chapter), except for any deductible or coinsurance amounts for which the beneficiary is liable under § 405.2410.

  (3) Refunds to beneficiaries. (i) The clinic agrees to refund as promptly as possible any money incorrectly collected from beneficiaries or from someone on their behalf.

     (ii) As used in this section, money incorrectly collected means sums collected in excess of the amount for which the beneficiary was liable under § 405.2410. It includes amounts collected at a time when the beneficiary was believed not to be entitled to Medicare benefits but:

     (A) The beneficiary is later determined to have been entitled to Medicare benefits; and

     (B) The beneficiary's entitlement period falls within the time the rural health clinic's agreement with the Secretary is in effect.
§ 405.2410 Application of Part B deductible and coinsurance.

(a) Application of deductible. (1) Medicare payment for rural health clinic services begins only after the beneficiary has incurred the deductible.

(b) Application of coinsurance. (1) The beneficiary is responsible for a coinsurance amount which cannot exceed 20 percent of the clinic's reasonable customary charge for the covered service; and

(ii) Has undergone a change of ownership.

(2) Notice of termination. The Secretary will give notice of termination to the rural health clinic at least 15 days before the effective date stated in the notice.

(3) Appeal by the rural health clinic. A rural health clinic may appeal the termination of its agreement in accordance with the provisions set forth in part 498 of this chapter.

(c) Effect of termination. Payment will not be available for rural health clinic services furnished on or after the effective date of termination.

(d) Notice to the public. Prompt notice of the date and effect of termination shall be given to the public, through publication in local newspapers:

(i) By the clinic, after the Secretary has approved or set a termination date; or

(ii) By the Secretary, when he has terminated the agreement.

(e) Conditions for reinstatement after termination of agreement by the Secretary. When an agreement with a rural health clinic is terminated by the Secretary, the rural health clinic may not file another agreement to participate in the Medicare program unless the Secretary:

(i) Finds that the reason for the termination of the prior agreement has been removed; and

(ii) Is assured that the reason for the termination will not recur.

§ 405.2404 Terminations of agreements.

(a) Termination by rural health clinic. (1) Notice to Secretary. If the clinic wishes to terminate its agreement it shall file with the Secretary a written notice stating the intended effective date of termination.

(2) Action by the Secretary. (i) The Secretary may approve the date proposed by the clinic, or set a different date no later than 6 months after the date of the clinic's notice.

(ii) The Secretary may approve a date which is less than 6 months after the date of notice if he determines that termination on that date would not:

(A) Unduly disrupt the furnishing of services to the community serviced by the clinic; or

(B) Otherwise interfere with the efficient and effective administration of the Medicare program.

(3) Cessation of business. If a clinic ceases to furnish services to the community, that shall be deemed to be a voluntary termination of the agreement by the clinic, effective on the last day of business.

(b) Termination by the Secretary. (1) Cause for termination. The Secretary may terminate an agreement if he determines that the rural health clinic:

(i) No longer meets the conditions for certification under part 491 of this chapter; or

(ii) Is not in substantial compliance with the provisions of the agreement, the requirements of this subpart, any other applicable regulations of this part, or any applicable provisions of title XVIII of the Act; or

(iii) Has undergone a change of ownership.

(2) Notice of termination. The Secretary will give notice of termination to the rural health clinic at least 15 days before the effective date stated in the notice.

(3) Appeal by the rural health clinic. A rural health clinic may appeal the termination of its agreement in accordance with the provisions set forth in part 498 of this chapter.

(c) Effect of termination. Payment will not be available for rural health clinic services furnished on or after the effective date of termination.

(d) Notice to the public. Prompt notice of the date and effect of termination shall be given to the public, through publication in local newspapers:

(1) By the clinic, after the Secretary has approved or set a termination date; or

(2) By the Secretary, when he has terminated the agreement.

(e) Conditions for reinstatement after termination of agreement by the Secretary. When an agreement with a rural health clinic is terminated by the Secretary, the rural health clinic may not file another agreement to participate in the Medicare program unless the Secretary:

(1) Finds that the reason for the termination of the prior agreement has been removed; and

(2) Is assured that the reason for the termination will not recur.

any one item or service furnished by the rural health clinic, may not exceed a reasonable amount customarily charged by the clinic for that particular item or service.

(ii) For any one item or service furnished by a Federally qualified health center, the coinsurance liability may not exceed 20 percent of a reasonable amount customarily charged by the center for that particular item or service.

[57 FR 24976, June 12, 1992]

§ 405.2411 Scope of benefits.

(a) Rural health clinic services reimbursable under this subpart are:

(1) The physicians' services specified in §405.2412;

(2) Services and supplies furnished as an incident to a physician's professional service;

(3) The nurse practitioner or physician assistant services specified in §405.2414;

(4) Services and supplies furnished as an incident to a nurse practitioner’s or physician assistant’s services; and

(5) Visiting nurse services.

(b) Rural health clinic services are reimbursable when furnished to a patient at the clinic, at a hospital or other medical facility, or at the patient’s place of residence.

§ 405.2412 Physicians' services.

(a) Physicians' services are professional services that are performed by a physician at the clinic or are performed away from the clinic by a physician whose agreement with the clinic provides that he or she will be paid by the clinic for such services.

§ 405.2413 Services and supplies incident to a physician's services.

(a) Services and supplies incident to a physician's professional service are reimbursable under this subpart if the service or supply is:

(1) Of a type commonly furnished in physicians’ offices;

(2) Of a type commonly rendered either without charge or included in the rural health clinic's bill;

(3) Furnished as an incidental, although integral, part of a physician’s professional services;

(4) Furnished under the direct, personal supervision of a physician; and

(5) In the case of a service, furnished by a member of the clinic's health care staff who is an employee of the clinic.

(b) Only drugs and biologicals which cannot be self-administered are included within the scope of this benefit.

§ 405.2414 Nurse practitioner and physician assistant services.

(a) Professional services are reimbursable under this subpart if:

(1) Furnished by a nurse practitioner, physician assistant, nurse midwife, or specialized nurse practitioner who is employed by, or receives compensation from, the rural health clinic;

(2) Furnished under the medical supervision of a physician;

(3) Furnished in accordance with any medical orders for the care and treatment of a patient prepared by a physician;

(4) They are of a type which the nurse practitioner, physician assistant, nurse midwife or specialized nurse practitioner who furnished the service is legally permitted to perform by the State in which the service is rendered; and

(5) They would be covered if furnished by a physician.

(b) The physician supervision requirement is met if the conditions specified in §491.8(b) of this chapter and any pertinent requirements of State law are satisfied.

(c) The services of nurse practitioners, physician assistants, nurse midwives or specialized nurse practitioners are not covered if State law or regulations require that the services be performed under a physician's order and no such order was prepared.

§ 405.2415 Services and supplies incident to nurse practitioner and physician assistant services.

(a) Services and supplies incident to a nurse practitioner’s or physician assistant’s services are reimbursable under this subpart if the service or supply is:

(1) Of a type commonly furnished in physicians’ offices;

(2) Of a type commonly rendered either without charge or included in the rural health clinic’s bill;
§ 405.2430 Basic requirements.

(a) Filing procedures.

(1) In response to a request from an entity that wishes to participate in the Medicare program, HCFA enters into an agreement with an entity when—

(3) Furnished as an incidental, although integral part of professional services furnished by a nurse practitioner, physician assistant, nurse midwife, or specialized nurse practitioner;

(4) Furnished under the direct, personal supervision of a nurse practitioner, physician assistant, nurse midwife, specialized nurse practitioner or a physician; and

(5) In the case of a service, furnished by a member of the clinic’s health care staff who is an employee of the clinic.

(b) The direct personal supervision requirement is met in the case of a nurse practitioner, physician assistant, nurse midwife, or specialized nurse practitioner only if such a person is permitted to supervise such services under the written policies governing the rural health clinic.

(c) Only drugs and biologicals which cannot be self-administered are included within the scope of this benefit.

§ 405.2416 Visiting nurse services.

(a) Visiting nurse services are covered if:

(1) The rural health clinic is located in an area in which the Secretary has determined that there is a shortage of home health agencies;

(2) The services are rendered to a homebound individual;

(3) The services are furnished by a registered nurse, licensed practical nurse, or licensed vocational nurse who is employed by, or receives compensation for the services from the clinic; and

(4) The services are furnished under a written plan of treatment that is:

(i) Established and reviewed at least every 60 days by a supervising physician of the rural health clinic or established by a nurse practitioner, physician assistant, nurse midwife, or specialized nurse practitioner and reviewed at least every 60 days by a supervising physician;

(ii) Signed by the nurse practitioner, physician assistant, nurse midwife, specialized nurse practitioner, or the supervising physician of the clinic.

(b) The nursing care covered by this section includes:

(1) Services that must be performed by a registered nurse, licensed practical nurse, or licensed vocational nurse if the safety of the patient is to be assured and the medically desired results achieved; and

(2) Personal care services, to the extent covered under Medicare as home health services. These services include helping the patient to bathe, to get in and out of bed, to exercise and to take medications.

(c) This benefit does not cover household and housekeeping services or other services that would constitute custodial care.

(d) For purposes of this section, homebound means an individual who is permanently or temporarily confined to his or her place of residence because of a medical or health condition. The individual may be considered homebound if he or she leaves the place of residence infrequently. For this purpose, “place of residence” does not include a hospital or long term care facility.

§ 405.2417 Visiting nurse services: Determination of shortage of agencies.

A shortage of home health agencies exists if the Secretary determines that the rural health clinic:

(a) Is located in a county, parish, or similar geographic area in which there is no participating home health agency or adequate home health services are not available to patients of the rural health clinic;

(b) Has (or expects to have) patients whose permanent residences are not within the area serviced by a participating home health agency; or

(c) Has (or expects to have) patients whose permanent residences are not within a reasonable traveling distance, based on climate and terrain, of a participating home health agency.

FEDERALLY QUALIFIED HEALTH CENTER SERVICES

SOURCE: 57 FR 24978, June 12, 1992, unless otherwise noted.

§ 405.2430 Basic requirements.

(a) Filing procedures. (1) In response to a request from an entity that wishes to participate in the Medicare program, HCFA enters into an agreement with an entity when—
§ 405.2434

(i) PHS recommends that the entity qualifies as a Federally qualified health center;

(ii) The Federally qualified health center assures HCFA that it meets the Federally qualified health center requirements specified in this subpart and part 491, as described in § 405.2434(a); and

(iii) The FQHC terminates other provider agreements, unless the FQHC assures HCFA that it is not using the same space, staff and resources simultaneously as a physician’s office or another type of provider or supplier. A corporate entity may own other provider types as long as the provider types are distinct from the FQHC.

(2) HCFA sends the entity a written notice of the disposition of the request.

(3) When the requirement of paragraph (a)(1) of this section is satisfied, HCFA sends the entity two copies of the agreement. The entity must sign and return both copies of the agreement to HCFA.

(4) If HCFA accepts the agreement filed by the Federally qualified health center, HCFA returns to the center one copy of the agreement with the notice of acceptance specifying the effective date (see § 405.2434).

(b) Recommendations by PHS about Federally qualified health centers.

(1) An entity must—

(i) Meet the applicable requirements of the PHS Act, as specified in § 405.2401(b); and

(ii) Be recommended by PHS to HCFA as a Federally qualified health center.

(2) The PHS notifies HCFA of entities that meet the requirements specified in § 405.2401(b).

(c) Provider-based and freestanding Federally qualified health centers. The requirements and benefits under Medicare for provider-based or freestanding Federally qualified health centers are the same, except that payment methodologies differ, as described in § 405.2462.

(d) Appeals. An entity is entitled to a hearing in accordance with part 498 of this chapter when HCFA fails to enter into an agreement with the entity.

§ 405.2434

Under the agreement, the Federally qualified health center must agree to the following:

(a) Maintain compliance with the requirements. (1) The Federally qualified health center must agree to maintain compliance with the Federally qualified health center requirements set forth in this subpart and part 491, except that the provisions of § 491.3 do not apply.

(2) Centers must promptly report to HCFA any changes that result in non-compliance with any of these requirements.

(b) Effective date of agreement. (1) Except as specified in paragraph (b)(2) of this section, the effective date of the agreement is the date HCFA accepts the signed agreement, which assures that all Federal requirements are met. (2) For facilities that met all requirements on October 1, 1991, the effective date of the agreement can be October 1, 1991.

(c) Charges to beneficiaries. (1) The beneficiary is responsible for payment of a coinsurance amount which is 20 percent of the amount of Part B payment made to the Federally qualified health center for the covered services. There is no coinsurance for a second or third opinion obtained in accordance with section 1164 of the Act or for pneumococcal vaccine and its administration. (2) The beneficiary is responsible for blood deductible expenses, as specified in § 410.161.

(3) The Federally qualified health center agrees not to charge the beneficiary (or any other person acting on behalf of a beneficiary) for any Federally qualified health center services for which the beneficiary is entitled to have payment made on his or her behalf by the Medicare program (or for which the beneficiary would have been entitled if the Federally qualified health center had filed a request for payment in accordance with § 410.165 of this chapter), except for coinsurance amounts.

(4) The Federally qualified health center may charge the beneficiary for items and services that are not Federally qualified health center services.

[57 FR 24078, June 12, 1992, as amended at 61 FR 14657, Apr. 3, 1996]
However, if the item or service is covered under Part B of Medicare, and the Federally qualified health center agrees to receive Part B payment under the assignment method, the Federally qualified health center may not charge the beneficiary more than 20 percent of the Part B payment.

(d) Refunds to beneficiaries. (1) The Federally qualified health center must agree to refund as promptly as possible any money incorrectly collected from Medicare beneficiaries or from someone on their behalf.

(2) As used in this section, “money incorrectly collected” means any amount for covered services that is greater than the amount for which the beneficiary was liable because of the coinsurance requirements specified in part 410, subpart E.

(3) Amounts also are considered incorrectly collected if the Federally qualified health center believed the beneficiary was not entitled to Medicare benefits but—

(i) The beneficiary was later determined to have been so entitled;

(ii) The beneficiary’s entitlement period fell within the time the Federally qualified health center’s agreement with HCFA was in effect; and

(iii) The amounts exceed the beneficiary’s coinsurance liability.

(e) Treatment of beneficiaries. (1) The Federally qualified health center must agree to accept Medicare beneficiaries for care and treatment.

(2) The Federally qualified health center may not impose any limitations with respect to care and treatment of Medicare beneficiaries that it does not also impose upon all other persons seeking care and treatment from the Federally qualified health center. Failure to comply with this requirement is a cause for termination of the Federally qualified health center’s agreement with HCFA in accordance with §405.2436(d).

(3) If the Federally qualified health center does not furnish treatment for certain illnesses and conditions to patients who are not Medicare beneficiaries, it need not furnish such treatment to Medicare beneficiaries.

§ 405.2436 Termination of agreement.

(a) Termination by Federally qualified health center. The Federally qualified health center may terminate its agreement by—

(1) Filing with HCFA a written notice stating its intention to terminate the agreement; and

(2) Notifying HCFA of the date on which the Federally qualified health center requests that the termination take effect.

(b) Effective date. (1) Upon receiving a Federally qualified health center’s notice of intention to terminate the agreement, HCFA will set a date upon which the termination takes effect. This effective date may be—

(i) The date proposed by the Federally qualified health center in its notice of intention to terminate, if that date is acceptable to HCFA; or

(ii) Except as specified in paragraph (2) of this section, a date set by HCFA, which is no later than 6 months after the date HCFA receives the Federally qualified health center’s notice of intention to terminate.

(2) The effective date of termination may be less than 6 months following HCFA’s receipt of the Federally qualified health center’s notice of intention to terminate if HCFA determines that termination on such a date would not—

(i) Unduly disrupt the furnishing of Federally qualified health center services to the community; or

(ii) Otherwise interfere with the effective and efficient administration of the Medicare program.

(3) The termination is effective at the end of the last day of business as a Federally qualified health center.

(c) Termination by HCFA. (1) HCFA may terminate an agreement with a Federally qualified health center—

(i) No longer meets the requirements specified in this subpart; or

(ii) Is not in substantial compliance with—

(A) The provisions of the agreement; or

(B) The requirements of this subpart, any other applicable regulations of this part, or any applicable provisions of title XVIII of the Act.
§ 405.2440 Conditions for reinstatement after termination by HCFA.

When HCFA has terminated an agreement with a Federally qualified health center, HCFA will not enter into another agreement with the Federally qualified health center to participate in the Medicare program unless HCFA—

(a) Finds that the reason for the termination no longer exists; and

(b) Is assured that the reason for the termination of the prior agreement will not recur.

§ 405.2442 Notice to the public.

(a) When the Federally qualified health center voluntarily terminates the agreement and an effective date is set for the termination, the Federally qualified health center must notify the public prior to a prospective effective date or on the actual day that business ceases, if no prospective date of termination has been set, through publication in at least one newspaper in general circulation in the area serviced by the Federally qualified health center.

(b) When HCFA terminates the agreement, HCFA will notify the public by publication in at least one newspaper in general circulation in the Federally qualified health center's service area.

§ 405.2444 Change of ownership.

(a) What constitutes change of ownership—(1) Incorporation. The incorporation of an unincorporated FQHC constitutes change of ownership.

(2) Merger. The merger of the center corporation into another corporation, or the consolidation of two or more corporations, one of which is the center corporation, resulting in the creation of a new corporation, constitutes a change of ownership. (The merger of another corporation into the center corporation does not constitute change of ownership.)

(3) Leasing. The lease of all or part of an entity constitutes a change of ownership of the leased portion.

(b) Notice to HCFA. A center which is contemplating or negotiating change of ownership must notify HCFA.

(c) Assignment of agreement. When there is a change of ownership as specified in paragraph (a) of this section, the agreement with the existing center is automatically assigned to the new owner if it continues to meet the conditions to be a Federally qualified health center.

(d) Conditions that apply to assigned agreements. An assigned agreement is subject to all applicable statutes and regulations and to the terms and conditions under which it was originally issued including, but not limited to, the following:

(1) Compliance with applicable health and safety standards.

(2) Compliance with the ownership and financial interest disclosure requirements of part 420, subpart C of this subchapter.

§ 405.2446 Scope of services.

(a) For purposes of this section, the terms rural health clinic and clinic when they appear in the cross references in paragraph (b) of this section also mean Federally qualified health centers.

(b) FQHC services that are paid for under this subpart are outpatient services that include the following:

(1) Physician services specified in § 405.2412.

(2) Services and supplies furnished as an incident to a physician's professional services, as specified in § 405.2413.

(3) Nurse practitioner or physician assistant services specified in § 405.2414.
(4) Services and supplies furnished as an incident to a nurse practitioner or physician assistant services, as specified in §405.2415.

(5) Clinical psychologist and clinical social worker services specified in §405.2450.

(6) Services and supplies furnished as an incident to a clinical psychologist or clinical social worker services, as specified in §405.2452.

(7) Visiting nurse services specified in §405.2416.

(8) Nurse-midwife services specified in §405.2401.

(9) Preventive primary services specified in §405.2448 of this subpart.

(c) Federally qualified health center services are covered when provided in outpatient settings only, including a patient’s place of residence, which may be a skilled nursing facility or a nursing facility or other institution used as a patient’s home.

(d) Federally qualified health center services are not covered in a hospital, as defined in section 1861(e)(1) of the Act.

§405.2448 Preventive primary services.

(a) Preventive primary services are those health services that—

(1) A center is required to provide as preventive primary health services under section 329, 330, and 340 of the Public Health Service Act;

(2) Are furnished by or under the direct supervision of a nurse practitioner, physician assistant, nurse midwife, specialized nurse practitioner, clinical psychologist, clinical social worker, or a physician;

(3) In the case of a service, are furnished by a member of the center’s health care staff who is an employee of the center or by a physician under arrangements with the center; and

(4) Except as specifically provided in section 1861(s) of the Act, include only drugs and biologicals that cannot be self-administered.

(b) Preventive primary services which may be paid for when provided by Federally qualified health centers are the following:

(1) Medical social services.

(2) Nutritional assessment and referral.

(3) Preventive health education.

(4) Children’s eye and ear examinations.

(5) Prenatal and post-partum care.

(6) Perinatal services.

(7) Well child care, including periodic screening.

(8) Immunizations, including tetanus-diphtheria booster and influenza vaccine.

(9) Voluntary family planning services.

(10) Taking patient history.

(11) Blood pressure measurement.

(12) Weight.

(13) Physical examination targeted to risk.

(14) Visual acuity screening.

(15) Hearing screening.

(16) Cholesterol screening.

(17) Stool testing for occult blood.

(18) Dipstick urinalysis.

(19) Risk assessment and initial counseling regarding risks.

(20) Tuberculosis testing for high risk patients.

(21) For women only.

(i) Clinical breast exam.

(ii) Referral for mammography; and

(iii) Thyroid function test.

(c) Preventive primary services do not include group or mass information programs, health education classes, or group education activities, including media productions and publications.

(d) Screening mammography is not considered a Federally qualified health center service, but may be provided at a Federally qualified health center if the center meets the requirements applicable to that service specified in §410.34 of this subchapter. Payment is made under applicable Medicare requirements.

(e) Preventive primary services do not include eyeglasses, hearing aids, or preventive dental services.

§405.2450 Clinical psychologist and clinical social worker services.

(a) For clinical psychologist or clinical social worker professional services to be payable under this subpart, the services must be—

[57 FR 24979, June 12, 1992, as amended at 61 FR 14657, Apr. 3, 1996]
§ 405.2452 Services and supplies incident to clinical psychologist and clinical social worker services.

(a) Services and supplies incident to a clinical psychologist’s or clinical social worker’s services are reimbursable under this subpart if the service or supply is—

(1) Of a type commonly furnished in a physician’s office;

(2) Of a type commonly furnished either without charge or included in the Federally qualified health center’s bill;

(3) Furnished as an incidental, although integral part of professional services furnished by a clinical psychologist or clinical social worker;

(4) Furnished under the direct, personal supervision of a clinical psychologist, clinical social worker or physician; and

(5) In the case of a service, furnished by a member of the center’s health care staff who is an employee of the center.

(b) The direct personal supervision requirement in paragraph (a)(4) of this section is met only if the clinical psychologist or clinical social worker is permitted to supervise such services under the written policies governing the Federally qualified health center.

Payment for Rural Health Clinic and Federally Qualified Health Center Services

§ 405.2460 Applicability of general payment exclusions.

The payment conditions, limitations, and exclusions set out in subpart C of this part, part 410 and part 411 of this chapter are applicable to payment for services provided by rural health clinics and Federally qualified health centers, except that preventive primary services, as defined in § 405.2448, are covered in Federally qualified health centers and not excluded by the provisions of section 1862(a) of the Act.

§ 405.2462 Payment for rural health clinic and Federally qualified health center services.

(a) Payment to provider-based rural health clinics and Federally qualified health centers. A rural health clinic or Federally qualified health center is paid in accordance with parts 405 and 413 of this subchapter, as applicable, if:

(1) The clinic or center is an integral and subordinate part of a hospital, skilled nursing facility or home health agency participating in Medicare (i.e., a provider of services); and

(2) The clinic or center is operated with other departments of the provider under common licensure, governance and professional supervision.

(b) Payment to independent rural health clinics and freestanding Federally qualified health centers. (1) All other clinics and centers will be paid on the basis of an all-inclusive rate for each beneficiary visit for covered services. This rate will be determined by the intermediary, in accordance with this subpart and general instructions issued by HCFA.

(2) The amount payable by the intermediary for a visit will be determined in accordance with paragraph (b)(3) and (4) of this section.

(3) Federally qualified health centers. For Federally qualified health center
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§ 405.2466 All-inclusive rate.

(a) Determination of rate. (1) An all-inclusive rate is determined by the intermediary at the beginning of the reporting period.

(2) The rate is determined by dividing the estimated total allowable costs by estimated total visits for rural health clinic or Federally qualified health center services.

(3) The rate determination is subject to any tests of reasonableness that may be established in accordance with this subpart.

(b) Adjustment of rate. (1) The intermediary, during each reporting period, periodically reviews the rate to assure that payments approximate actual allowable costs and visits for rural health clinic or Federally qualified health center services.

(i) There is a significant change in the utilization of clinic or center services;

(ii) Actual allowable costs vary materially from the clinic or center’s allowable costs; or

(iii) Other circumstances arise which warrant an adjustment.

(2) The clinic or center may request the intermediary to review the rate to determine whether adjustment is required.

§ 405.2466 Annual reconciliation.

(a) General. Payments made to a rural health clinic or a Federally qualified health center during a reporting period are subject to reconciliation to
§ 405.2468 Allowable costs.

(a) Applicability of general Medicare principles. In determining whether and to what extent a specific type or item of cost is allowable, such as interest, depreciation, bad debts and owner compensation, the intermediary applies the principles for reimbursement of provider costs, as set forth in part 413 of this subchapter.

(b) Typical rural health clinic and Federally qualified health center costs. The following types and items of cost are included in allowable costs to the extent that they are covered and reasonable:

(1) Compensation for the services of a physician, physician assistant, nurse practitioner, nurse-midwife, visiting nurse, qualified clinical psychologist,
and clinical social worker who owns, is employed by, or furnishes services under contract to an FQHC. (RHCs are not paid for services furnished by contracted individuals other than physicians.)

(2) Compensation for the duties that a supervising physician is required to perform under the agreement specified in §491.8 of this chapter.

(3) Costs of services and supplies incident to the services of a physician, physician assistant, nurse practitioner, nurse-midwife, qualified clinical psychologist, or clinical social worker.

(4) Overhead costs, including clinic or center administration, costs applicable to use and maintenance of the entity, and depreciation costs.

(5) Costs of services purchased by the clinic or center.

(c) Tests of reasonableness for rural health clinic cost and utilization. Tests of reasonableness authorized by sections 1833(a) and 1861(v)(1)(A) of the Act may be established by HCFA or the carrier with respect to direct or indirect overall costs, costs of specific items and services, or costs of groups of items and services. Those tests include, but are not limited to, screening guidelines and payment limitations.

(d) Screening guidelines. (1) Costs in excess of amounts established by the guidelines are not included unless the clinic or center provides reasonable justification satisfactory to the intermediary.

(2) Screening guidelines are used to assess the costs of services, including the following:

(i) Compensation for the professional and supervisory services of physicians and for the services of physician assistants, nurse practitioners, and nurse-midwives.

(ii) Services of physicians, physician assistants, nurse practitioners, nurse-midwives, visiting nurses, qualified clinical psychologists, and clinical social workers.

(iii) The level of administrative and general expenses.

(iv) Staffing (for example, the ratio of other clinic or center personnel to physicians, physician assistants, and nurse practitioners).

(v) The reasonableness of payments for services purchased by the clinic or center, subject to the limitation that the costs of physician services purchased by the clinic or center may not exceed amounts determined under the applicable provisions of subpart E of part 405 or part 415 of this chapter.

(e) Payment limitations. Limits on payments may be set by HCFA, on the basis of costs estimated to be reasonable for the provision of such services.

(f) Graduate medical education. (1) Effective for that portion of cost reporting periods occurring on or after January 1, 1999, if an RHC or an FQHC incurs “all or substantially all” of the costs for the training program in the nonhospital setting as defined in §413.86(b) of this chapter, the RHC or FQHC may receive direct graduate medical education payment for those residents.

(2) Direct graduate medical education costs are not included as allowable cost under §405.2466(b)(1)(i); and therefore, are not subject to the limit on the all-inclusive rate for allowable costs.

(3) Allowable graduate medical education costs must be reported on the RHC’s or the FQHC’s cost report under a separate cost center.

(4) Allowable graduate medical education costs are non-reimbursable if payment for these costs are received from a hospital or a Medicare+Choice organization.

(5) Allowable direct graduate medical education costs under paragraphs (f)(6) and (f)(7)(i) of this section, are subject to reasonable cost principles under part 413 and the reasonable compensation equivalency limits in §§415.60 and 415.70 of this chapter.

(6) The allowable direct graduate medical education costs are those costs incurred by the nonhospital site for the educational activities associated with patient care services of an approved program, subject to the redistribution and community support principles in §413.85(c).

(i) The following costs are allowable direct graduate medical education costs to the extent that they are reasonable—

(A) The costs of the residents’ salaries and fringe benefits (including travel and lodging expenses where applicable).
§ 405.2470 Reports and maintenance of records.

(a) Maintenance and availability of records. The rural health clinic or Federally qualified health center must:

(1) Maintain adequate financial and statistical records, in the form and containing the data required by HCFA, to allow the intermediary to determine payment for covered services furnished to Medicare beneficiaries in accordance with this subpart;

(2) Make the records available for verification and audit by HHS or the General Accounting Office;

(3) Maintain financial data on an accrual basis, unless it is part of a governmental institution that uses a cash basis of accounting. In the latter case, appropriate depreciation on capital assets is allowable rather than the expenditure for the capital asset.

(b) Adequacy of records. (1) The intermediary may suspend reimbursement if it determines that the clinic or center does not maintain records that provide an adequate basis to determine payments under Medicare.

(2) The suspension continues until the clinic or center demonstrates to the intermediary's satisfaction that it does, and will continue to, maintain adequate records.

(c) Reporting requirements—(1) Initial report. At the beginning of its initial reporting period, the clinic or center must submit an estimate of budgeted costs and visits for rural health clinic or Federally qualified health center services for the reporting period, in the form and detail required by HCFA, and such other information as HCFA may require to establish the payment rate.

(2) Annual reports. Within 90 days after the end of its reporting period, the clinic or center must submit, in such form and detail as may be required by HCFA, a report of:

(i) Its operations, including the allowable costs actually incurred for the period and the actual number of visits for rural health clinic or Federally qualified health center services furnished during the period; and

(ii) The estimated costs and visits for rural health clinic services or Federally qualified health center services for the succeeding reporting period and such other information as HCFA may require to establish the payment rate.

(3) Late reports. If the clinic or center does not submit an adequate annual report on time, the intermediary may reduce or suspend payments to preclude excess payment to the clinic or center.

(4) Inadequate reports. If the clinic or center does not furnish a report or furnishes a report that is inadequate for the intermediary to make a determination of program payment, HCFA may deem all payments for the reporting period to be overpayments.

(5) Postponement of due date. For good cause shown by the clinic or center, the intermediary may, with HCFA's approval, grant a 30-day postponement of the due date for the annual report.
Health Care Financing Administration, HHS § 406.2

(6) Reports following termination of agreement or change of ownership. The report from a clinic or center which voluntarily or involuntarily ceases to participate in the Medicare program or experiences a change in ownership (see §§ 405.2436-405.2438) is due no later than 45 days following the effective date of the termination of agreement or change of ownership.

§ 405.2472 Beneficiary appeals.

A beneficiary may request a hearing by an intermediary (subject to the limitations and conditions set forth in subpart H of this part) if:

(a) The beneficiary is dissatisfied with an intermediary’s determination denying a request for payment made on his or her behalf by a rural health clinic or Federally qualified health center; or

(b) The beneficiary is dissatisfied with the amount of payment; or

(c) The beneficiary believes the request for payment is not being acted upon with reasonable promptness.


PART 406—HOSPITAL INSURANCE ELIGIBILITY AND ENTITLEMENT

Subpart A—General Provisions

§ 406.1 Statutory basis.

Sections 226, 226A, 1818 and 1818A of the Social Security Act and section 103 of Public Law 89-97 establish the conditions for entitlement to hospital insurance benefits. Sections 202 (t) and (u) of the Act specify limitations that apply to certain aliens and to persons convicted of certain offenses.

[48 FR 15236, Mar. 25, 1983, unless otherwise noted. Redesignated at 51 FR 41338, Nov. 14, 1986]

§ 406.2 Scope.

Subparts A through D of this part specify the conditions of eligibility for hospital insurance and set forth certain specific conditions that affect entitlement to benefits. Hospital insurance is authorized under Part A of title XVIII and is also referred to as Medicare Part A. It includes inpatient hospital care, posthospital SNF care, home health services, and hospice care.

§ 406.3 Definitions.

First month of eligibility means the first month in which an individual meets all the requirements for entitlement to hospital insurance except application or enrollment if that is required.

First month of entitlement means the first month for which the individual meets all the requirements for entitlement to Part A benefits.

Insured individual means an individual who has the number of quarters of coverage required for monthly social security benefits.

Quarter of coverage means a calendar quarter that is counted toward the number of covered quarters required to make the individual eligible for monthly social security benefits. A quarter is counted if during that quarter (or that calendar year) the individual earned a required minimum amount of money. (For details, see 20 CFR part 404, subpart B.)

§ 406.5 Basis of eligibility and entitlement.

(a) Hospital insurance without premiums. Hospital insurance is available to most individuals without payment of a premium if they:

(1) Are age 65 or over,

(2) Have received social security or railroad retirement disability benefits for 25 months; or

(3) Have end-stage renal disease. Subpart B of this part explains the requirements such individuals must meet to obtain hospital insurance without premiums.

(b) Premium hospital insurance. Many individuals who are age 65 or over, but do not meet the requirements set forth in subpart B of this part, and certain individuals under age 65, may obtain the benefits by paying a premium. Section 406.20 of this part explains the requirements such individuals must meet to obtain premium hospital insurance.


§ 406.6 Application or enrollment for hospital insurance.

(a) Basic provision. In most cases, eligibility for Medicare Part A is a result of entitlement to monthly social security or railroad retirement cash benefits or eligibility for monthly social security cash benefits. This section specifies the individuals who need not file an application to become entitled to hospital insurance, those who must file an application, and those who must enroll.

(b) Individuals who need not file an application for hospital insurance. An individual who meets any of the following conditions need not file an application for hospital insurance:

(1) Is under age 65 and has been entitled, for more than 24 months, to monthly social security or railroad retirement benefits based on disability.

(2) At the time of attainment of age 65, is entitled to monthly social security or railroad retirement benefits.

(3) Establishes entitlement to monthly social security or railroad retirement benefits at any time after attaining age 65.

(c) Individuals who must file an application for hospital insurance. An individual must file an application for hospital insurance if he or she seeks entitlement to hospital insurance on the basis of—

(1) The transitional provisions set forth in §406.11;

(2) Deemed entitlement to disabled widow’s or widower’s benefit under certain circumstances as provided in §406.12;

(3) A diagnosis of end-stage renal disease, as specified in §406.13;

(4) Effective January 1, 1981, eligibility for social security cash benefits, as specified in §406.10(a)(3), if the individual has attained age 65 without applying for those benefits; or

(5) The special provisions applicable to government employment as set forth in §406.15.

(d) When application is deemed to be filed. (1) An application based on the transitional provisions or on ESRD is deemed to be filed in the first month of eligibility if it is filed not more than 3 months before the first month, and is retroactive to that month if filed within 12 months after the first month. An application filed more than 12 months after the first month of eligibility is retroactive to the 12th month before the month it is filed.
§ 406.10

(2) An application for deemed entitlement to disabled widow's or widower's benefits, that is filed before the first month in which the individual meets all conditions of entitlement for this benefit, will be deemed a valid application if those conditions are met before an initial determination, reconsideration, or hearing decision is made on the application. If the conditions are met after the date of any hearing decision, a new application will have to be filed. An application validly filed within 12 months after the first month of eligibility is retroactive to that first month. If filed more than 12 months after that first month, it is retroactive to the 12th month before the month of filing.

(3) Effective June 8, 1980, an application based on eligibility for social security benefits at or after age 65, that is filed before the first month in which the individual meets all eligibility conditions for this benefit, will be deemed a valid application if those conditions are met before an initial determination, reconsideration, or hearing decision is made on the application. If the conditions are met after the date of any hearing decision, a new application will have to be filed.

(4) Effective March 1, 1981, an application under § 406.10 that is validly filed within 6 months after the first month of eligibility is retroactive to that first month. If filed more than 6 months after that first month, it is retroactive to the 8th month before the month of filing.

(e) Individuals who must enroll for hospital insurance. An individual who must pay a monthly premium for hospital insurance must enroll in accordance with the procedures set forth in § 406.21.

§ 406.7 Forms to apply for entitlement under Medicare Part A

The following forms, available free of charge by mail from HCFA or at any Social Security branch or district office, are used to apply for Medicare entitlement under the circumstances indicated:

HCFA-18-F-5—Application for Hospital Insurance Entitlement. (For use by individuals who are not eligible for retirement benefits under Title II of the Social Security Act or under the Railroad Retirement Act. This form may also be used for enrollment in the supplementary medical insurance program.)

HCFA-43—Application for Health Insurance Benefits under Medicare for Individuals with End Stage Renal Disease (ESRD). (An initial application for entitlement by individuals with ESRD).

As an alternative, an individual may use the application for monthly social benefits to apply also for Medicare entitlement if he or she is eligible for hospital insurance at that time.

[53 FR 6633, Mar. 2, 1988]

Subpart B—Hospital Insurance Without Monthly Premiums

§ 406.10 Individual age 65 or over who is entitled to social security or railroad retirement benefits, or who is eligible for social security benefits.

(a) Requirements. An individual is entitled to hospital insurance benefits under section 226 of the Act if he or she has attained aged 65 and is:

(1) Entitled to monthly social security benefits under section 202 of the Social Security Act;

(2) A qualified railroad retirement beneficiary who has been certified as such to the Social Security Administration by the Railroad Retirement Board in accordance with section 7(d) of the Railroad Retirement Act of 1974; or

(3) Effective January 1, 1981, eligible for monthly social security benefits under section 202 of the Act and has filed an application for hospital insurance.

(b) Beginning and end of entitlement.

(1) Entitlement begins with the first day of the first month in which the individual meets the requirements of paragraph (a) of this section.

(2) Entitlement continues until the individual dies or no longer meets the requirements of paragraph (a) of this section. An individual is not entitled to railroad retirement benefits and is neither entitled to, nor eligible for, monthly social security benefits in the month in which he or she dies. However, an individual who meets all other
requirements for hospital insurance entitlement is entitled to hospital insurance in the month in which he or she dies if he or she—

(i) Would have been entitled to monthly railroad retirement benefits or social security benefits in that month if he or she had not died; or

(ii) Has filed an application for hospital insurance and would have been eligible for monthly social security benefits in that month if he or she had not died.

§ 406.11 Individual age 65 or over who is not eligible as a social security or railroad retirement benefits beneficiary, or on the basis of government employment.

(a) Basis. Section 103 of the law that established the Medicare program in 1965 (Pub. L. 89-97) provided for eligibility for certain individuals who were age 65 or would soon attain age 65 but would not be able to qualify for social security or railroad retirement benefits.

(b) Requirements. Unless he or she is excluded under paragraph (c) of this section, an individual age 65 or over who does not meet the requirements of § 406.10 or § 406.15 (and who would not meet those requirements if he or she filed an application), is entitled to Medicare Part A benefits if he or she meets the following requirements:

(1) Age and quarters of coverage. (i) He or she attained age 65 before 1968; or

(ii) If he or she attained age 65 in 1968 or later, he or she must have at least 3 quarters of coverage for each year that elapsed after 1966 and before the year in which he or she attained age 65. (The quarters of coverage may have been acquired at any time, not necessarily during the elapsed years.)

(2) Residence and citizenship. He or she is a resident of the United States and—

(i) A citizen of the United States; or

(ii) An alien lawfully admitted for permanent residence who has continuously resided in the United States for 5 years immediately preceding the first month in which he or she meets all other requirements for entitlement to hospital insurance.

(3) Application. He or she has filed an application for Medicare Part A no earlier than the third month before the first month of eligibility.

(c) Bases for exclusion. An individual who meets the requirements of paragraph (b) of this section is excluded from Medicare Part A if he or she—

(1) Has been convicted of spying, sabotage, or treason, sedition, and subversive action under chapter 37, 105, or 115 of title 18 of the United States Code;

(2) Has been convicted of conspiracy to establish a dictatorship under section 4 of the Internal Security Act of 1950;

(3) On February 16, 1965, was or could have been covered under the Federal Employees Health Benefits Act (F E H B A) of 1959; or

(4) In his or her first month of eligibility,

(i) Is covered by an enrollment under the F E H B A; or

(ii) Could have been covered by an enrollment under that Act if he or she (or any other person who could provide him or her with coverage) was a Federal employee at any time after February 15, 1965, and had enrolled and retained coverage under that Act.

(d) End of exclusion. An individual excluded under paragraph (c)(3) or (4) of this section can become entitled beginning with the first month in which he or she loses the right to F E H B A coverage solely because he or she or the other person leaves Federal employment.

(e) Beginning and end of entitlement. (1) Entitlement begins—

(i) In the first month of eligibility if the application is filed no later than 12 months after the first month of eligibility:

(ii) In the 12th month before the month of application if the application is filed more than 12 months after the first month of eligibility.

(2) Entitlement continues until death or until the month before the month in which the individual becomes entitled under § 406.10 or § 406.15.

§ 406.12 Individual under age 65 who is entitled to social security or railroad retirement disability benefits.

(a) Basic requirements. An individual under age 65 is entitled to hospital insurance benefits if, for 25 months, he or she has been—

(1) Entitled or deemed entitled to social security disability benefits as an insured individual, child, widow, or widower who is “under a disability” or

(2) A disabled qualified beneficiary certified under Section 7(d) of the Railroad Retirement Act.

(b) Previous periods of disability benefits entitlement. Months of a previous period of entitlement or deemed entitlement to disability benefits count toward the 25-month requirement if any of the following conditions is met:

(1) Entitlement was as an insured individual or a disabled qualified railroad retirement beneficiary, and the previous period ended within the 60 months preceding the month in which the current disability began.

(2) Entitlement was as a disabled child, widow, or widower, and the previous period ended within the 84 months preceding the month in which the current disability began.

(3) The previous period ended on or after March 1, 1988 and the current impairment is the same as, or directly related to, the impairment on which the previous period of entitlement was based.

(c) Deemed entitlement to disabled widow’s or widower’s monthly benefits.

(1) Purpose. The provisions of paragraphs (c) (2), (3), and (4) of this section are intended to enable individuals—

(i) To meet the 25-month requirement of paragraph (a) of this section;

(ii) To retain hospital insurance entitlement when they are no longer entitled to monthly disability benefits.

(2) Deemed entitlement for certain individuals entitled to old-age insurance benefits. An individual who becomes entitled to monthly old-age insurance benefits before age 65, is, by law, precluded from establishing or retaining entitlement to disabled widow’s or widower’s monthly benefits. However, for purposes of meeting the 25-month requirement, a widow or widower who meets all other requirements for disability benefits and is excluded solely because of entitlement to old-age insurance benefits, shall be deemed to be (or to continue to be) entitled to disability benefits. A widow or widower who is not entitled to disability benefits for the month before attaining age 60 must file two applications, one for old-age insurance benefits and one for hospital insurance.

(3) Deemed entitlement for certain individuals entitled to mother’s benefits. An individual entitled to mother’s insurance benefits under section 202(g) of the Social Security Act cannot at the same time be entitled to disabled widow’s benefits. However, if she applies for hospital insurance, she shall be deemed to be entitled to disabled widow’s monthly benefits in the first month (of the 12 months before application) in which she would have been entitled to those benefits if she had filed an application for them.

(4) Deemed entitlement for certain individuals entitled to father’s benefits. An individual who is entitled to father’s insurance benefits under section 202(g) of the Act cannot at the same time be entitled to disabled widow’s benefits. However, if he applies for hospital insurance benefits, he shall be deemed to be entitled to disabled widow’s monthly benefits as follows:

(i) If he applied for hospital insurance benefits before May 1984, he was deemed entitled to disabled widow’s benefits for any month after April 1981 for which he would have been entitled to those benefits if he had filed an application for them.

(ii) If he applies for hospital insurance benefits in or after May 1984, he is deemed entitled to disabled widow’s benefits for any month, up to 12 months before the month of application, for which he would have been entitled to those benefits if he had filed an application for them.

(iii) Hospital insurance entitlement under this paragraph (c)(4) could not begin before May 1983.

(5) Deemed retroactive entitlement for certain disabled widows and widowers. In some cases, disabled widows or widowers cannot become entitled to monthly cash benefits before the month in which they file application. However, for purposes of meeting the
25-month requirement, disability benefit entitlement will be deemed to have begun with the earliest month (of the 12 months before the application for cash benefits) in which the individual met all the requirements except the filing of an application. (This provision is effective for applications filed on or after January 1, 1978.)

(d) When entitlement begins and ends.
(1) Entitlement to hospital insurance begins with the 25th month of an individual’s entitlement or deemed entitlement to disability benefits. Although an individual is not entitled to disability benefits for the month in which he or she dies, for purposes of this paragraph the individual will be deemed to be entitled for the month of death.

(2) Except as provided in paragraph (e) of this section, entitlement to hospital insurance ends with the earliest of the following:
   (i) The last day of the last month in which he or she was entitled or deemed entitled to disability benefits or was qualified as a disabled railroad retirement beneficiary, if he or she was notified of the termination of entitlement before that month.
   (ii) The last day of the month in which he or she is mailed a notice that his or her entitlement or deemed entitlement to disability benefits, or his or her status as a qualified disabled railroad retirement beneficiary, has ended.
   (iii) The last day of the month before the month he or she attains age 65. (An individual who is entitled to social security or railroad retirement cash benefits for the month of attainment of age 65 is automatically entitled to hospital insurance under § 406.10.)
   (iv) The day of death.

(e) Continuation of Medicare entitlement when disability benefit entitlement ends because of substantial gainful activity (SGA)—(1) Definitions. As used in this section—
   Trial work period means the 9-month period provided under title II of the Act and as defined 20 CFR 404.1592, during which the individual may test his or her ability to work and still receive disability cash benefits; and
   Reentitlement period means a period as defined in 20 CFR 404.1592a that begins with the first month after the trial work period and ends with the 36th month after the trial work period or, if earlier, with the first month in which the impairment no longer exists or is no longer disabling. (During the reentitlement period, benefits may be discontinued because of SGA. However, if SGA is later discontinued, benefits may be reinstated without a new application and a new disability determination.)

(2) Duration of continued Medicare entitlement. Effective January 1, 1988, if an individual’s entitlement to disability benefits or status as a qualified disabled railroad retirement beneficiary ends because he or she engaged in, or demonstrated the ability to engage in, substantial gainful activity after the 36 months following the end of the trial work period, Medicare entitlement continues until the earlier of the following:
   (i) The last day of the 24th month following the first month of SGA occurring after the 15th month of the individual’s reentitlement period or, if later, the end of the month following the month the individual’s disability benefit entitlement ends.
   (ii) The last day of the month in which notice is mailed to the individual indicating that he or she is no longer entitled to hospital insurance because of an event or circumstance (for example, there has been medical improvement, or the disabled widow has remarried) that would terminate disability benefit entitlement if it had not already been terminated because of substantial gainful activity.

§ 406.13 Individual who has end-stage renal disease.

(a) Statutory basis and applicability. This section explains the conditions of entitlement to hospital insurance benefits on the basis of end-stage renal disease, and specifies the beginning and end of the period of entitlement. It implements section 226A of the Social Security Act.
§ 406.13  
Definitions. As used in this section:
End-stage renal disease (ESRD) means that stage of kidney impairment that appears irreversible and permanent and requires a regular course of dialysis or kidney transplantation to maintain life.
Child or spouse means a child or spouse whose relationship to the parent or spouse meets the relationship requirements for entitlement to child's monthly social security benefits or to wife's, husband's, widow's, widower's, mother's or father's monthly benefits, as set forth in 20 CFR part 404. However, the duration of relationship requirements apply only to divorced spouses. (See 20 CFR 404.331.)
Dependent child means a person who, on the first day he or she has end-stage renal disease, is unmarried and meets the dependency requirements for entitlement to child's social security benefits on the basis of a parent's earnings (see 20 CFR 404.350-404.365) and who—
(1) Is under age 22;
(2) Is under a disability that began before age 22; or
(3) Is under age 26, is receiving at least one-half support from that parent, and has continuously received at least one-half support from that parent since the day before attaining age 22.

(c) Requirements. An individual is entitled to hospital insurance benefits if—
(1) He or she is medically determined to have ESRD;
(2) He or she is:
(i) Fully or currently insured under the social security program (title II of the Act) or would be fully or currently insured if his or her employment (after 1936) as defined under the Railroad Retirement Act were considered "employment" under the Social Security Act;
(ii) Entitled to monthly social security or railroad retirement benefits; or
(iii) The spouse or dependent child of a person who meets the requirements of paragraph (c)(2)(i) or (c)(2)(ii) of this section;
(3) He or she has filed an application for Medicare Part A; and
(4) He or she has satisfied the waiting period explained in paragraph (e) of this section.
(d) Filing an application. (1) An individual may obtain an application form, and help in completing it, from any social security office.
(2) An application is not valid if it is filed earlier than the third month before the month in which the individual meets the conditions of paragraphs (c)(3), (c)(2), and (c)(4) of this section.
(3) If an individual who has ESRD dies before he or she has filed an application, or is unable to file because of physical or mental condition, a relative or other person responsible for his or her affairs may file in his or her behalf. If a responsible person is not available, the hospital or dialysis facility that furnished treatment may file the application.
(e) Beginning of entitlement—(1) Basic limitations. Entitlement can begin no earlier than the first month in which the individual meets the conditions specified in paragraph (c) of this section, or the 12th month before the month of application, whichever is later.
(2) Waiting period. Entitlement begins on the first day of the third month after the month in which the individual initiates a regular course of renal dialysis, if the course is maintained throughout the waiting period, unless entitlement would begin earlier under paragraph (e) (3) or (4) of this section. This means that if dialysis began in January, entitlement would begin April 1.
(3) Exceptions: Early kidney transplant. If the individual receives a transplant, entitlement begins with the first day of the month in which the transplant was performed. However, if the individual is admitted as an inpatient to a hospital that is an approved renal transplantation center or renal dialysis center (see §405.2102) for procedures preliminary to transplant surgery, entitlement begins—
(i) On the first day of the month in which he or she initially enters the hospital, if the transplant is performed in that month or in either of the next 2 months; or
(ii) On the first day of the second month before the month of kidney
transplantation, if the transplant is delayed more than 2 months after the month of initial hospital stay. For example, if an individual enters the hospital in January, and the transplant is performed in January, February, or March, entitlement would begin January 1. However, if the transplant is performed in April, entitlement would begin February 1.

(4) Exceptions: Self-dialysis training. Entitlement begins on the first day of the month in which a regular course of renal dialysis began if:

(i) Before the end of the waiting period, the individual participates in a self-dialysis training program offered by a participating Medicare facility that is approved to provide such training;

(ii) The patient’s physician has certified that it is reasonable to expect the individual will complete the training program and will self-dialyze on a regular basis; and

(iii) The regular course of dialysis is maintained throughout the time that would otherwise be the waiting period (unless it is terminated earlier because the individual dies).

(f) End of entitlement. Entitlement ends with—

(1) The end of the 12th month after the month in which a regular course of dialysis ends; or

(2) The end of the 36th month after the month in which the individual has received a kidney transplant.

(g) Resumption of entitlement. Entitlement is resumed under the following conditions:

(1) An individual who initiates a regular course of renal dialysis or has a kidney transplant during the 12-month period after the previous course of dialysis ended is entitled to Part A benefits and eligible to enroll in Part B with the month the regular course of dialysis is resumed or the month the kidney is transplanted.

(2) An individual who initiates a regular course of renal dialysis or has a kidney transplant during the 36-month period after an earlier kidney transplant is entitled to Part A benefits and eligible to enroll in Part B with the month the regular course of dialysis begins or with the month the subsequent kidney transplant occurs.

(3) An individual who initiates a regular course of renal dialysis more than 12 months after the previous course of regular dialysis ended or more than 36 months after the month of a kidney transplant is eligible to enroll in Part A and Part B with the month in which the regular course of dialysis is resumed. If he or she is otherwise entitled under the conditions specified in paragraph (c) of this section, including the filing of an application, entitlement begins with the month in which dialysis is initiated or resumed, without a waiting period, subject to the limitations of paragraph (e)(1) of this section.

[48 FR 12536, Mar. 25, 1983, as amended at 60 FR 22535, May 8, 1995]

§ 406.15 Special provisions applicable to Medicare qualified government employment.

(a) Definition. As used in this section, Medicare-qualified government employment means Federal, State, or local government employment that is subject only to the hospital insurance portion of the tax imposed by the Federal Insurance Contributions Act (F.I.C.A.). This includes—


(2) Wages paid to State and local government employees hired after March 31, 1996.

(3) Wages paid to State and local government employees hired before April 1, 1986 but whose employment after March 31, 1996 is covered, for Medicare purposes only, under an agreement under section 218 of the Act.

(b) Crediting of wages that are taxable only for Medicare purposes. Medicare qualified government employment is credited in the same way and in the same amount as social security covered employment is credited for monthly social security cash benefit purposes. However, since only the Medicare portion (not the social security portion) of the F.I.C.A. tax is imposed, Medicare qualified government employment does not help qualify the individual for monthly Social Security cash benefits.

(c) Required quarters of coverage. (1) To qualify for hospital insurance on the basis of Medicare qualified government employment, an individual must
have the number of quarters of coverage necessary to qualify for hospital insurance under §406.10, §406.12, or §406.13.

(2) An individual who has worked in Medicare qualified government employment may qualify for hospital insurance on the basis of Medicare qualified government employment exclusively, or a combination of Medicare qualified government employment and social security covered employment.

(d) Transitional provision for Federal employment. Any individual who was a Federal employee at any time both during and before January 1983 will receive credit for quarters of Federal employment before January 1983 without paying tax. This transitional provision applies even if the Federal employee did not receive Federal wages for January 1983, for instance, because he or she was on approved leave without pay or on loan to a State or foreign agency.

(e) Conditions of entitlement. An individual who has worked in Medicare qualified government employment (or any related individual who would be entitled to social security cash benefits on the employee’s record if Medicare qualified government employment qualified for those benefits) is entitled to hospital insurance benefits if he or she—

(1) Would meet the requirements of §406.10, §406.12, or §406.13 if Medicare qualified government employment were social security covered employment; and

(2) Has filed an application for hospital insurance.

For purposes of this section not more than 12 months before the month of application may be counted towards the 25-month qualifying period specified in §406.12(a).

(f) Beginning and end of entitlement—

(1) Basic rule. Subject to the limitations specified in paragraph (f)(2) and (f)(3) of this section, entitlement begins and ends as specified in §406.10, §406.12 or §406.13, whichever is used to establish hospital insurance entitlement for the Federal, State, or local government employee or related individual.

(2) Limitations: Federal government employment. (i) Hospital insurance entitlement based on Federal employment could not begin before January 1983.

(ii) No months before January 1983 may be used to satisfy the qualifying period required for entitlement based on disability.

(3) Limitations: State and local government employment. (i) Hospital insurance entitlement based on State or local government employment cannot begin before April 1986.

(ii) No months before April 1986 may be used to satisfy the qualifying period required for entitlement based on disability.

§ 406.20—Premium Hospital Insurance

(a) General provisions. Hospital insurance benefits are available to most individuals age 65 or over and to certain individuals under age 65 who do not qualify for those benefits under subpart B of this part and are willing to pay a monthly premium. This is called premium hospital insurance.

(b) Eligibility of individuals age 65 or over to enroll for premium hospital insurance. Any individual is eligible to enroll for Medicare Part A if he or she—

(1) Has attained age 65;

(2) Is a resident of the United States and is either—

(i) A citizen of the United States; or

(ii) An alien lawfully admitted for permanent residence who has resided in the United States continuously for the 5-year period immediately preceding the month in which he or she meets all other requirements;

(3) Is not eligible for Part A benefits under subpart B of this part; and

(4) Is entitled to supplementary medical insurance (Part B of Medicare) or is eligible and has enrolled for it during an enrollment period.

(c) Eligibility of individuals under age 65 to enroll for premium hospital insurance. An individual who has not attained age 65 is eligible to enroll for Medicare Part A if he or she meets the following conditions:

(1) Has been entitled to Medicare Part A (under §406.12 or §406.15) on the
§ 406.21 Individual enrollment.

(a) Basic provision. An individual who meets the requirements of § 406.20(b) or (c) may enroll for premium hospital insurance only during his or her “initial enrollment period”, a “general enrollment period”, a “special enrollment period”, or, for HMO/CMP enrollees, a “transfer enrollment period”, as set forth in paragraphs (b) through (f) of this section.

(b) Initial enrollment periods—(1) Initial enrollment period for individual age 65 or over. The initial enrollment period extends for 7 months, from the third month before the month the individual first meets the requirements of § 406.20(b)(1) through (b)(3) through the third month after that first month of eligibility.

(2) Initial enrollment period for individual under age 65. The initial enrollment period begins with the month in which the individual receives notice that entitlement to Medicare Part A will end because he or she has lost entitlement to disability benefits solely because of earnings in excess of the amounts allowed under the social security regulations pertaining to “substantial gainful activity” (20 CFR 404.1571–404.1574); and (4) is not otherwise entitled to Medicare Part A.


§ 406.21 Individual enrollment.

(a) Basic provision. An individual who meets the requirements of § 406.20(b) or (c) may enroll for premium hospital insurance only during his or her “initial enrollment period”, a “general enrollment period”, a “special enrollment period”, or, for HMO/CMP enrollees, a “transfer enrollment period”, as set forth in paragraphs (b) through (f) of this section.

(b) Initial enrollment periods—(1) Initial enrollment period for individual age 65 or over. The initial enrollment period extends for 7 months, from the third month before the month the individual first meets the requirements of § 406.20(b)(1) through (b)(3) through the third month after that first month of eligibility.

(2) Initial enrollment period for individual under age 65. The initial enrollment period begins with the month in which the individual receives notice that entitlement to Medicare Part A will end because he or she has lost entitlement to disability benefits solely because of earnings in excess of the amounts allowed under the social security regulations pertaining to “substantial gainful activity” (20 CFR 404.1571–404.1574); and (4) is not otherwise entitled to Medicare Part A.

HMO or CMP, or during the first month that he or she is no longer enrolled in the HMO or CMP, part A coverage will begin on the first day of the month of part A enrollment, or, at the option of the individual, on the first day of any of the following 3 months.

(ii) If the individual enrolls in premium hospital insurance during any of the last 7 months of the transfer enrollment period, coverage will begin on the first day of the month after the month of enrollment.

§ 406.22 Effect of month of enrollment on entitlement.

(a) Individual age 65 or over. For an individual who has attained age 65, the following rules apply:

(1) If the individual enrolls during the 3 months before the first month of eligibility, entitlement begins with the first month of eligibility.

(2) If the individual enrolls in the first month of eligibility, entitlement begins with the following month.

(3) If the individual enrolls during the month after the first month of eligibility, entitlement begins with the second month after the month of enrollment.

(4) If the individual enrolls in either of the last 2 months of the enrollment period, entitlement begins with the third month after the month of enrollment.

(b) Individual under age 65. For an individual who has not attained age 65, the following rules apply:

(1) If the individual enrolls before the month in which he or she meets the requirements of §406.20(c), entitlement begins with the month in which the individual meets those requirements.

(2) If the individual enrolls in the month in which he or she first meets the requirements of §406.20(c), entitlement begins with the following month.

(3) If the individual enrolls in the month following the month in which he or she meets the requirements of §406.20(c), entitlement begins with the second month after the month of enrollment.

§ 406.24 Special enrollment period.¹

(a) Terminology. As used in this subpart, the following terms have the indicated meanings.

(1) Current employment status has the meaning given this term in §411.104 of this chapter.

(2) Family member has the meaning given this term in §411.201 of this chapter.

(3) Group health plan (GHP) and large group health plan (LGHP) have the meanings given those terms in §411.101 of this chapter, except that the “former employee” language of those definitions does not apply with respect to SEPs because—

(i) Section 1837(i)(1)(A) of the Act explicitly requires that GHP coverage of an individual age 65 or older, be by reason of the individual’s (or the individual’s spouse’s) current employment status;

(ii) The sentence following section 1837(i)(1)(B) of the Act refers to “large group health plan”. Under section 1862(b)(1)(B)(i), as amended by OBRA ’93, LGHP coverage of a disabled individual must be “by virtue of the individual’s or a family member’s current employment status with an employer”.

(b) Duration of SEP.² (1) The SEP includes any month during any part of which—

¹Before August 1996, SEPs were available only for enrollment in supplementary medical insurance, not for enrollment in premium hospital insurance.

²Before March 1995, SEPs began on the first day of the first month the individual was no longer covered under a GHP or LGHP by reason of current employment status.
§ 406.26 Enrollment under State buy-in.

(a) Enrollment of QMBs under a State buy-in agreement—(1) Effective date. Beginning with calendar year 1990, a State may request and be granted a modification of its buy-in agreement to include enrollment and payment of Part A premiums for QMBs (as defined in section 1905(p)(1) of the Act) who can become entitled to Medicare Part A only by paying a premium.

(2) Amount of premium. Premiums paid under State buy-in are not subject to increase because of late enrollment or reenrollment.

(b) Beginning of coverage under buy-in. The coverage period begins with the latest of the following:

(1) The third month following the month in which the agreement modification covering QMBs is effectuated.

(2) The first month in which the individual is entitled to premium hospital insurance under § 406.20(b) and has QMB status.

(3) The date specified in the agreement modification.

(c) End of coverage under buy-in. Buy-in coverage ends with the earlier of the following:

(1) Death. Coverage ends on the last day of the month in which the QMB dies.

(2) Loss of QMB status. If the individual loses eligibility for QMB status, coverage ends on the last day of the month in which HCFA receives the State’s notice of ineligibility.

Before August 10, 1993, an individual under age 65 could qualify for a SEP only if he or she had LGHP coverage as an “active individual”, which the statute defined as “an employee, employer, self-employed individual (such as the employer), individual associated with the employer in a business relationship, or as a member of the family of any of those persons”.

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(3) Termination of buy-in agreement. If the State's buy-in agreement is terminated, coverage ends on the last day of the last month for which the agreement is in effect.

(4) Entitlement to premium-free Part A. If the individual becomes entitled to premium-free Part A, buy-in coverage ends on the last day of entitlement to premium Part A.

(d) Continuation of coverage: Individual enrollment following termination of buy-in coverage—(1) Deemed enrollment. If coverage under a buy-in agreement ends because the agreement is terminated or the individual loses QMB status, the individual—
   (i) is considered to have enrolled during his or her initial enrollment period; and
   (ii) is entitled to Part A benefits and liable for Part A premiums beginning with the first month for which he or she is no longer covered under the buy-in agreement.

(2) Voluntary termination. (i) An individual may voluntarily terminate entitlement acquired under paragraph (d)(1) of this section by filing, with SSA or HCFA, a request for disenrollment.
   (ii) Voluntary disenrollment is effective as follows:
      (A) If the individual files a request within 30 days after the date of HCFA's notice that buy-in coverage has ended, the individual's entitlement ends on the last day of the last month for which the State paid the premium.
      (B) If the individual files the request more than 30 days but not more than 6 months after buy-in coverage ends, entitlement ends on the last day of the month in which the request is filed.
      (C) If the individual files the request later than the 6th month after buy-in coverage ends, entitlement ends at the end of the month after the month in which request is filed.

[56 FR 38080, Aug. 12, 1991]

§ 406.28 End of entitlement.

Any of the following actions or events ends entitlement to premium hospital insurance:

(a) Filing of request for termination. The beneficiary may at any time give HCFA or the Social Security Administration written notice that he or she no longer wishes to participate in the premium hospital insurance program.

(1) If he or she files the notice before entitlement begins, he or she will be deemed not to have enrolled.

(2) If he or she files the notice after entitlement begins, that entitlement will end at the close of the month following the month in which he or she filed the notice.

(b) Eligibility for hospital insurance without premiums. (1) If an individual meets the eligibility requirements for hospital insurance specified in § 406.10, § 406.11, § 406.13 or § 406.15, entitlement to premium hospital insurance ends with the month before the month in which he or she meets those requirements.

(2) If an individual meets the requirements of § 406.10, § 406.11, § 406.13, or § 406.15, he or she will be deemed to have filed the required application for hospital insurance benefits in his or her first month of eligibility under that section.

(c) End of entitlement to supplementary medical insurance (SMI) for individual who has attained age 65. In the case of an individual enrolled on the basis of § 406.20(b), entitlement to premium hospital insurance ends on the same date that entitlement to SMI ends.

(d) Nonpayment of premium. (1) If an individual fails to pay the premium bill, entitlement will end on the last day of the third month after the billing month.

(2) HCFA may reinstate entitlement if the individual shows good cause for failure to pay on time, and pays all overdue premiums within 3 calendar months after the date specified in paragraph (d)(1) of this section.

(e) Death. Entitlement ends with the day of death. (A premium is due for the month of death.)

(f) End of disabling impairment for individual under age 65. In the case of an individual enrolled on the basis of § 406.20(c), entitlement to premium hospital insurance ends on the last day of the month after the month in which the individual is notified that he or she no longer has a disabling impairment.

§ 406.32 Monthly premiums.

(a) Promulgation and effective date. Beginning with 1984, premiums are promulgated each September, effective for the succeeding calendar year.

(b) Monthly premiums: Determination of dollar amount.

(1) Effective for calendar years beginning January 1989, the dollar amount is determined based on an estimate of one-twelfth of the average per capita costs for benefits and administrative costs that will be payable with respect to individuals age 65 or over from the Federal Hospital Insurance Trust Fund during the succeeding calendar year.

(2) Before 1989, the dollar amount was determined by multiplying $33 by the ratio of the next year's inpatient deductible to $76, which was the inpatient deductible determined for 1973. (Because of cost controls, the deductible actually charged for that year was $72.)

(3) Effective for months beginning January 1994, if an individual meets the requirements in paragraph (c) of this section, the monthly premium determined under paragraph (b)(1) of this section is reduced in each month in which the individual meets the requirements by 25 percent in 1994, 30 percent in 1995, 35 percent in 1996, 40 percent in 1997 and 45 percent in 1998 and thereafter.

(4) The amount determined under paragraphs (b)(1), (2), or (3) of this section is rounded to the next nearest multiple of $1. (Fifty cents is rounded to the next higher dollar.)

(c) Qualifying for a reduction in monthly premium. An individual who qualifies for the reduction described in paragraph (b)(3) of this section must be an individual who—

(1) Has 30 or more quarters of coverage (QCs) as defined in 20 CFR 404.140 through 404.146;

(2) Has been married for at least the previous one year period to a worker who has 30 or more QCs;

(3) Had been married to a worker who had 30 or more QCs for a period of at least one year before the death of the worker;

(4) Is divorced from, after at least 10 years of marriage to, a worker who subsequently died and who had 30 or more QCs at the time the divorce became final;

(5) Is divorced from, after at least 10 years of marriage to, a worker who subsequently died and who had 30 or more QCs at the time the divorce became final.

(d) Monthly premiums: Increase for late enrollment and for reenrollment. For an individual who enrolls after the close of the initial enrollment period or reenrolls, the amount of the monthly premium, as determined under paragraph (b) of this section, is increased by 10 percent for each full 12 months in the periods described in §§ 406.33 and 406.34. Effective beginning with premiums due for July 1986, the premium increase is limited to 10 percent and is payable for twice the number of full 12-month periods determined under those sections.

(e) Collection of monthly premiums. (1) HCFA will bill the enrollee on a monthly basis and include an addressed return envelope with the bill.

(2) The enrollee must pay by check or money order that is payable to “HCFA Medicare Insurance,” and shows his or her name and the claim number that appears on his or her Medicare card. He or she must return the bill with the check or money order.

(f) Months for which payment is due.

(1) A premium payment is due for each month beginning with the first month of coverage and continuing through the month of death or if earlier, the month in which coverage ends.

(2) A premium is due for the month of death if coverage is still in effect, even if the individual dies on the first day of the month.

(g) Option for group payments. A public or private organization may pay the premiums on behalf of one or more enrollees under a contract or other arrangement with HCFA if HCFA determines that this method of payment is administratively feasible. (The rules set forth in subpart E of part 408 of this chapter, for SMI premiums, also apply to group payment of Part A premiums.)

§ 406.33 Determination of months to be counted for premium increase: Enrollment.

(a) Enrollment before April 1, 1981, or after September 30, 1981. The months to be counted for premium increase are the months from the end of the initial enrollment period through the end of the general enrollment period, the special enrollment period, or the transfer enrollment period in which the individual enrolls, excluding the following:

(1) Any months before September 1973.

(2) For premiums due for months after May 1986, any months beginning with January 1983 during which the individual was enrolled in an employer group health plan based on the current employment of the individual or the individual’s spouse.

(3) Any months during the 7-month special enrollment period under § 406.21(e) during which premium hospital insurance coverage is in effect.

(4) Any months that the individual was enrolled in an HMO or CMP under part 417, subpart K of this chapter as described in § 406.21(f).

(b) Enrollment during the period April 1 through September 30, 1981. The months to be counted for premium increase are the months from the end of the initial enrollment period through the month in which the individual enrolled, excluding any months before September 1973.

(c) Examples.

(1) John F’s initial enrollment period ended July 1979 but he did not enroll until January 1980. The months to be counted are August 1979 through March 1980. Since only 8 months elapsed, there is no premium increase.

(2) Mary T’s initial enrollment period ended in April 1980 but she did not enroll until May 1981. The months to be counted are August 1979 through March 1980. Since only 8 months elapsed, there is no premium increase.

§ 406.34 Determination of months to be counted for premium increase: Reenrollment.

(a) First reenrollment before April 1, 1981, or after September 30, 1981. The months to be counted for premium increase are:

(1) The months specified in § 406.33(a) or (b); plus

(2) The months from the end of the first period of entitlement through the end of the general enrollment period in which the individual reenrolled.

(b) First reenrollment during the period April 1, 1981 through September 30, 1981. The months to be counted for premium increase are:

(1) The months specified in § 406.33(a); plus

(2) The months from the end of the first period of entitlement through the month in which the individual reenrolled.

(c) Subsequent reenrollment during the period April 1, 1981 through September 30, 1981. The months to be counted for premium increase are:

(1) The months specified in paragraph (a) of this section; plus

(2) The months from April 1981 through the month in which the individual reenrolled for the second time.

(3) Since only one reenrollment was permitted before April 1981, any months from the end of the individual’s first enrolment period to the month in which the individual reenrolled are counted.

(4) Vincent C’s initial enrollment period ended August 31, 1986. He was covered under his wife’s employer group health plan until she retired on May 31, 1989. He enrolled during June 1989, the first month of the special enrollment period under § 406.21(e). No months are countable for premium increase purposes because the exclusions of paragraph (a) of this section apply to all months.

(5) Terry P enrolled in the 1987 general enrollment period, with coverage effective July 1987. There were 28 months after the end of his initial enrollment period through the end of the 1987 general enrollment period. His premium is increased by 10 percent. The increase will be eliminated after he has paid the additional 10 percent for 48 months.
§ 406.38 Prejudice to enrollment rights because of Federal Government error.

(a) If an individual's enrollment or nonenrollment for premium hospital insurance is unintentional, inadvertent, or erroneous because of the error, misrepresentation, or inaction of a Federal employee, or any person authorized by the Federal Government to act on its behalf, the Social Security Administration or HCFA may take whatever action it determines is necessary to provide appropriate relief.

(b) The action may include—

(1) Designation of a special initial or general enrollment period;

(2) Designation of an entitlement period;

(3) Adjustment of premiums;

(4) Any combination of the actions specified in paragraph (b) (1) through (3) of this section; or

(5) Any other remedial action which may be necessary to correct or eliminate the effects of such error, misrepresentation, or inaction.


Subpart D—Special Circumstances That Affect Entitlement to Hospital Insurance

§ 406.50 Nonpayment of benefits on behalf of certain aliens.

(a) Hospital insurance benefit payments may not be made for services furnished to an alien in any month in which his or her monthly social security benefits are suspended (or would be suspended if he or she were entitled to those benefits) because the alien remains outside the United States for more than 6 months.

(b) Benefits will be payable beginning with services furnished in the first full calendar month the alien is back in the United States.


§ 406.52 Conviction of certain offenses.

(a) Penalty that affects entitlement.

(1) If an individual is convicted of any of the crimes listed in § 406.11(c) (1) and (2), the court may impose, in addition to all other penalties, a penalty that affects entitlement to hospital insurance, beginning with the month of conviction.

(2) The additional penalty is that the individual's income (or the income of the insured individual on whose earnings record he or she became or seeks to become entitled) for the year of conviction and any previous year may not be counted in determining the insured status necessary for entitlement to hospital insurance.

(b) Effect of pardon. If the President of the United States pardons the convicted individual, that individual regains (or may again seek) entitlement effective with the month following the month in which the pardon is granted.

PART 407—SUPPLEMENTARY MEDICAL INSURANCE (SMI) ENROLLMENT AND ENTITLEMENT

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AUTHORITY: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh).

SOURCE: 53 FR 47204, Nov. 22, 1988, unless otherwise noted.

Subpart A—General Provisions

§ 407.1 Basis and scope.
(a) Statutory basis. The supplementary medical insurance (SMI) program is authorized by Part B of title XVIII of the Social Security Act.
(b) Sections 1836 and 1837 set forth the eligibility and enrollment requirements.
(c) Section 1838 specifies the entitlement periods, which vary depending on the time and method of enrollment and on the basis for termination.
(d) Section 1843 sets forth the requirements for State buy-in agreements under which States may enroll, and pay the SMI premiums for, eligible individuals who are also eligible for cash assistance or Medicaid.
(e) Section 104(b) of the Social Security Amendments of 1965 (Pub. L. 89-97) specifies the limitations that apply to certain aliens and persons convicted of subversive activities.
(f) Scope. This part sets forth the eligibility, enrollment, and entitlement requirements and procedures for supplementary medical insurance. (The rules about premiums are in part 408 of this chapter.)

§ 407.2 General description of program.
Part B of Title XVIII of the Act provides for voluntary “supplementary medical insurance” available to most individuals age 65 or over and to disabled individuals who are under age 65 and entitled to hospital insurance. The SMI program is financed by premiums paid by (or for) each individual enrolled in the program, plus contributions from Federal funds. It covers certain physicians’ services, outpatient services, home health services, services furnished by rural health clinics (RHCs), Federally qualified health centers (FQHCs), ambulatory surgical centers (ASCs), and comprehensive outpatient rehabilitation facilities (CORFs), and other medical and other health services.

[57 FR 24980, June 12, 1992]

§ 407.4 Basic requirements for entitlement.
(a) An individual must meet the following requirements to be entitled to SMI:
§ 407.10 Eligibility to enroll.
(a) Basic rule. Except as specified in paragraph (b) of this section, an individual is eligible to enroll for SMI if he or she—
(1) Is entitled to hospital insurance under any of the rules set forth in §§ 406.10 through 406.15 of this chapter; or
(2) Meets the following requirements:
   (i) Has attained age 65. (An individual is considered to have attained age 65 on the day before the 65th anniversary of his or her birth.)
   (ii) Is a resident of the United States.
   (iii) Is a citizen of the United States, or an alien lawfully admitted for permanent residence who has resided continuously in the United States during the 5 years preceding the month in which he or she applies for enrollment.
(b) Exception. An individual is not eligible to enroll for SMI if he or she has been convicted of—
(1) Spying, sabotage, treason, or subversive activities under chapter 37, 105, or 115 of title 18 of the United States Code; or

§ 407.12 General enrollment provisions.
(a) Opportunity to enroll. (1) An individual who is eligible to enroll for SMI may do so during an initial enrollment period or a general enrollment period as specified in §§ 407.14, and 407.15. An individual who meets the conditions specified in § 407.20 may enroll during a special enrollment period, as provided in that section.
(2) An individual who fails to enroll during his or her initial enrollment period or whose enrollment has been terminated may enroll or reenroll during a general enrollment period, or, if he or she meets the specified conditions, during a special enrollment period.
(b) Enrollment periods ending on a non-workday. (1) If an enrollment period ends on a Federal nonworkday, that period is automatically extended to the next succeeding workday.
(2) A Federal nonworkday is any Saturday, Sunday, or Federal legal holiday or a day that is declared by statute or executive order to be a day on which...
Federal employees are not required to work.

§ 407.14 Initial enrollment period.
(a) Duration. (1) The initial enrollment period is the 7-month period that begins 3 months before the month an individual first meets the eligibility requirements of § 407.10 and ends 3 months after that first month of eligibility.
(2) In determining the initial enrollment period of an individual who is age 65 or over and eligible for enrollment solely because of entitlement to hospital insurance, the individual is considered as first meeting the eligibility requirements for SMI in the first day he or she becomes entitled to hospital insurance or would have been entitled if he or she had filed an application for that program.
(b) Deemed initial enrollment period. (1) SSA or HCFA will establish a deemed initial enrollment period for an individual who fails to enroll during the initial enrollment period because of a belief, based on erroneous documentary evidence, that he or she had not yet attained age 65. The period will be established as though the individual had attained age 65 on the date indicated by the incorrect information.
(2) A deemed initial enrollment period established under paragraph (b)(1) of this section is used to determine the individual's premium and right to enroll in a general enrollment period if that is advantageous to the individual.

§ 407.15 General enrollment period.
(a) Except as specified in paragraph (b) of this section, the general enrollment period is January through March of each calendar year.
(b) An unlimited general enrollment period existed between April 1 and September 30, 1981. Any eligible individual whose initial enrollment period had ended, or whose previous period of entitlement had terminated, could have enrolled or reenrolled during any month of that 6-month period.

§ 407.17 Automatic enrollment.
(a) Who is automatically enrolled. An individual is automatically enrolled for SMI if he or she:
(1) Resides in the United States, except in Puerto Rico;
(2) Becomes entitled to hospital insurance under any of the provisions set forth in §§ 406.10 through 406.15 of this chapter; and
(3) Does not decline SMI enrollment.
(b) Opportunity to decline automatic enrollment. (1) SSA will notify an individual that he or she is automatically enrolled under paragraph (a) of this section and grant the individual a specified period (at least 2 months after the month the notice is mailed) to decline enrollment.
(2) The individual may decline enrollment by submitting to SSA or HCFA a signed statement that he or she does not wish SMI.
(3) The statement must be submitted before entitlement begins, or if later, within the time limits set in the notice of enrollment.

§ 407.18 Determining month of automatic enrollment.
(a) An individual who is automatically enrolled in SMI under § 407.17 will have the month of enrollment determined in accordance with paragraphs (b) through (f) of this section. The month of enrollment determines the month of entitlement.
(b) An individual is automatically enrolled in the third month of the initial enrollment period if he or she—
(1) Is entitled to social security benefits under section 202 of the Act on the first day of the initial enrollment period;
(2) Is entitled to hospital insurance based on end-stage renal disease; on entitlement to disability benefits as a social security or railroad retirement beneficiary; or on deemed entitlement to disability benefits on the basis of Medicare-qualified government employment; or
(3) Establishes entitlement to hospital insurance by filing an application and meeting all other requirements (as set forth in subpart B of part 406 of this chapter) during the first 3 months of the initial enrollment period.
(c) If an individual establishes entitlement to hospital insurance on the basis of an application filed in the last 4 months of the SMI initial enrollment...
Under the current statute, the SEP provision applicable to disabled individuals covered under an LGHP expires on September 1998. Unless Congress changes that date, the last SEP available under those provisions will begin with June 1998.

§ 407.20 Special enrollment period related to coverage under group health plans.

(a) Terminology—(1) Group health plan (GHP) and large group health plan (LGHP). These terms have the meanings given them in §411.101 of this chapter except that the “former employee” language of those definitions does not apply with respect to SEPs for the reasons specified in §406.24(a)(3) of this chapter.

(2) Special enrollment period (SEP). This term has the meaning set forth in §406.24(a)(4) of this chapter. In order to use a SEP, an individual must meet the conditions of paragraph (b) and of paragraph (c) or (d) of this section, as appropriate.

(b) General rule. All individuals must meet the following conditions:

(1) They are eligible to enroll for SMI on the basis of age or disability, but not on the basis of end-stage renal disease.

(2) When first eligible for SMI coverage (4th month of their initial enrollment period), they were covered under a GHP or LGHP on the basis of current employment status or, if not so covered, they enrolled in SMI during their initial enrollment period; and

(3) For all months thereafter, they maintained coverage under either SMI or a GHP or LGHP. (Generally, if an individual fails to enroll in SMI during any available SEP, he or she is not entitled to any additional SEPs. However, if an individual fails to enroll during a SEP because coverage under the same or a different GHP or LGHP was restored before the end of that particular SEP, that failure to enroll does not preclude additional SEPs.)

(c) Special rule: Individual age 65 or over. For an individual who is or was covered under a GHP, coverage must be by reason of the current employment status of the individual or the individual’s spouse.

(d) Special rules: Disabled individual. Individuals entitled on the basis of disability (but not on the basis of end-stage renal disease) must meet conditions that vary depending on whether they were covered under a GHP or an LGHP.

(1) For a disabled individual who is or was covered under a GHP, coverage must be on the basis of the current employment status of the individual or the individual’s spouse.

(2) For a disabled individual who is or was covered under an LGHP, coverage must be as follows:

(i) Before August 10, 1993, as an “active individual”, that is, as an employee, employer, self-employed individual (such as the employer), individual associated with the employer in a business relationship, or as a member of the family of any of those persons.

(ii) On or after August 10, 1993, by reason of current employment status of the individual or a member of the individual’s family.

(e) Effective date of coverage. The rule set forth in §406.24(d) for Medicare Part A applies equally to Medicare Part B.

[61 FR 40346, Aug. 2, 1996]
§ 407.22 Request for individual enrollment.

(a) A request for enrollment is required of an individual who meets the eligibility requirements of § 407.10 and desires SMI, if the individual—

(1) Is not entitled to hospital insurance;

(2) Has previously declined enrollment in SMI;

(3) Has had a previous period of SMI entitlement which terminated;

(4) Resides in Puerto Rico or outside the United States; or

(5) Is enrolling or reenrolling during a special enrollment period under § 407.20.

(b) A request for enrollment under paragraph (a) of this section must:

(1) Be signed by the individual or someone acting in his or her behalf; and

(2) Be filed with SSA or HCFA during the initial enrollment period, a general enrollment period, or a special enrollment period as provided in § 407.20.


The following apply whether an individual is self-enrolled or automatically enrolled in SMI:

(a) Enrollment during initial enrollment period. (1) If an individual enrolls during the first three months of the initial enrollment period, entitlement begins with the first month of eligibility.

(2) If an individual enrolls during the fourth month of the initial enrollment period, entitlement begins with the following month.

(3) If an individual enrolls during the fifth month of the initial enrollment period, entitlement begins with the second month after the month of enrollment.

(4) If an individual enrolls in either of the last two months of the initial enrollment period, entitlement begins with the third month after the month of enrollment.

(5) Example. An individual first meets the eligibility requirements for enrollment in April. The initial enrollment period is January through July. The month in which the individual enrolls determines the month that begins the period of entitlement, as follows:

<table>
<thead>
<tr>
<th>Enrolls in initial enrollment period</th>
<th>Entitlement begins on—</th>
</tr>
</thead>
<tbody>
<tr>
<td>January</td>
<td>April 1 (month eligibility requirements first met).</td>
</tr>
<tr>
<td>February</td>
<td>April 1.</td>
</tr>
<tr>
<td>March</td>
<td>April 1.</td>
</tr>
<tr>
<td>April</td>
<td>May 1 (month following month of enrollment).</td>
</tr>
<tr>
<td>May</td>
<td>July 1 (second month after month of enrollment).</td>
</tr>
<tr>
<td>June</td>
<td>September 1 (third month after month of enrollment).</td>
</tr>
<tr>
<td>July</td>
<td>October 1 (third month after month of enrollment).</td>
</tr>
</tbody>
</table>

(b) Enrollment on reenrollment during general enrollment period. (1) If an individual enrolls or reenrolls during a general enrollment period before April 1, 1981 or after September 30, 1981, entitlement begins on July 1 of that calendar year.

(2) If an individual enrolled or reenrolled during the general enrollment period between April 1, 1981 and September 20, 1981, entitlement began with the third month after the month in which the enrollment request was filed.

(c) Enrollment or reenrollment during a SEP. The rules set forth in § 406.24(d) of this chapter apply.

§ 407.27 Termination of entitlement: Individual enrollment.

An individual’s entitlement will terminate for any of the following reasons:

(a) Death. Entitlement to SMI ends on the last day of the month in which the individual dies.

(b) Termination of hospital insurance benefits. If an individual’s entitlement to hospital insurance ends before the month in which he or she attains age 65, entitlement to SMI will end on the same day unless it has been previously terminated in accordance with paragraph (c) or (d) of this section.

(c) Request by individual. An individual may at any time give HCFA or SSA written notice that he or she no longer wishes to participate in SMI, and request disenrollment.

(1) Before July 1987, entitlement ended at the end of the calendar quarter after the quarter in which the individual filed the disenrollment request.

(2) For disenrollment requests filed in or after July 1987, entitlement ends
§ 407.30 Limitations on enrollment.

(a) Initial enrollment periods—(1) Individual under age 65. An individual who has not attained age 65 may have one or more periods of entitlement to hospital insurance, based on disability. Since each period of disability entitlement entitles the individual to hospital insurance and since entitlement to hospital insurance makes the individual eligible for SMI enrollment, an individual may have an SMI initial enrollment period for each continuous period of entitlement to hospital insurance.

(2) Individuals who have attained age 65. An individual who has attained age 65 may not have more than one initial enrollment period on the basis of age. However, if the individual develops ESRD after age 65, he or she may have another initial enrollment period based on meeting the requirements of §406.13 of this chapter.

(b) Number of enrollments. There is no limitation on the number of enrollments.

(c) Coverage under buy-in agreements. For purposes of paragraph (a) of this section, the continued enrollment of an individual following the end of coverage under a State buy-in agreement is considered an initial enrollment.

§ 407.32 Prejudice to enrollment rights because of Federal Government misrepresentation, inaction, or error.

If an individual’s enrollment or non-enrollment in SMI is unintentional, inadvertent, or erroneous because of the error, misrepresentation, on inaction of a Federal employee or any person authorized by the Federal Government to act in its behalf, the Social Security Administration or HCFA may take whatever action it determines is necessary to provide appropriate relief. The action may include:

(a) Designation of a special initial or general enrollment period;

(b) Designation of an entitlement period based on that enrollment period;

(c) Adjustment of premiums;

(d) Any combination of actions under paragraphs (a) through (c) of this section; or

(e) Any other remedial action that may be necessary to correct or eliminate the effects of the error, misrepresentation, or inaction.

Subpart C—State Buy-In Agreements

§ 407.40 Enrollment under a State buy-in agreement.

(a) Statutory basis. (1) Section 1843 of the Act, as amended through 1969, permitted a State to enter into an agreement with the Secretary to enroll in the SMI program certain individuals who are eligible for SMI and who are members of the buy-in group specified in the agreement. A buy-in group could include certain individuals receiving Federally-aided State cash assistance (with the option of excluding individuals also entitled to social security benefits or railroad retirement benefits) or could include all individuals eligible for Medicaid. Before 1981, December 31, 1969 was the last day on which a State could request a buy-in agreement or a modification to include a coverage group broader than the one originally selected.

(2) Section 945(e) of the Omnibus Reconciliation Act of 1980 (Pub. L. 96–499) further amended section 1843 to provide that, during calendar year 1981, a State could request a buy-in agreement if it did not already have one, or request a broader coverage group for an existing agreement.

(3) Several laws enacted during 1980-1987 had the effect of requiring that the buy-in groups available under section 1843 of the Act be expanded to include certain individuals who lose eligibility for cash assistance payments but are treated as if they were cash assistance recipients for Medicaid eligibility purposes.

(4) Section 301(e)(1) of the Medicare Catastrophic Coverage Act of 1988 (Pub. L. 100–360) amends section 1843 of the Act to restore the 1961 provisions on a permanent basis, effective “after 1988.”
(5) The same section 301, as amended by section 608(d)(14)(H) of the Family Support Act of 1988 (Pub. L. 100–485), further amended section 1843 of the Act, beginning January 1, 1989, to establish a new buy-in category consisting of Qualified Medicare Beneficiaries and to provide that a State may request a buy-in agreement if it does not already have one, or request a broader buy-in group for the existing agreement.

(b) Definitions. As used in this section, unless the context indicates otherwise—

Cash assistance means any of the following kinds of monthly cash benefits, authorized by specified titles of the Act and, for convenience, represented by initials, as follows:

AABD stands for aid to the aged, blind or disabled under the first title XVI of the Act in effect until December 31, 1973.

AB stands for aid to the blind under title X of the Act.

AFDC stands for aid to families with dependent children under Part A of title IV of the Act.

APTD stands for aid to the permanently and totally disabled under title XIV of the Act.

OAA stands for old-age assistance under title I of the Act.

SSI stands for supplemental security income for the aged, blind, and disabled under the second title XVI of the Act, effective January 1, 1974.

SSP stands for State supplementary payments, whether mandatory or optional, to an aged, blind, or disabled individual under the second title XVI or the Act.

Qualified Medicare Beneficiary or QMB means an individual who meets the definition in §400.200 of this chapter and, therefore, is eligible to have the State Medicaid agency pay Medicare cost sharing amounts on his or her behalf.

Railroad retirement beneficiary means an individual entitled to receive an annuity under the Railroad Retirement Act of 1974.

State buy-in agreement or buy-in agreement means an agreement authorized by section 1843 of the Act, under which a State secures SMI or premium HI coverage for individuals who are members of the buy-in group specified in the agreement, by enrolling them and paying the premiums on their behalf.

(c) Basic rules.

(1) A State that has a buy-in agreement in effect must enroll any individual who is eligible to enroll in SMI under §407.10.

(2) Any State that does not have a buy-in agreement in effect may request buy-in for any one of the groups specified in §§407.42 and 407.43.

(3) Any State that does have an agreement may request a modification to cover a broader buy-in group or cancel its current agreement and request a new agreement to cover a narrower group.

§ 407.42 Buy-in groups available to the 50 States, the District of Columbia, and the Northern Mariana Islands.

(a) Categories included in the buy-in groups. The buy-in groups that are available to the 50 States, the District of Columbia, and the Northern Mariana Islands are specified in paragraph (b) of this section in terms of the following categories:

(1) Category A: Individuals who—

(i) Receive SSI or SSP or both; and

(ii) Are covered under the State's Medicaid plan as categorically needy.

(2) Category B: Individuals who—

(i) Under the Act or any other provision of Federal law are treated, for Medicaid eligibility purposes, as though they were receiving SSI or SSP; and

(ii) Are covered under the State's Medicaid plan as categorically needy.

(3) Category C: Individuals who are receiving AFDC.

(4) Category D: Individuals who, under the Act or any other provision of Federal law, are treated, for Medicaid eligibility purposes, as though they were receiving AFDC.

(5) Category E: Individuals who, in accordance with §§435.114 or §435.134 of this chapter, are covered under the...
Rules for buy-in for premium hospital insurance for QMBs are set forth in §406.26 of this chapter.

(6) Category F: Individuals who are Qualified Medicare Beneficiaries.

(7) Category G: All other individuals who are eligible for Medicaid.

(b) Buy-in groups available. Any of the 50 States, the District of Columbia, and the Northern Mariana Islands may buy-in for one of the following groups:

(1) Group 1: Categories A through G.

(2) Group 2: Categories A through F.

(3) Group 3: Categories A through E.

(4) Group 4: Categories A, B, and F, individuals in categories C and D who are not social security or railroad retirement beneficiaries, and individuals in category E who are included in that category (in accordance with §435.134 of this chapter) because they received OAA, AB, APTD, or AABD in August 1972 or would have been eligible to receive such cash assistance for that month if they had applied or had not been institutionalized.

(5) Group 5: Categories A and B, individuals in categories C and D who are not social security or railroad retirement beneficiaries, and individuals in category E who are included in that category (in accordance with §435.134 of this chapter) because they received OAA, AB, APTD, or AABD in August 1972 or would have been eligible to receive such cash assistance for that month if they had applied or had not been institutionalized.

(6) Group 6: Categories A, B, and F, and individuals in category E who are included in that category (in accordance with §435.134 of this chapter) because they received AABD for that month if they had applied or had not been institutionalized. This option is available only to those States that had an AABD program as of December 31, 1973.

[56 FR 38081, Aug. 12, 1991]

§ 407.43 Buy-in groups available to Puerto Rico, Guam, the Virgin Islands, and American Samoa.

(a) Categories included in buy-in groups. The buy-in groups that are available to Puerto Rico, Guam, the Virgin Islands, and American Samoa, which are not covered by the SSI program, are described in paragraph (b) of this section in terms of the following categories:

(1) Category A: Individuals receiving OAA, AB, APTD, or AFDC.

(2) Category B: Individuals who, under the Act or any other provision of Federal law, are treated, for Medicaid eligibility purposes, as though they were receiving AFDC.

(3) Category C: Individuals who, in accordance with §436.112 of this chapter, are covered under the State’s Medicaid plan despite the increase in social security benefits provided by Public Law 92-336.

(4) Category D: Individuals who are Qualified Medicare Beneficiaries.

(5) Category E: All other individuals who are eligible for Medicaid.

(b) Buy-in groups available. Puerto Rico, Guam, the Virgin Islands, and American Samoa may choose any of the following coverage groups:

(1) Group 1: Categories A through E.

(2) Group 2: Categories A through D.

(3) Group 3: Categories A through C.

(4) Group 4: Individuals in category D, and individuals in categories A and B who are not social security or railroad retirement beneficiaries.

(5) Group 5: Individuals in categories A and B who are not social security or railroad retirement beneficiaries.

(6) Group 6: Individuals in category D, individuals in category A who are receiving OAA, and individuals in category C who are included in that category (in accordance with §435.134 of this chapter) because they received AABD for that month if they had applied or had not been institutionalized. This option is available only to those States that had an AABD program as of December 31, 1973.

[56 FR 38081, Aug. 12, 1991]
§ 407.47 Beginning of coverage under a State buy-in agreement.

(a) General rule. The beginning of an individual’s coverage period depends on two factors:

(1) The individual’s meeting the SMI eligibility requirements and the requirements for being a member of the buy-in group; and

(2) The effective date of the buy-in agreement or agreement modification that covers the group to which the individual belongs, and which may not be earlier than the third month after the month in which the agreement or modification is executed.

(b) Application of general rule: Medicaid eligibles who are, or are treated as, cash assistance recipients. For Medicaid eligibles who are, or are treated as, cash assistance recipients (that is, are members of categories A through E of § 407.42(a) or categories A through C of § 407.43(a)), coverage begins with the later of the following:

(1) The first month in which the individual—

(i) Meets the SMI eligibility requirements specified in § 407.10; and

(ii) Is a member of one of those categories.

(2) The month in which the buy-in agreement or agreement modification covering QMBs is effective.

(c) Application of general rule: Qualified Medicare Beneficiaries. For individuals who are QMBs (that is, are members of category F of § 407.42 or category D of § 407.43(a)), coverage begins with the later of the following:

(1) The first month in which the individual meets the SMI eligibility requirements specified in § 407.10, and has QMB status.

(2) The month in which the buy-in agreement or agreement modification covering QMBs is effective.

(d) Application of general rule: Other individuals eligible for Medicaid. For individuals who are members of category G of § 407.42(a) or category E of § 407.43(a), coverage begins with the later of the following:

(1) The second month after the month in which the individual—

(i) Meets the SMI eligibility requirements specified in § 407.10; and

(ii) Is determined to be eligible for Medicaid.
§ 407.48 Termination of coverage under a State buy-in agreement.

An individual’s coverage under a buy-in agreement terminates with the earliest of the following events:

(a) Death. Coverage ends on the last day of the month in which the individual dies.

(b) Loss of entitlement to hospital insurance benefits before age 65. If an individual loses entitlement to hospital insurance benefits before attaining age 65, coverage ends on the last day of the last month for which he or she is entitled to hospital insurance.

(c) Loss of eligibility for the buy-in group. If an individual loses eligibility for inclusion in the buy-in group, buy-in coverage ends as follows:

(1) On the last day of the last month for which he or she is eligible for inclusion in the group, if HCFA determines ineligibility or receives a State ineligibility notice by the 25th day of the second month after the month in which the individual becomes ineligible for inclusion in the group.

(2) On the last day of the second month before the month in which HCFA receives a State ineligibility notice later than the time specified in paragraph (c)(1) of this section. A notice received by HCFA after the 25th day of the month is considered to have been received in the following month.

(d) Termination or modification of buy-in agreement. If the State’s buy-in agreement is terminated, or modified to substitute a narrower buy-in group, coverage ends on the last day of the month for which the agreement was in effect, or covered the broader buy-in group.

[53 FR 47204, Nov. 22, 1988, as amended at 56 FR 38082, Aug. 12, 1991]

§ 407.50 Continuation of coverage: Individual enrollment following end of coverage under a State buy-in agreement.

(a) Deemed enrollment. When coverage under a buy-in agreement ends because the agreement terminates, or is modified to substitute a narrower buy-in group, or because the individual is no longer eligible for inclusion in the buy-in group, the individual—

(1) Is considered to have enrolled during his or her initial enrollment period; and

(2) Will be entitled to SMI on this basis and liable for SMI premiums beginning with the first month for which he or she is no longer covered under the buy-in agreement.

(b) Voluntary termination. (1) An individual may voluntarily terminate entitlement acquired under paragraph (a) of this section by filing, with SSA or HCFA, a request for disenrollment.

(2) Voluntary disenrollment is effective as follows:

(i) If the individual files a request within 30 days after the date of HCFA’s notice that buy-in coverage has ended, the individual’s entitlement ends on the last day of the last month for which the State paid the premium.

(ii) If the individual files the request more than 30 days but not more than 6 months after buy-in coverage ends, entitlement ends on the last day of the month in which the request is filed.

(iii) If the individual files the request later than the 6th month after buy-in coverage ends, entitlement ends at the end of the month in which request is filed.¹

[53 FR 47204, Nov. 22, 1988, as amended at 56 FR 38082, Aug. 12, 1991]

¹For requests filed before July 1987, entitlement ended on the last day of the calendar quarter after the quarter in which the disenrollment request was filed.
PART 408—PREMIUMS FOR SUPPLEMENTARY MEDICAL INSURANCE

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AUTHORITY: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh).
SOURCE: 52 FR 48115, Dec. 18, 1987, unless otherwise noted.

Subpart A—General Provisions

§ 408.1 Statutory basis.

(a) This part implements certain provisions of sections 1837 through 1840 and 1881(d) of the Social Security Act (the Act) and conforms to other regulations that implement section 1843 of the Act. Section 1838(b) requires regulations to establish when an individual’s coverage ends because of nonpayment of premiums. It also specifies that those regulations may provide a grace period for payment of overdue premiums without loss of coverage. Section 1839 sets forth the specific procedures for determining the amount of the monthly premium and section 1840 establishes the rules for payment of premiums. Section 1843 provides that a State may enter into a buy-in agreement to secure SMI coverage for certain individuals by enrolling them in the SMI program and paying the premiums on their behalf. Section 1881(d) provides that Medicare payment, for the reasonable charges incurred in connection with a kidney donation, shall
§ 408.2 Scope and purpose.

(a) This part sets forth the policies and procedures for determining the amount of monthly supplementary medical insurance (SMI) premiums, for the payment, collection, or refund of premiums, for termination of coverage because of nonpayment of premiums, and for reinstatement of coverage if certain conditions are met. It conforms to subpart C of part 407 of this chapter, which sets forth the requirements for State buy-in agreements. These policies are intended to protect enrollee coverage to the maximum degree compatible with maintaining the integrity of the SMI program.

(b) Policies that apply to premiums that certain individuals must pay in order to become entitled to Medicare Part A hospital insurance benefits, are set forth in part 406 of this chapter.

§ 408.3 Definitions.

As used in this part, unless the context indicates otherwise—

Enrollee means an individual who is enrolled in the SMI program under Medicare Part B.

Taxable year means the 12-month period (calendar or fiscal year) for which the individual files his or her income tax return.

§ 408.4 Payment obligations.

(a) Month for which payment is due. (1) A payment is due for each month, beginning with the first month of SMI coverage and continuing through the month of death or, if earlier, the month in which coverage terminates.

(2) A premium is due for the month of death, if SMI coverage is still in effect, even though the individual dies on the first day of the month.

(b) Overdue premiums. (1) Overdue premiums constitute an obligation enforceable against the enrollee or the enrollee's estate.

(2) Overdue premiums are collected—

(i) By deduction from social security or railroad retirement benefits or Federal civil service annuities;

(ii) Directly from the enrollee or the enrollee's estate; or

(iii) By offset against any SMI payments payable to the enrollee or the enrollee's estate.

(3) Interest is not charged on overdue premiums, except under a State buy-in agreement, as provided in §408.6(c)(4).

(c) Premiums not required for certain kidney donors. (1) No premiums are required for SMI benefits related to the donation of a kidney if the donor is not an enrollee.

(2) A kidney donor who is an enrollee is not relieved of the obligation for premiums.

§ 408.6 Methods and priorities for payment.

(a) Methods of payment—(1) General rules. Premiums are paid by one of the following four methods:

(i) Payment by a State under a buy-in agreement.

(ii) Deduction from monthly railroad retirement of social security cash benefits or Federal civil service annuities.

(iii) Direct remittance on an individual basis, by or on behalf of the enrollee.

(iv) Direct remittance on a group basis, by an employer, union, lodge or other organization, or by an entity of State or local government.
§ 408.8 Grace period and termination date.

(a) Grace period. (1) For all initial premium payments (monthly or quarterly), and subsequent monthly or quarterly payments, the grace period ends with the last day of the third month after the billing month.

(2) For payments required because the monthly benefit is less than the monthly premium, the grace period ends on April 30 of the year following the calendar year which the premiums are due.

(b) Extension of grace period: Last day is nonwork day. If the last day of the grace period is a Saturday, Sunday, legal holiday, or a day that, by statute or executive order, is a nonwork day for Federal employees, the grace period is extended to the next succeeding work day.

(c) Termination date. The end of the grace period is the termination date for SMI coverage if overdue premiums have not been paid by that date in accordance with § 408.68.

(d) Extension of grace period for good cause. (1) HCFA may reinstate entitlement, without interruption of coverage, if the individual shows good cause for failure to pay within the initial grace period, and pays all overdue premiums within three calendar months after the termination date.

(2) Good cause will be found if the individual establishes, by a credible statement, that failure to pay premiums within the initial grace period was due to conditions over which he or she had no control, or which he or she could not reasonably have been expected to foresee.

§ 408.10 Claim for monthly benefits pending concurrently with request for SMI enrollment.

(a) If it is clear that an individual who applies for social security or railroad retirement benefits and for SMI will be entitled to monthly benefits, the application for monthly benefits is processed simultaneously with the request for SMI enrollment.

(1) If monthly benefits are paid, the SMI premiums are deducted from those benefits.

(2) If monthly benefits are suspended (for instance, because the individual’s earnings exceed the maximum allowed by law), the enrollee is billed for direct remittance.

(b) If it is clear that an individual will be entitled to SMI, but there is substantial question as to eligibility for monthly benefits, the request for SMI enrollment is processed separately.

(1) When SMI enrollment is approved, the enrollee is billed for direct remittance.

(2) When the application for monthly benefits is adjudicated, the following rules apply:

(i) If monthly benefits are paid, the SMI premiums are deducted from those benefits, with appropriate adjustments for any premiums already paid by direct remittance.

(ii) If the application for monthly benefits is approved but the benefits are suspended, the grace period is as set forth in §408.8(a).

(iii) If the application for monthly benefits is denied, the grace period is as set forth in §408.8(a)(1).


Subpart B—Amount of Monthly Premiums

§ 408.20 Monthly premiums.

(a) Statutory provisions. (1) The law established a monthly premium of $3 for the initial period of the program. It also set forth criteria and procedures for the Secretary to follow each December, beginning with December 1968, to determine and promulgate the standard monthly premium for the 12-month period beginning with July of the following year.

(2) The law was amended in 1983 to require that the Secretary promulgate the standard monthly premium in September of that year, and each year thereafter, to be effective for the 12 months beginning with the following January.

(3) The standard monthly premium applies to individuals who enroll during their initial enrollment periods. In other situations, that premium may be increased or decreased as specified in this subpart.

(b) Criteria and procedures for the period from July 1976 through December 1983, the period from January 1991 through December 1995, and for periods after December 1998.

(1) For periods from July 1976 through December 1983, and after December 1998, the Secretary determines and promulgates as the standard monthly premium (for disabled as well as aged enrollees) the lower of the following:

(i) The actuarial rate for the aged.

(ii) The monthly premium promulgated the previous December for the year beginning July 1, increased by a percentage that is the same as the latest cost-of-living increase in old age insurance benefits that occurred before the current promulgation. (Because of the change in the effective dates of the premium amount (under paragraph (a)(2) of this section), there was no increase in the standard monthly premium for the period July 1983 through December 1983.)

(2) For periods after December 1998, the Secretary determines the standard monthly premium in the manner specified in paragraph (b)(1) of this section, but promulgates it in September for the following calendar year.

(3) The premiums for calendar years 1991 through 1995 are those amounts as specified by section 1839(e)(1)(B) of the Act as follows:

(i) In 1991, $29.90;

(ii) In 1992, $31.80;

(iii) In 1993, $36.60;

(iv) In 1994, $40.80;

(v) In 1995, $43.00.
§ 408.24

(iv) In 1994, $41.10; and
(v) In 1995, $46.10.

(c) Premiums for calendar years 1984 through 1990 and 1996 through 1998. For calendar years 1984 through 1990 and 1996 through 1998, the standard monthly premium for all enrollees—

(1) Is equal to 50 percent of the actuarial rate for enrollees age 65 or over, that is, is calculated on the basis of 25 percent of program costs without regard to any cost-of-living increase in old age insurance benefits; and
(2) Is promulgated in the preceding September.

(d) Limitation on increase of standard premium: 1987 and 1988. If there is no cost-of-living increase in old age or disability benefits for December 1985 or December 1986, the standard monthly premiums for 1987 and 1988 (promulgated in September 1986 and September 1987, respectively) may not be increased.

(e) Nonstandard premiums for certain cases—(1) Basic rule. A nonstandard premium may be established in individual cases only if the individual is entitled to old age or disability benefits for the months of November and December, and actually receives the corresponding benefit checks in December and January.
(2) Special rules: Calendar years 1987 and 1988. For calendar years 1987 and 1988, the following rules apply:
(i) A nonstandard premium may be established if there is a cost-of-living increase in old age or disability benefits but, because the increase in the standard premium is greater than the cost-of-living increase, the beneficiary would receive a lower cash benefit in January than he or she received in December.
(ii) A nonstandard premium may not be established if the reduction in the individual's benefit would result, in whole or in part, from any circumstance other than the circumstance described in paragraph (e)(2)(i) of this section.
(3) Special rule: Calendar years after 1988. (i) Beginning with calendar year 1989, a premium increase greater than the cost-of-living increase is still a prerequisite for a nonstandard premium.
(ii) However, a nonstandard premium is not precluded solely because the cash benefit is further reduced as a result of government pension offset or workers' compensation payment.
(4) Amount of nonstandard premium. The nonstandard premium is the greater of the following:
(i) The premium paid for December,
(ii) The standard premium promulgated for January, reduced as necessary to compensate for—
(A) The fact that the cost-of-living increase was less than the increase in the standard premium; or
(B) The further reduction in benefit because of government pension offset or workers' compensation payments.
(5) Effective dates of nonstandard premium. A nonstandard premium established under this paragraph (e) continues in effect for the rest of the calendar year even if later there are retroactive adjustments in benefit payments. (The nonstandard premium could be affected by a determination that the individual had not established, or had lost, entitlement to monthly benefits for November or December, or both.)
(6) Effect of late enrollment or reenrollment. A nonstandard premium is subject to increase for late enrollment or reenrollment as required under other sections of this subpart. The increase is computed on the basis of the standard premium and added to the nonstandard premium.

§ 408.22 Increased premiums for late enrollment and for reenrollment.

For an individual who enrolls after expiration of his or her initial enrollment period or reenrolls after termination of a coverage period, the standard monthly premium determined under §408.20 is increased by ten percent for each full twelve months in the periods specified in §§408.24 and 408.25.

§ 408.24 Individuals who enrolled or reenrolled before April 1, 1981 or after September 30, 1981.

(a) Enrollment. For an individual who first enrolled before April 1, 1981 or after September 30, 1981, the period includes the number of months elapsed between the close of the individual's initial enrollment period and the close
§ 408.25 of the enrollment period in which he or she first enrolled, and excludes the following:

(1) The three months of January through March 1968, if the individual first enrolled before April 1968.

(2) Any months before January 1973 during which the individual was precluded from enrolling or reenrolling by the 3-year limitation on enrollment or reenrollment that was in effect before October 30, 1972.

(3) Any months in or before a period of coverage under a State buy-in agreement.

(4) For an individual under age 65, any month before his or her current continuous period of entitlement to hospital insurance.

(5) For an individual age 65 or older, any month before the month he or she attained age 65.

(6) For premiums due for months beginning with September 1984 and ending with May 1986, the following:

(i) Any months after December 1982 during which the individual was—
(A) Age 65 to 69;
(B) Entitled to hospital insurance (Medicare Part A); and
(C) Covered under a group health plan (GHP) by reason of current employment status.

(ii) Any months of SMI coverage for which the individual enrolled during a special enrollment period as provided in §407.20 of this chapter.

(7) For premiums due for months beginning with June 1986, the following:

(i) Any months after December 1982 during which the individual was:
(A) Age 65 or over; and
(B) Covered under a GHP by reason of current employment status.

(ii) Any months of SMI coverage for which the individual enrolled during a special enrollment period as provided in §407.20 of this chapter.

(8) For premiums due for months beginning with January 1987, the following:

(i) Any months after December 1986 during which the individual was:
(A) A disabled Medicare beneficiary under age 65;
(B) Not eligible for Medicare on the basis of end stage renal disease, under §406.13 of this chapter; and
(C) Covered under an LGHP as described in §407.20 of this chapter.

(ii) Any months of SMI coverage for which the individual enrolled during a special enrollment period as provided in §407.20 of this chapter.

(9) For premiums due for months beginning with July 1990, the following:

(i) Any months after December 1986 during which the individual met the conditions of paragraphs (a)(8)(i)(A) and (a)(8)(i)(B) of this section, and was covered under a GHP by reason of the current employment status of the individual or the individual’s spouse.

(ii) Any months of SMI coverage for which the individual enrolled during a special enrollment period as provided in §407.20 of this chapter.

(b) Reenrollment. For an individual who reenrolled before April 1, 1981 or after September 30, 1981, the period:

(1) Includes the following:

(i) The number of months elapsed between the close of the individual’s initial enrollment period and the close of the enrollment period in which he or she first enrolled; plus

(ii) The number of months elapsed between the individual’s initial period of coverage and the close of the enrollment period in which he or she reenrolled; plus

(iii) The number of months elapsed between each subsequent period of coverage and the close of the enrollment period in which he or she reenrolled.

(2) Excludes the following:

(i) The periods specified in paragraphs (a)(1) through (a)(9) of this section; and

(ii) Any month before April 1, 1981 during which the individual was precluded from reenrolling by the two-enrollment limitation in effect before that date.

§ 408.25 Individuals who enrolled or reenrolled between April 1 and September 30, 1981.

(a) Basic rules. Except as specified in paragraph (b) of this section, the rules set forth in §408.24 apply to an individual who enrolled or reenrolled between April 1 and September 30, 1981.

(b) Exception. For an individual who enrolled or reenrolled between April 1 and September 30, 1981, the months to
be counted ran through the month in which he or she reenrolled. (During those 6 months, continuous open enrollment was in effect and there was no 3-month "general enrollment period".)

§ 408.26 Examples.

Example 1. Mr. J, who became age 65 and otherwise eligible for enrollment in November 1965, first enrolls in March 1968. The months to be included in determining the amount of the increase in Mr. J's premiums begin with June 1966 (the first month after the close of his initial enrollment period) and extend through December 1967 (the period January through March of 1968 is excluded in determining the total months) for a total of 19 months. Since there is only one full 12-month period in 19 months, Mr. J's premiums will be 10 percent greater than if he had enrolled in his initial enrollment period.

Example 2. Mr. V, who enrolled in December 1965, voluntarily terminates his enrollment effective midnight December 31, 1967. He enrolls for a second time in January 1969. The months to be included in determining the amount of the increase in Mr. V's premiums are January 1968 through March 1969, a total of 15 months. Since this totals one full 12-month period, Mr. V's monthly premium will be increased by 10 percent.

Example 3. Ms. N becomes age 65 in July 1965 and first enrolls in December 1967. She pays premiums increased by 10 percent above the regular rate, beginning July 1968, the first month of her SMI coverage. Ms. N fails to pay the premiums for the calendar quarter ending June 30, 1970, and her coverage is terminated on that date, the end of her grace period. Ms. N enrolls for a second time in January 1971. The months to be included in determining the amount of the increase in Ms. N's premiums are June 1966 through March 1969, a total of 19 months. Since this totals one full 12-month period, Ms. N's monthly premium will be increased by 10 percent.

Example 4. Mr. X attained age 65 in August 1966 and enrolled during his initial enrollment period. His coverage was terminated effective June 30, 1968, for nonpayment of premiums. He reenrolls in March 1973. For purposes of computing any applicable premium increase, he will not be charged any months between March 1971 (the end of the last general enrollment period during which he was eligible to reenroll under the law in effect before October 30, 1972) and January 1973. Therefore, he will be charged 36 months (July 1968-March 1971 plus January 1973-March 1973) and his premiums for his second period of coverage will be increased 30 percent.

Example 5. Ms. C, who attained age 65 in August 1973, had two periods of supplementary medical insurance coverage, both of which were terminated because of nonpayment of premiums: August 1973 through April 1975 and July 1977 through August 1978. She reenrolls in July 1981. The months to be included in determining the amount of premium increase are May 1975 through March 1977 (23 months) and April 1981 through July 1981 (4 months) for a total of 27 months. The 31 months from September 1978 through March 1981 may not be counted because Ms. C was prevented from reenrolling by the two-enrollment limitation in effect before April 1, 1981. For Ms. C, the standard monthly premium would be increased by 20 percent.

§ 408.27 Rounding the monthly premium.

Any monthly premium that is not a multiple of 10 cents is rounded to the nearest multiple of 10 cents, and any odd multiple of 5 cents is rounded to the next higher multiple of 10 cents.

Subpart C—Deduction From Monthly Benefits

§ 408.40 Deduction from monthly benefits: Basic rules.

(a) Deduction from monthly benefits. (1) Enrollees who are receiving monthly benefits do not have the option of paying by direct remittance to avoid deduction.
§ 408.42 Deduction from railroad retirement benefits.

(a) Responsibility for deductions. If an enrollee is entitled to railroad retirement benefits, his or her SMI premiums are deducted from those benefits by the Railroad Retirement Board (RRB) even though he or she is also entitled to social security benefits or a civil service annuity, or both.

(b) Action when benefits are suspended. If the railroad retirement benefits are suspended, the RRB sends premium notices requesting direct remittance, to be made in accordance with the rules set forth in Subpart D of this part.

§ 408.43 Deduction from social security benefits.

SSA, acting as HCFA's agent, deducts the premiums from the monthly social security benefits if the enrollee is not entitled to railroad retirement benefits. (If the benefit is less than the monthly premium, the benefit is withheld and the enrollee is required to pay the balance through direct remittance.)

§ 408.44 Deduction from civil service annuities.

(a) Responsibility for deductions. If an enrollee is not entitled to railroad retirement benefits or social security benefits, and is receiving a civil service annuity, the premiums are deducted from that annuity by the Office of Personnel Management (OPM) on the basis of a notice from SSA indicating that the annuitant is entitled to SMI.

(b) Deduction of spouse’s premiums. If the annuitant’s spouse is also enrolled for SMI and is not entitled to a civil service annuity or to social security or railroad retirement benefits, and the annuitant gives written consent, OPM also deducts the spouse’s premium from the annuitant’s monthly check.

(c) Withdrawal of annuitant’s consent. (1) If an annuitant wishes to withdraw consent for deduction of the spouse’s premium, he or she must send written notice of withdrawal to OPM.

(2) The withdrawal notice is effective with the third month after the month in which it is received, or with the month specified in the notice, whichever is later.

§ 408.45 Deduction from age 72 special payments.

(a) Deduction of premiums. SMI premiums are deducted from age 72 special payments made under section 228 of the Act or the payments are withheld under procedures that correspond to the rules set forth in §§ 408.40 and 408.43.

(b) Collection of premiums while age 72 special payments are suspended. If the age 72 special payments are suspended, HCFA or its agent notifies the enrollee to pay premiums by direct remittance, in accordance with the rules set forth in § 408.60.

(c) Grace period. The grace period ends with the last day of the third month after the billing month.

(d) Resumption of age 72 special payments. (1) If age 72 special payments are resumed before the end of the grace period and all premium arrears can be deducted from those special payments, SMI coverage continues and the enrollee need not pay by direct remittance.

(2) Subsequent special payments are reduced by the amount of the premium for as long as the enrollee receives special payments.

§ 408.46 Effect of suspension of social security benefits.

(a) Benefit payments to be resumed during the taxable year. (1) If social security benefit payments are scheduled to
be resumed during the enrollee’s current taxable year, the enrollee is not billed.

(2) The enrollee may, if he or she wishes, pay the premiums during suspension of benefits.

(b) Benefit payments not to be resumed during the enrollee’s current taxable year.

(1) If social security benefits are suspended for a period that will not permit collection of all premiums due from monthly benefits payable in the enrollee’s current taxable year, HCFA or its agents bill the enrollee and require direct remittance in accordance with subpart D of this part.

(2) The first billing is for whatever premiums are necessary to place the enrollee in a quarterly cycle.

(3) Thereafter, the billing is on a quarterly basis. (Quarters for different enrollees are staggered throughout the year.)

(4) The enrollee has the option of paying premiums for more than one quarter at the same time.

§ 408.47 [Reserved]

§ 408.50 When premiums are considered paid.

(a) Actual deduction. A premium is considered paid if it is actually deducted from a monthly benefit check. Therefore—

(1) The premium is “paid” even if SSA later finds that the benefit was paid in error; but

(2) A finding that a monthly benefit was erroneously withheld does not constitute payment of the premium for that month. Since there was no payment, there was no deduction. The enrollee is billed and continuance of coverage depends on payment of premiums before the end of the grace period or extended grace period.

(b) Payment within the grace period. Overdue premiums are considered paid within the grace period in the following situations:

(1) Benefits are resumed during the grace period. (i) Monthly cash benefit payments are payable for the last month of the initial grace period or for earlier months on the basis of a notice filed by the enrollee before the initial grace period ends; and

(ii) Those payments are sufficient to permit deduction of all overdue premiums.

(2) Annual earnings report or other report submitted during the grace period shows a benefit is due. (i) Before the end of the grace period, the enrollee submits a report clearly showing that monthly cash benefits, previously withheld, are payable; and

(ii) Those benefits are sufficient to permit deduction of the full amount of the overdue premiums.

(3) Premium arrears are paid by direct remittance. The enrollee makes a direct remittance payment of all overdue premiums before the end of the grace period.


§ 408.52 Change from direct remittance to deduction.

If a direct remittance enrollee becomes entitled to monthly benefits—

(a) The SMI premiums are deducted from those benefits; and

(b) The enrollee is notified of the deduction and of any adjustment of the initial benefit check that is required to collect overdue premiums or refund premiums paid in advance.

§ 408.53 Change from partial direct remittance to full deduction.

If a benefit that was less than the premium (and therefore required direct remittance of the difference) is increased to an amount equal to, or greater than, the premium—

(a) The full premium is paid from the benefit; and

(b) Any amounts the enrollee had paid toward premiums not yet due are refunded.

Subpart D—Direct Remittance: Individual Payment

§ 408.60 Direct remittance: Basic rules.

(a) Premiums not deducted from monthly benefits under Subpart C of this part or paid by a State buy-in agreement must be paid by direct remittance to HCFA or its agents, by or on behalf of the enrollee.
§ 408.62 Initial and subsequent billings.

(a) Monthly billing. (1) The first premium bill is for the period from the first month of coverage (or the first month of change from deduction or State buy-in payment) through the end of the first month after the month of billing.

(b) Quarterly billing. (1) The first premium bill is for the period from the first month of coverage (or of change from deduction or State buy-in payment) through the third month after the month of billing.

(c) Subsequent billings are for periods of one month.

§ 408.63 Billing procedures when monthly benefits are less than monthly premiums.

If monthly benefits are less than monthly premiums, the following procedures apply:

(a) Notice of amount due. At the beginning of SMI entitlement, and at the beginning of each succeeding calendar year, SSA—

(1) Notifies the enrollee of the amount of benefits payable for the rest of the year and the total premiums due for those same months; and

(2) Bills the enrollee for the difference.

(b) Notice of amount overdue. At the beginning of each succeeding calendar year, SSA—

(1) Notifies the enrollee of any amounts overdue for premiums for the preceding calendar year; and

(2) Indicates that if the amount still overdue on April 30 is equal to or greater than the premium for 3 months, SMI coverage will terminate on that date.

§ 408.65 Payment options.

(a) The enrollee is not asked to pay premiums at the time of enrollment but is instructed to pay them upon receipt of a premium bill from HCFA or its agents.

(b) However, if the enrollee wishes, he or she may pay from one to 12 months or from one to four quarters at the time of enrollment.
(2) The premiums can no longer be deducted from the civil service annuity of the enrollee or the enrollee's spouse. (3) The enrollee no longer qualifies for coverage under a State buy-in agreement, and is not entitled to social security or railroad retirement monthly benefits.

(b) Billing. When any of the events specified in paragraph (a) of this section occurs (or as soon thereafter as possible), HCFA or its agents bill the enrollee for direct remittance, in accordance with this subpart.

§ 408.82 Conditions for group billing.

HCFA agrees to a group billing arrangement only if the following conditions are met:

(a) Conditions the group payer must meet. The group payer submits a written request for group billing—

(1) Showing that all or part of the payments are made from the payer's funds or from funds due the enrollees and in the payer's possession; and

(2) Agreeing not to charge the enrollees for the service of paying the premiums or for the administrative costs such as recordkeeping and postage.

(b) Enrollees eligible for group payment.

(1) Group payment may be made only on behalf of individuals who are already enrolled and are being billed for direct remittance.

(2) Group payment may not be made for enrollees whose premiums are being deducted from monthly benefits in accordance with Subpart C of this part or being paid by the State under a buy-in agreement.

(c) Protection of enrollee's rights. The use of group billing must not jeopardize the enrollees' right—

(1) To confidentiality of personal information;

(2) To terminate enrollment;

(3) To resume individual payment of premiums if he or she wishes; and

(4) To receive notice of any action that affects the SMI benefits.

(d) Authorization by the enrollee. (1) To ensure maximum feasible protection of the rights specified in paragraph (c) of this section, each enrollee must give written authorization as specified in § 408.84(a)(2).

(2) A group payer that is not an entity of State or local government must submit all enrollee authorizations to HCFA.

(3) A group payer that is an entity of State or local government may retain the authorizations and certify to HCFA that it has on file an authorization for each enrollee included in the group.

(4) It is on the basis of the enrollee's authorization that HCFA sends the group payer information about each enrollee, as necessary to carry out the group payment function.

(e) Size of group. The number of enrollees must be at least 20, which is the minimum size sufficient to make group billing efficient. (Smaller groups may use the informal procedure described in § 408.80(b).)
§ 408.84 Billing and payment procedures.

(a) Initial premium notice. (1) HCFA or its agent always sends the initial premium notice to the enrollee.

(2) An enrollee who wishes to have the premiums paid on a group basis must give the notice to the group payer, along with written authorization for sending subsequent notices to the group payer and for release of the information required for the group payment process.

(b) Monthly billings. Group premiums are billed on a monthly basis. However, the group payer may pay up to 12 months in advance.

(c) Group payers must make their payments within 30 days after billing, to avoid infringing on the 90-day grace period during which the premiums may be paid by the enrollee if he or she is dropped from the group.

(d) Effect of group payment. Payment by a group payer is considered payment by the enrollee.

§ 408.86 Responsibilities under group billing arrangement.

(a) Enrollee responsibilities. (1) The enrollee is still responsible for premium payments; the group payer simply acts as his agent. If the agent fails to pay, or identifies the payment incorrectly, SSA notifies both the agent and the enrollee that the enrollee's account is delinquent. If an enrollee fails to take action on that notice, entitlement is terminated for nonpayment of premiums.

(2) The enrollee must promptly notify both SSA and the group payer of any change of address.

(b) Group payer's responsibilities. The group payer must—

(1) Make premium payments promptly upon receipt of notices;

(2) Promptly notify both HCFA and the enrollee when it drops an enrollee from the group;

(3) Make payments in a way that facilitates efficient and economical processing; and

(4) Maintain the confidentiality of the personal information obtained from HCFA for the group payment process.

(c) HCFA responsibilities. HCFA—

(1) Sends the bill to the group payer upon authorization from the enrollee;

(2) Notifies both the payer and the enrollee if the payer fails to make timely payments; and

(3) Refunds excess premiums in accordance with §408.88.

§ 408.88 Refund of group payments.

(a) Basis for refund. Group payments are refunded only in the following circumstances:

(1) The premium was for a month after the month in which the enrollee's SMI coverage terminated or the enrollee died.

(2) The premium was for a month after the month in which the group payer gave notice (before the 26th day of that month) that the enrollee was no longer eligible for group payment and was being dropped from the group.

(b) Example. F is the wife of J who is a retiree of Corporation X. That corporation pays premiums on behalf of all of its retirees and their dependents. F obtains a divorce from J on October 20 and thus disqualifies herself for further premium payments by the corporation. The corporation gives notice on November 10 that a refund is due because F has been dropped from the list of persons for whom it has agreed to pay premiums. The premium paid for December would be refunded to the group payer.

(c) To whom refund is made. (1) HCFA ordinarily refunds to the group payer the premiums specified in paragraph (a) of this section.

(2) However, if HCFA has information that clearly shows those premiums were paid from the enrollee's funds, it sends the refund to the enrollee.

§ 408.90 Termination of group billing arrangement.

(a) A group billing arrangement may be terminated either by the group payer or by HCFA upon 30 days' notice.

(b) HCFA may terminate the arrangement if it finds that the group payer is not acting in the best interest of the enrollees or that, for any other reason, the arrangement has proved inconvenient for HCFA.

§ 408.92 Change from group payment to deduction or individual payment.

(a) Enrollee excluded from group payment arrangement because of entitlement
to monthly benefits. (1) When an enrollee becomes entitled to monthly benefits from which premiums can be deducted as specified in subpart C of this part, HCFA notifies the group payer to discontinue payment for that enrollee.

(2) In order to maintain confidentiality, HCFA does not explain to the group payer the reason for excluding the enrollee from the group payment arrangement.

(3) The enrollee’s premiums are thereafter deducted from the monthly benefits, in accordance with subpart C of this part.

(b) Enrollee no longer eligible for the group. (1) When an enrollee is no longer eligible to be included in the group (for instance because he or she is no longer employed by the group payer or has terminated union or lodge membership), the group payer must promptly notify HCFA and the enrollee.

(2) HCFA or its agents resume sending individual bills to the enrollee, for direct remittance subject to the grace period and termination dates specified in §408.8.

Subpart F—Termination and Reinstatement of Coverage

§ 408.100 Termination of coverage for nonpayment of premiums.

(a) Effective date of termination. Termination is effective on the last day of the grace period. The determination is not made until 15 days after that day to allow for processing of remittances mailed late in the grace period, as provided in §408.68.

(b) Notice of termination. (1) SSA sends the enrollee notice of termination between 15 and 30 days after the end of the grace period and includes information regarding the enrollee’s right of appeal.

(2) HCFA notifies any intermediary or carrier that had previously been informed that the enrollee had met the SMI deductible for the year in which the termination is effective.

§ 408.102 Reconsideration of termination.

(a) Basic rules. Coverage may be reinstated without interruption of benefits if the following conditions are met:

(1) The enrollee appeals the termination by the end of the month following the month in which SSA sent the notice of termination.

(2) The enrollee alleges and it is found that the enrollee did not receive timely and adequate notice that the premiums were overdue.

(3) The enrollee pays, within 30 days after SSA’s subsequent request for payment, all premiums due through the month in which he or she appealed the termination.

(b) Basis for reinstating coverage. Coverage may be reinstated if the evidence establishes one of the following:

(1) The enrollee acted diligently to pay the premiums or to request relief upon receiving a premium notice very late in the grace period or shortly after its end, and the delayed notice was not the enrollee’s fault. (For example, if the billing notice was misaddressed or lost in the mail, it would not be the enrollee’s fault; if the enrollee had moved and not notified SSA of the new address, he or she would be responsible for the delay.)

(2) On the basis of information given by SSA, the enrollee could reasonably have believed that the premiums were being paid by deduction from benefits or by some other means. (An example would be a notice indicating that premiums would be paid by a State Medicaid agency or a group payer or would be deducted from the spouse’s civil service annuity.)

(c) No basis for reinstating coverage. Coverage may not be reinstated if the enrollee—

(1) Received timely and adequate notice but failed to pay within the grace period, for example because of insufficient income or resources; or

(2) Appealed the termination more than one month after the month in which SSA sent the termination notice.

§ 408.104 Reinstatement procedures.

(a) Request for payment. If the conditions of §408.102(a) (1) and (2) are met, SSA sends written notice requesting the enrollee to pay, within 30 days, all premiums due through the month in which the enrollee appealed the termination.
§ 408.110 Collection of unpaid premiums.

(a) Basis and scope—(1) Basis. Under the Federal Claims Collection Act of 1966 (31 U.S.C. 3711), HCFA is required to collect any debts due it but is authorized to suspend or terminate collection action on debts of less than $20,000 when certain conditions are met. (See 4 CFR, parts 101-105 for general rules implementing the Federal Claims Collection Act.) As indicated in § 408.4, unpaid premiums are debts owed the Federal government by the enrollee or the enrollee’s estate.

(2) Scope. This section sets forth the methods of collection used by HCFA and the circumstances under which HCFA terminates or renews collection action. The regulations in this section apply to hospital insurance premiums as well as SMI premiums.

(b) Collection of unpaid premiums. Generally, HCFA will attempt to collect unpaid premiums by one of the following methods:

(1) By billing enrollees who pay the premiums directly to HCFA or to a designated agent in accordance with § 408.60.

(2) By deduction from any benefits payable to the enrollee or the estate of a deceased enrollee under Title II or XVIII of the Social Security Act, the Railroad Retirement Act or any act administered by the Office of Personnel Management in accordance with § 408.4(b) and Subpart C of this part (Deduction from Monthly Benefits); or

(3) By billing the estate of a deceased enrollee.

(c) Termination of collection action. HCFA terminates collection action on unpaid premiums under either of the following circumstances, if the cost of collection exceeds the amount of overdue premiums:

(1) The individual is not entitled to benefits under the Acts listed in paragraph (b)(2) of this section, is not currently enrolled for SMI or premium hospital insurance, and demonstrates, to HCFA’s satisfaction, that he or she is unable to pay the debt within a reasonable time.

(2) The individual has been dead more than 27 months (the maximum time allowed for claiming SMI benefits), and the legal representative of his or her estate demonstrates, to HCFA’s satisfaction, that the estate is unable to pay the debt within a reasonable time.

(d) Renewal of collection efforts. HCFA renews collection efforts in either of the following circumstances, if the cost of collection does not exceed the amount of the overdue premiums:

(1) The individual enrolls again for premium hospital insurance or SMI. (Payment of overdue premiums is not a prerequisite for reenrollment.)

(2) The individual becomes entitled or reentitled to social security or railroad retirement benefits or a Federal civil service annuity.

§ 408.112 Refund of excess premiums after the enrollee dies.

If HCFA has received premiums for months after the enrollee’s death, HCFA refunds those premiums as follows:

(a) To the person or persons who paid the premiums or, if the premiums were paid by the enrollee, to the representative of the enrollee’s estate, if any.

(b) If refund cannot be made under paragraph (a) of this section, HCFA refunds the premiums to the enrollee’s survivors in the following order of priority:

(1) The surviving spouse, if he or she was either living in the same household with the deceased at the time of death, or was, for the month of death, entitled to monthly social security or railroad retirement benefits on the basis of the same earnings record as the deceased beneficiary.

(2) The child or children who were, for the month of death, entitled to monthly social security or railroad retirement benefits on the basis of the same earnings record as the deceased (and, if there is more than one child, in equal parts to each child);
(3) The parent or parents who were, for the month of death, entitled to monthly social security or railroad retirement benefits on the basis of the same earnings record as the deceased (and, if there is more than one parent, in equal parts to each parent);

(4) The surviving spouse who was not living in the same household with the deceased at the time of death and was not, for the month of death, entitled to monthly social security or railroad retirement benefits on the basis of the same earnings record as the deceased beneficiary;

(5) The child or children who were not entitled to monthly social security or railroad retirement benefits on the basis of the same earnings record as the deceased (and, if there is more than one child, in equal parts to each child);

(6) The parent or parents who were not entitled to monthly social security or railroad retirement benefits on the basis of the same earnings record as the deceased (and, if there is more than one parent, in equal parts to each parent).

If none of the listed relatives survives, no refund can be made.

PART 409—HOSPITAL INSURANCE BENEFITS

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§ 409.1 Statutory Basis.

This part is based on the identified provisions of the following sections of the Social Security Act:

(a) Sections 1812 and 1813 establish the scope of benefits of the hospital insurance program under Medicare Part A and set forth deductible and coinsurance requirements.

(b) Sections 1814 and 1815 establish conditions for, and limitations on, payment for services furnished by providers.

(c) Section 1820 establishes the critical access hospital program.

(d) Section 1861 describes the services covered under Medicare Part A, and benefit periods.

(e) Section 1862(a) specifies exclusions from coverage; and section 1862(h) requires a registry of pacemakers.

(f) Section 1881 sets forth the rules for individuals who have end-stage renal disease (ESRD), for organ donors, and for dialysis, transplantation, and other services furnished to ESRD patients.

409.2 Scope.

Subparts A through G of this part describe the benefits available under Medicare Part A and set forth the limitations on those benefits, including certain amounts of payment for which beneficiaries are responsible.


§ 409.3 Definitions.

As used in this part, unless the context indicates otherwise—

Arrangements means arrangements which provide that Medicare payment made to the provider that arranged for the services discharges the liability of the beneficiary or any other person to pay for those services.

Covered refers to services for which the law and the regulations authorize Medicare payment.

Nominal charge provider means a provider that furnishes services free of charge or at a nominal charge and is either a public provider, or another provider that (1) demonstrates to HCFA’s satisfaction that a significant portion of its patients are low-income, and (2) requests that payment for its services be determined accordingly.

Participating refers to a hospital or other facility that meets the conditions of participation and has in effect a Medicare provider agreement.

Qualified hospital means a facility that—

(a) Is primarily engaged in providing, by or under the supervision of doctors of medicine or osteopathy, inpatient services for the diagnosis, treatment, and care or rehabilitation of persons who are sick, injured, or disabled;

(b) Is not primarily engaged in providing skilled nursing care and related services for inpatients who require medical or nursing care;

(c) Provides 24-hour nursing service in accordance with Sec. 1861(e)(5) of the Act;

(d) If it is a U.S. hospital, is licensed, or approved as meeting the standards for licensing, by the State or local licensing agency; and

(e) If it is a foreign hospital, is licensed, or approved as meeting the standard for licensing, by the appropriate Canadian or Mexican licensing agency, and for purposes of furnishing non-emergency services to U.S. residents, is accredited by the Joint Commission on Accreditation of Hospitals.
§ 409.11 Bed and board.

(a) Semiprivate and ward accommodations. Except for applicable deductible and coinsurance amounts, Medicare Part A pays in full for bed and board and semiprivate (2 to 4 beds), or ward (5 or more beds) accommodations.

(b) Private accommodations—(1) Conditions for payment in full. Except for applicable deductible and coinsurance amounts, Medicare Part A pays in full for a private room if—

(i) The patient's condition requires him or her to be isolated;

(ii) The hospital or CAH has no semi-private or ward accommodations; or

(iii) The hospital's or CAH's semi-private and ward accommodations are fully occupied by other patients, were so occupied at the time the patient was

§ 409.10 Included services.

(a) Subject to the conditions, limitations, and exceptions set forth in this subpart, the term “inpatient hospital or inpatient CAH services” means the following services furnished to an inpatient of a participating hospital or of a participating CAH or, in the case of emergency services or services in foreign hospitals, to an inpatient of a qualified hospital:

(1) Bed and board.

(2) Nursing services and other related services.

(3) Use of hospital or CAH facilities.

(4) Medical social services.

(5) Drugs, biologicals, supplies, appliances, and equipment.

(6) Certain other diagnostic or therapeutic services.

(7) Medical or surgical services provided by certain interns or residents-in-training.

(8) Transportation services, including transport by ambulance.

(b) Inpatient hospital services does not include the following types of services:

(1) Posthospital SNF care, as described in §409.20, furnished by a hospital or a critical access hospital that has a swing-bed approval.

(2) Nursing facility services, described in §440.155 of this chapter, that may be furnished as a Medicaid service under title XIX of the Act in a swing-bed hospital that has an approval to furnish nursing facility services.

(3) Physician services that meet the requirements of §415.102(a) of this chapter for payment on a fee schedule basis.

(4) Physician assistant services, as defined in section 1861(s)(2)(K)(i) of the Act.

(5) Nurse practitioner and clinical nurse specialist services, as defined in section 1861(s)(2)(K)(ii) of the Act.

(6) Certified nurse mid-wife services, as defined in section 1861(gg) of the Act.

(7) Qualified psychologist services, as defined in section 1861(ii) of the Act.

(8) Services of an anesthetist, as defined in §410.69.
admitted to the hospital or CAH, respectively, for treatment of a condition that required immediate inpatient hospital or inpatient CAH care, and have been so occupied during the interval.

(2) Period of payment. In the situations specified in paragraph (b)(1)(i) and (iii) of this section, Medicare pays for a private room until the patient's condition no longer requires isolation or until semiprivate or ward accommodations are available.

(3) Conditions for patient's liability. The hospital or CAH may charge the patient the difference between its customary charge for the private room and its most prevalent charge for a semiprivate room if—

(i) None of the conditions of paragraph (b)(1) of this section is met; and

(ii) The private room was requested by the patient or a member of the family, who, at the time of the request, was informed what the hospital's or CAH's charge would be.

§ 409.12 Nursing and related services, medical social services; use of hospital or CAH facilities.

(a) Except as provided in paragraph (b) of this section, Medicare pays for nursing and related services, use of hospital or CAH facilities, and medical social services as inpatient hospital or inpatient CAH services only if those services are ordinarily furnished by the hospital or CAH, respectively, for the care and treatment of inpatients.

(b) Exception. Medicare does not pay for the services of a private duty nurse or attendant. An individual is not considered to be a private duty nurse or attendant if he or she is a hospital or CAH employee at the time the services are furnished.

§ 409.13 Drugs and biologicals.

(a) Except as specified in paragraph (b) of this section, Medicare pays for drugs and biologicals as inpatient hospital or inpatient CAH services only if—

(1) They represent a cost to the hospital or CAH;

(2) They are ordinarily furnished by the hospital or CAH for the care and treatment of inpatients; and

(3) They are furnished to an inpatient for use in the hospital or CAH.

(b) Exception. Medicare pays for a limited supply of drugs for use outside the hospital or CAH if it is medically necessary to facilitate the beneficiary's departure from the hospital and required until he or she can obtain a continuing supply.

§ 409.14 Supplies, appliances, and equipment.

(a) Except as specified in paragraph (b) of this section, Medicare pays for supplies, appliances, and equipment as inpatient hospital or inpatient CAH services only if—

(1) They represent a cost to the hospital or CAH;

(2) They are ordinarily furnished by the hospital or CAH to inpatients; and

(3) They are furnished to inpatients for use in the hospital or CAH.

(b) Exceptions. Medicare pays for items to be used beyond the hospital or CAH stay if—

(1) The item is one that the beneficiary must continue to use after he or she leaves the hospital or CAH, for example, heart valves or a heart pacemaker, or

(2) The item is medically necessary to permit or facilitate the beneficiary's departure from the hospital or CAH and is required until the beneficiary can obtain a continuing supply. Tracheostomy or draining tubes are examples.

§ 409.15 Services furnished by an intern or a resident-in-training.

Medical or surgical services provided by an intern or a resident-in-training are included as "inpatient hospital or inpatient CAH services" if they are provided—

(a) By an intern or a resident-in-training under a teaching program approved by the Council on Medical Education of the American Medical Association, or the Bureau of Professional Education of the American Osteopathic Association;
(b) By an intern or a resident-in-training in the field of dentistry under a teaching program approved by the Council on Dental Education of the American Dental Association; or
(c) By an intern or a resident-in-training in the field of podiatry under a teaching program approved by the Council on Podiatry Education of the American Podiatry Association.

§ 409.16 Other diagnostic or therapeutic services.

Diagnostic or therapeutic services other than those provided for in §§409.12, 409.13, and 409.14 are considered as inpatient hospital or inpatient CAH services if—
(a) They are furnished by the hospital or CAH, or by others under arrangements made by the hospital or CAH;
(b) Billing for those services is through the hospital or CAH; and
(c) The services are of a kind ordinarily furnished to inpatients either by the hospital or CAH or under arrangements made by the hospital or CAH.

§ 409.18 Services related to kidney transplantations.

(a) Kidney transplants. Medicare pays for kidney transplantation surgery only if performed in a renal transplantation center approved under subpart U of part 405 of this chapter.
(b) Services in connection with kidney donations. Medicare pays for services related to the evaluation or preparation of a potential or actual donor, to the donation of the kidney, or to post-operative recovery services directly related to the kidney donation—
(1) If the kidney is intended for an individual who has ESRD and is entitled to Medicare benefits or can be expected to become so entitled within a reasonable time; and
(2) Regardless of whether the donor is entitled to Medicare.

§ 409.19 Services related to cardiac pacemakers and pacemaker leads.

(a) Requirement. (1) Providers that request or receive Medicare payment for the implantation, removal, or replacement of permanent cardiac pacemakers and pacemaker leads must submit to HCFA the information required for the pacemaker registry.
(2) The required information is set forth under 21 CFR part 805 of the FDA regulations and must be submitted in accordance with general instructions issued by HCFA.

(b) Denial of payment. If HCFA finds that a provider has failed to comply with paragraph (a) of this section, HCFA denies payment for the implantation, removal, or replacement of any permanent cardiac pacemaker or pacemaker lead, effective 45 days after sending the provider written notice in accordance with paragraph (c) of this section.

(c) Notice of denial of payment. The notice of denial of payment—
(1) States the reasons for the determination;
(2) Grants the provider 45 days from the date of the notice to submit the information or evidence showing that the determination is in error; and
(3) Informs the provider of its right to hearing.
(d) Right to hearing. If the denial of payment determination goes into effect at the expiration of the 45-day period, it constitutes an “initial determination” subject to administrative and judicial review under part 498 of this chapter.

Subpart C—Posthospital SNF Care

§ 409.20 Coverage of services.

(a) Included services. Subject to the conditions and limitations set forth in this subpart and subpart D of this part, “posthospital SNF care” means the following services furnished to an inpatient of a participating SNF, or of a participating hospital or critical access hospital (CAH) that has a swing-bed approval:
(1) Nursing care provided by or under the supervision of a registered professional nurse.
(2) Bed and board in connection with the furnishing of that nursing care.
(3) Physical, occupational, or speech therapy.
§ 409.21 Nursing care.

(a) Basic rule. Medicare pays for nursing care as posthospital SNF care when provided by or under the supervision of a registered professional nurse.

(b) Exception. Medicare does not pay for the services of a private duty nurse or attendant. An individual is not considered to be a private duty nurse or attendant if he or she is an SNF employee at the time the services are furnished.

§ 409.22 Bed and board.

(a) Semiprivate and ward accommodations. Except for applicable deductible and coinsurance amounts Medicare Part A pays in full for semiprivate (2 to 4 beds), or ward (5 or more beds) accommodations.

(b) Private accommodations—(1) Conditions for payment in full. Except for applicable coinsurance amounts, Medicare pays in full for a private room if—

(i) The patient’s condition requires him to be isolated;

(ii) The SNF has no semiprivate or ward accommodations; or

(iii) The SNF semiprivate and ward accommodations are fully occupied by other patients, were so occupied at the time the patient was admitted to the SNF for treatment of a condition that required immediate inpatient SNF care, and have been so occupied during the interval.

(2) Period of payment. In the situations specified in paragraph (b)(1)(i) and (iii) of this section, Medicare pays for a private room until the patient’s condition no longer requires isolation or until semiprivate or ward accommodations are available.

(c) Conditions for patient’s liability. The facility may charge the patient the difference between its customary charge for the private room furnished and its most prevalent charge for a semiprivate room if:

(i) None of the conditions of paragraph (b)(1) of this section is met, and

(ii) The private room was requested by the patient or a member of the family who, at the time of request was informed what the charge would be.

§ 409.23 Physical, occupational, and speech therapy.

Medicare pays for physical, occupational, or speech therapy as posthospital SNF care if—

(a) It is furnished by the facility or under arrangements made by the facility.

(b) Billing for the therapy is by or through the facility.

§ 409.24 Medical social services.

Medicare pays for medical social services as posthospital SNF care, including—
Health Care Financing Administration, HHS

§ 409.27 Other services generally provided by (or under arrangements made by) SNFs.

In addition to those services specified in §§ 409.21 through 409.26, Medicare pays as posthospital SNF care for such other diagnostic and therapeutic services as are generally provided by (or under arrangements made by) SNFs, including—

(a) Medical and other health services as described in subpart B of part 410 of this chapter, subject to any applicable limitations or exclusions contained in that subpart or in § 409.20(b);

(b) Respiratory therapy services prescribed by a physician for the assessment, diagnostic evaluation, treatment, management, and monitoring of patients with deficiencies and abnormalities of cardiopulmonary function; and—

§ 409.26 Transfer agreement hospital services.

(a) Services furnished by an intern or a resident-in-training. Medicare pays for medical services that are furnished by an intern or a resident-in-training (under a hospital teaching program approved in accordance with the provisions of § 409.15) as posthospital SNF care, if the intern or resident is in—

(1) A participating hospital with which the SNF has in effect an agreement under § 409.21 of this chapter for the transfer of patients and exchange of medical records; or

(2) A hospital that has a swing-bed approval, and is furnishing services to an SNF-level inpatient of that hospital.

(b) Other diagnostic or therapeutic services. Medicare pays for other diagnostic or therapeutic services as posthospital SNF care if they are provided—

(1) By a participating hospital with which the SNF has in effect a transfer agreement as described in paragraph (a)(1) of this section; or

(2) By a hospital or a CAH that has a swing-bed approval, to its own SNF-level inpatient.

§ 409.25 Drugs, biologicals, supplies, appliances, and equipment.

(a) Drugs and biologicals. Except as specified in paragraph (b) of this section, Medicare pays for drugs and biologicals as posthospital SNF care only if—

(1) They represent a cost to the facility;

(2) They are ordinarily furnished by the facility for the care and treatment of inpatients; and

(3) They are furnished to an inpatient for use in the facility.

(b) Exception. Medicare pays for a limited supply of drugs for use outside the facility if it is medically necessary to facilitate the beneficiary's departure from the facility and required until he or she can obtain a continuing supply.

(c) Supplies, appliances, and equipment. Except as specified in paragraph (d) of this section, Medicare pays for supplies, appliances, and equipment as posthospital SNF care only if they are—

(1) Ordinarily furnished by the facility to inpatients; and

(2) Furnished to inpatients for use in the facility.

(d) Exception. Medicare pays for items to be used after the individual leaves the facility if—

(1) The item is one that the beneficiary must continue to use after leaving, such as a leg brace; or

(2) The item is necessary to permit or facilitate the beneficiary's departure from the facility and is required until he or she can obtain a continuing supply, for example, sterile dressings.
§ 409.30 Basic requirements.

Posthospital SNF care, including SNF-type care furnished in a hospital or CAH that has a swing-bed approval, is covered only if the beneficiary meets the requirements of this section and only for days when he or she needs and receives care of the level described in § 409.31. A beneficiary in an SNF is also considered to meet the level of care requirements of § 409.31 up to and including the assessment reference date for the 5-day assessment prescribed in § 413.343(b) of this chapter, when assigned to one of the Resource Utilization Groups that is designated (in the annual publication of Federal prospective payment rates described in § 413.345 of this chapter) as representing the required level of care. For the purposes of this section, the assessment reference date is defined in accordance with § 483.315(d) of this chapter, and must occur no later than the eighth day of posthospital SNF care.

(a) Pre-admission requirements. The beneficiary must—

(1) Have been hospitalized in a participating or qualified hospital or participating CAH, for medically necessary inpatient hospital or inpatient CAH care, for at least 3 consecutive calendar days, not counting the date of discharge; and

(2) Have been discharged from the hospital or CAH in or after the month he or she attained age 65, or in a month for which he or she was entitled to hospital or CAH insurance benefits on the basis of disability or end-stage renal disease, in accordance with part 406 of this chapter.

(b) Date of admission requirements.

(1) Except as specified in paragraph (b)(2)

(2) The beneficiary must be in need of posthospital SNF care, be admitted to the facility, and receive the needed care within 30 calendar days after the date of discharge from a hospital or CAH.

(2) Exception. A beneficiary for whom posthospital SNF care would not be medically appropriate within 30 days after discharge from the hospital or CAH may be admitted at the time it would be medically appropriate to begin an active course of treatment.

§ 409.31 Level of care requirements.

(a) Definition. As used in this section, skilled nursing and skilled rehabilitation services means services that:

(1) Are ordered by a physician;

(2) Require the skills of technical or professional personnel such as registered nurses, licensed practical (vocational) nurses, physical therapists, occupational therapists, and speech pathologists or audiologists; and

(3) Are furnished directly by, or under the supervision of, such personnel.

(b) Specific conditions for meeting level of care requirements.

(1) The beneficiary must require skilled nursing or skilled rehabilitation services, or both, on a daily basis.

(2) Those services must be furnished for a condition—

(i) For which the beneficiary received inpatient hospital or inpatient CAH care; or

(ii) Which arose while the beneficiary was receiving care in a SNF or swing-bed hospital for a condition for which he or she received inpatient hospital or inpatient CAH services.

(3) The daily skilled services must be ones that, as a practical matter, can

14 days after discharge from the hospital or CAH and permitted admission up to 28 days after discharge if a SNF bed was not available in the geographic area in which the patient lived, or at the time it would be medically appropriate to begin an active course of treatment, if SNF care would not be medically appropriate within 14 days after discharge.
only be provided in a SNF, on an inpatient basis.

§ 409.32 Criteria for skilled services and the need for skilled services.

(a) To be considered a skilled service, the service must be so inherently complex that it can be safely and effectively performed only by, or under the supervision of, professional or technical personnel.

(b) A condition that does not ordinarily require skilled services may require them because of special medical complications. Under those circumstances, a service that is usually nonskilled (such as those listed in § 409.33(d)) may be considered skilled because it must be performed or supervised by skilled nursing or rehabilitation personnel. For example, a plaster cast on a leg does not usually require skilled care. However, if the patient has a preexisting acute skin condition or needs traction, skilled personnel may be needed to adjust traction or watch for complications. In situations of this type, the complications, and the skilled services they require, must be documented by physicians' orders and nursing or therapy notes.

(c) The restoration potential of a patient is not the deciding factor in determining whether skilled services are needed. Even if full recovery or medical improvement is not possible, a patient may need skilled services to prevent further deterioration or preserve current capabilities. For example, a terminal cancer patient may need some of the skilled services described in § 409.33.

§ 409.33 Examples of skilled nursing and rehabilitation services.

(a) Services that could qualify as either skilled nursing or skilled rehabilitation services—(1) Overall management and evaluation of care plan. (i) When overall management and evaluation of care plan constitute skilled services. The development, management, and evaluation of a patient care plan based on the physician's orders constitute skilled services when, because of the patient's physical or mental condition, those activities require the involvement of technical or professional personnel in order to meet the patient's needs, promote recovery, and ensure medical safety. Those activities include the management of a plan involving a variety of personal care services only when, in light of the patient's condition, the aggregate of those services requires the involvement of technical or professional personnel.

(ii) Example. An aged patient with a history of diabetes mellitus and angina pectoris who is recovering from an open reduction of a fracture of the neck of the femur requires, among other services, careful skin care, appropriate oral medications, a diabetic diet, an exercise program to preserve muscle tone and body condition, and observation to detect signs of deterioration in his or her condition or complications resulting from restricted, but increasing, mobility. Although any of the required services could be performed by a properly instructed person, such a person would not have the ability to understand the relationship between the services and evaluate the ultimate effect of one service on the other. Since the nature of the patient's condition, age, and immobility create a high potential for serious complications, such an understanding is essential to ensure the patient's recovery and safety. Under these circumstances, the management of the plan of care would require the skills of a nurse even though the individual services are not skilled. Skilled planning and management activities are not always specifically identified in the patient's clinical record. Therefore, if the patient's overall condition supports a finding that recovery and safety can be ensured only if the total care is planned, managed, and evaluated by technical or professional personnel, it is appropriate to infer that skilled services are being provided.

(2) Observation and assessment of the patient's changing condition—(i) When observation and assessment constitute skilled services. Observation and assessment constitute skilled services when the skills of a technical or professional person are required to identify and
evaluate the patient's need for modification of treatment or for additional medical procedures until his or her condition is stabilized.

(ii) Examples. A patient with congestive heart failure may require continuous close observation to detect signs of decompensation, abnormal fluid balance, or adverse effects resulting from prescribed medication(s) that serve as indicators for adjusting therapeutic measures. Similarly, surgical patients transferred from a hospital to an SNF while in the complicated, unstabilized postoperative period, for example, after hip prosthesis or cataract surgery, may need continued close skilled monitoring for postoperative complications and adverse reaction. Patients who, in addition to their physical problems, exhibit acute psychological symptoms such as depression, anxiety, or agitation, may also require skilled observation and assessment by technical or professional personnel to ensure their safety or the safety of others, that is, to observe for indications of suicidal or hostile behavior. The need for services of this type must be documented by physicians' orders or nursing or therapy notes.

(3) Patient education services—(i) When patient education services constitute skilled services. Patient education services are skilled services if the use of technical or professional personnel is necessary to teach a patient self-maintenance.

(ii) Examples. A patient who has had a recent leg amputation needs skilled rehabilitation services provided by technical or professional personnel to provide gait training and to teach prosthesis care. Similarly, a patient newly diagnosed with diabetes requires instruction from technical or professional personnel to learn the self-administration of insulin or foot-care precautions.

(b) Services that qualify as skilled nursing services. (1) Intravenous or intramuscular injections and intravenous feeding.

(2) Enteral feeding that comprises at least 26 per cent of daily calorie requirements and provides at least 501 milliliters of fluid per day.

(3) Nasopharyngeal and tracheostomy aspiration;

(4) Insertion and sterile irrigation and replacement of suprapubic catheters;

(5) Application of dressings involving prescription medications and aseptic techniques;

(6) Treatment of extensive decubitus ulcers or other widespread skin disorder;

(7) Heat treatments which have been specifically ordered by a physician as part of active treatment and which require observation by nurses to adequately evaluate the patient's progress;

(8) Initial phases of a regimen involving administration of medical gases;

(9) Rehabilitation nursing procedures, including the related teaching and adaptive aspects of nursing, that are part of active treatment, e.g., the institution and supervision of bowel and bladder training programs.

(c) Services which would qualify as skilled rehabilitation services. (1) Ongoing assessment of rehabilitation needs and potential: Services concurrent with the management of a patient care plan, including tests and measurements of range of motion, strength, balance, coordination, endurance, functional ability, activities of daily living, perceptual deficits, speech and language or hearing disorders;

(2) Therapeutic exercises or activities: Therapeutic exercises or activities which, because of the type of exercises employed or the condition of the patient, must be performed by or under the supervision of a qualified physical therapist or occupational therapist to ensure the safety of the patient and the effectiveness of the treatment;

(3) Gait evaluation and training: Gait evaluation and training furnished to restore function in a patient whose ability to walk has been impaired by neurological, muscular, or skeletal abnormality;

(4) Range of motion exercises: Range of motion exercises which are part of the active treatment of a specific disease state which has resulted in a loss of, or restriction of, mobility (as evidenced by a therapist's notes showing the degree of motion lost and the degree to be restored);

(5) Maintenance therapy; Maintenance therapy, when the specialized
knowledge and judgment of a qualified therapist is required to design and establish a maintenance program based on an initial evaluation and periodic reassessment of the patient's needs, and consistent with the patient's capacity and tolerance. For example, a patient with Parkinson's disease who has not been under a rehabilitation regimen may require the services of a qualified therapist to determine what type of exercises will contribute the most to the maintenance of his present level of functioning.

(6) Ultrasound, short-wave, and microwave therapy treatment by a qualified physical therapist;

(7) Hot pack, hydrocollator, infrared treatments, paraffin baths, and whirlpool; Hot pack hydrocollator, infrared treatments, paraffin baths, and whirlpool in particular cases where the patient's condition is complicated by circulatory deficiency, areas of desensitization, open wounds, fractures, or other complications, and the skills, knowledge, and judgment of a qualified physical therapist are required; and

(8) Services of a speech pathologist or audiologist when necessary for the restoration of function in speech or hearing.

(d) Personal care services. Personal care services which do not require the skills of qualified technical or professional personnel are not skilled services except under the circumstances specified in §409.32(b). Personal care services include, but are not limited to, the following:

(1) Administration of routine oral medications, eye drops, and ointments;

(2) General maintenance care of colostomy and ileostomy;

(3) Routine services to maintain satisfactory functioning of indwelling bladder catheters;

(4) Changes of dressings for non-infected postoperative or chronic conditions;

(5) Prophylactic and palliative skin care, including bathing and application of creams, or treatment of minor skin problems;

(6) Routine care of the incontinent patient, including use of diapers and protective sheets;

(7) General maintenance care in connection with a plaster cast;

(8) Routine care in connection with braces and similar devices;

(9) Use of heat as a palliative and comfort measure, such as whirlpool and hydrocollator;

(10) Routine administration of medical gases after a regimen of therapy has been established;

(11) Assistance in dressing, eating, and going to the toilet;

(12) Periodic turning and positioning in bed; and

(13) General supervision of exercises which have been taught to the patient; including the actual carrying out of maintenance programs, i.e., the performance of the repetitive exercises required to maintain function do not require the skills of a therapist and would not constitute skilled rehabilitation services (see paragraph (c) of this section). Similarly, repetitious exercises to improve gait, maintain strength, or endurance; passive exercises to maintain range of motion in paralyzed extremities, which are not related to a specific loss of function; and assistive walking do not constitute skilled rehabilitation services.


§ 409.34 Criteria for “daily basis”.

(a) To meet the daily basis requirement specified in §409.31(b)(1), the following frequency is required:

(1) Skilled nursing services or skilled rehabilitation services must be needed and provided 7 days a week; or

(2) As an exception, if skilled rehabilitation services are not available 7 days a week those services must be needed and provided at least 5 days a week.

(b) A break of one or two days in the furnishing of rehabilitation services will not preclude coverage if discharge would not be practical for the one or two days during which, for instance, the physician has suspended the therapy sessions because the patient exhibited extreme fatigue.

§ 409.35 Criteria for “practical matter”.

(a) General considerations. In making a “practical matter” determination, as required by §409.31(b)(3), consideration
§ 409.36 Effect of discharge from posthospital SNF care.

If a beneficiary is discharged from a facility after receiving posthospital SNF care, he or she is not entitled to additional services of this kind in the same benefit period unless—

(a) He or she is readmitted to the same or another facility within 30 calendar days following the day of discharge (or, before December 5, 1980, within 14 calendar days after discharge); or

(b) He or she is again hospitalized for at least 3 consecutive calendar days.

Subpart E—Home Health Services Under Hospital Insurance

§ 409.40 Basis, purpose, and scope.

This subpart implements sections 1814(a)(2)(C), 1835(a)(2)(A), and 1861(m) of the Act with respect to the requirements that must be met for Medicare payment to be made for home health services furnished to eligible beneficiaries.

[59 FR 65493, Dec. 20, 1994]

§ 409.41 Requirement for payment.

In order for home health services to qualify for payment under the Medicare program the following requirements must be met:

(a) The services must be furnished to an eligible beneficiary by, or under arrangements with, an HHA that—

(1) Meets the conditions of participation for HHAs at part 484 of this chapter; and

(2) Has in effect a Medicare provider agreement as described in part 489, subparts A, B, C, D, and E of this chapter.

(b) The physician certification and recertification requirements for home health services described in §424.22.

(c) All requirements contained in §§409.42 through 409.47.

[59 FR 65494, Dec. 20, 1994]

§ 409.42 Beneficiary qualifications for coverage of services.

To qualify for Medicare coverage of home health services, a beneficiary must meet each of the following requirements:

(a) Confined to the home. The beneficiary must be confined to the home or in an institution that is not a hospital, SNF or nursing facility as defined in section 1861(e)(1), 1819(a)(1) or 1919(a)(1) of the Act, respectively.

(b) Under the care of a physician. The beneficiary must be under the care of a physician who establishes the plan of care. A doctor of podiatric medicine may establish a plan of care only if that is consistent with the functions he or she is authorized to perform under State law.

(c) In need of skilled services. The beneficiary must meet at least one of the following skilled services as certified by a physician in accordance with the physician certification and recertification requirements for home health services under §424.22 of this chapter.

(1) Intermittent skilled nursing services that meet the criteria for skilled services and the need for skilled services found in §409.32. (Also see...
§ 409.33(a) and (b) for a description of examples of skilled nursing and rehabilitation services.)

(2) Physical therapy services that meet the requirements of § 409.44(c).

(3) Speech-language pathology services that meet the requirements of § 409.44(c).

(4) Continuing occupational therapy services that meet the requirements of § 409.44(c) if the beneficiary's eligibility for home health services has been established by virtue of a prior need for intermittent skilled nursing care, speech-language pathology services, or physical therapy in the current or prior certification period.

(d) Under a plan of care. The beneficiary must be under a plan of care that meets the requirements for plans of care specified in § 409.43.

(e) By whom the services must be furnished. The home health services must be furnished by, or under arrangements made by, a participating HHA.

§ 409.43 Plan of care requirements.

(a) Contents. The plan of care must contain those items listed in § 484.18(a) of this chapter that specify the standards relating to a plan of care that an HHA must meet in order to participate in the Medicare program.

(b) Physician's orders. The physician's orders for services in the plan of care must specify the medical treatments to be furnished as well as the type of home health discipline that will furnish the ordered services and at what frequency the services will be furnished. Orders for services to be provided “as needed” or “PRN” must be accompanied by a description of the beneficiary's medical signs and symptoms that would occasion the visit and a specific limit on the number of those visits to be made under the order before an additional physician order would have to be obtained. Orders for care may indicate a specific range in frequency of visits to ensure that the most appropriate level of services is furnished. If a range of visits is ordered, the upper limit of the range is considered the specific frequency.

(c) Physician signature. (1) Request for Anticipated payment signature requirements. If the physician signed plan of care is not available at the time the HHA requests an anticipated payment of the initial percentage prospective payment in accordance with § 484.205, the request for the anticipated payment must be based on—

(i) A physician's verbal order that—

(A) is recorded in the plan of care;

(B) includes a description of the patient's condition and the services to be provided by the home health agency;

(C) includes an attestation (relating to the physician's orders and the date received) signed and dated by the registered nurse or qualified therapist (as defined in 42 CFR 484.4) responsible for furnishing or supervising the ordered service in the plan of care, and

(ii) A referral prescribing detailed orders for the services to be rendered that is signed and dated by a physician.

(2) Reduction or disapproval of anticipated payment requests. HCFA has the authority to reduce or disapprove requests for anticipated payments in situations when protecting Medicare program integrity warrants this action. Since the request for anticipated payment is based on verbal orders as specified in paragraph (c)(1)(i) and/or a prescribing referral as specified in (c)(1)(ii) of this section and is not a Medicare claim for purposes of the Act (although it is a “claim” for purposes of Federal, civil, criminal, and administrative law enforcement authorities, including but not limited to the Civil Monetary Penalties Law (as defined in 42 U.S.C. 1320a-7a (i) (2)), the Civil False Claims Act (as defined in 31 U.S.C. 3729(c)), and the Criminal False Claims Act (18 U.S.C. 287)), the request for anticipated payment will be canceled and recovered unless the claim is submitted within the greater of 60 days from the end of the episode or 60 days from the issuance of the request for anticipated payment.

(3) Final percentage payment signature requirements. The plan of care must be signed and dated—

(i) By a physician as described who meets the certification and recertification requirements of § 424.22 of this chapter; and
(ii) Before the claim for each episode for services is submitted for the final percentage prospective payment.

(4) Changes to the plan of care signature requirements. Any changes in the plan must be signed and dated by a physician.

(d) Oral (verbal) orders. If any services are provided based on a physician’s oral orders, the orders must be put in writing and be signed and dated with the date of receipt by the registered nurse or qualified therapist (as defined in § 484.4 of this chapter) responsible for furnishing or supervising the ordered services. Oral orders may only be accepted by personnel authorized to do so by applicable State and Federal laws and regulations as well as by the HHA’s internal policies. The oral orders must also be countersigned and dated by the physician before the HHA bills for the care.

(e) Frequency of review. (1) The plan of care must be reviewed by the physician (as specified in § 409.42(b)) in consultation with agency professional personnel at least every 60 days or more frequently when there is a—
   (i) Beneficiary elected transfer;
   (ii) Significant change in condition resulting in a change in the case-mix assignment; or
   (iii) Discharge and return to the same HHA during the 60-day episode.

   (2) Each review of a beneficiary’s plan of care must contain the signature of the physician who reviewed it and the date of review.

(f) Termination of the plan of care. The plan of care is considered to be terminated if the beneficiary does not receive at least one covered skilled nursing, physical therapy, speech-language pathology services, or occupational therapy visit in a 60-day period unless the physician documents that the interval without such care is appropriate to the treatment of the beneficiary’s illness or injury.


§ 409.44 Skilled services requirements.

(a) General. The intermediary’s decision on whether care is reasonable and necessary is based on information provided on the forms and in the medical record concerning the unique medical condition of the individual beneficiary. A coverage denial is not made solely on the basis of the reviewer’s general inferences about patients with similar diagnoses or on data related to utilization generally but is based upon objective clinical evidence regarding the beneficiary’s individual need for care.

(b) Skilled nursing care. (1) Skilled nursing care consists of those services that must, under State law, be performed by a registered nurse, or practical (vocational) nurse, as defined in § 484.4 of this chapter, and meet the criteria for skilled nursing services specified in § 409.32. See § 409.33(a) and (b) for a description of skilled nursing services and examples of them.

   (i) In determining whether a service requires the skill of a licensed nurse, consideration must be given to the inherent complexity of the service, the condition of the beneficiary, and accepted standards of medical and nursing practice.

   (ii) If the nature of a service is such that it can safely and effectively be performed by the average nonmedical person without direct supervision of a licensed nurse, the service cannot be regarded as a skilled nursing service.

   (iii) The fact that a skilled nursing service can be or is taught to the beneficiary or to the beneficiary’s family or friends does not negate the skilled aspect of the service when performed by the nurse.

   (iv) If the service could be performed by the average nonmedical person, the absence of a competent person to perform it does not cause it to be a skilled nursing service.

   (2) The skilled nursing care must be provided on a part-time or intermittent basis.

   (3) The skilled nursing services must be reasonable and necessary for the treatment of the illness or injury.

   (i) To be considered reasonable and necessary, the services must be consistent with the nature and severity of the beneficiary’s illness or injury, his or her particular medical needs, and accepted standards of medical and nursing practice.

   (ii) The skilled nursing care provided to the beneficiary must be reasonable within the context of the beneficiary’s condition.
(iii) The determination of whether skilled nursing care is reasonable and necessary must be based solely upon the beneficiary's unique condition and individual needs, without regard to whether the illness or injury is acute, chronic, terminal, or expected to last a long time.

(c) Physical therapy, speech-language pathology services, and occupational therapy. To be covered, physical therapy, speech-language pathology services, and occupational therapy must satisfy the criteria in paragraphs (c)(1) through (4) of this section. Occupational therapy services initially qualify for home health coverage only if they are part of a plan of care that also includes intermittent skilled nursing care, physical therapy, or speech-language pathology services as follows:

(1) Speech-language pathology services and physical or occupational therapy services must relate directly and specifically to a treatment regimen (established by the physician, after any needed consultation with the qualified therapist) that is designed to treat the beneficiary's illness or injury. Services related to activities for the general physical welfare of beneficiaries (for example, exercises to promote overall fitness) do not constitute physical therapy, occupational therapy, or speech-language pathology services for Medicare purposes.

(2) Physical and occupational therapy and speech-language pathology services must be reasonable and necessary. To be considered reasonable and necessary, the following conditions must be met:

(i) The services must be considered under accepted standards of medical practice to be a specific, safe, and effective treatment for the beneficiary's condition.

(ii) The services must be of such a level of complexity and sophistication or the condition of the beneficiary must be such that the services required can safely and effectively be performed only by a qualified physical therapist or by a qualified physical therapy assistant under the supervision of a qualified physical therapist, by a qualified speech-language pathologist, or by a qualified occupational therapist or a qualified occupational therapy assistant under the supervision of a qualified occupational therapist (as defined in §484.4 of this chapter). Services that do not require the performance or supervision of a physical therapist or occupational therapist are not considered reasonable or necessary physical therapy or occupational therapy services, even if they are performed by or supervised by a physical therapist or occupational therapist. Services that do not require the skills of a speech-language pathologist are not considered to be reasonable and necessary speech-language pathology services even if they are performed by or supervised by a speech-language pathologist.

(iii) There must be an expectation that the beneficiary's condition will improve materially in a reasonable (and generally predictable) period of time based on the physician's assessment of the beneficiary's restoration potential and unique medical condition, or the services must be necessary to establish a safe and effective maintenance program required in connection with a specific disease, or the skills of a therapist must be necessary to perform a safe and effective maintenance program. If the services are for the establishment of a maintenance program, they may include the design of the program, the instruction of the beneficiary, family, or home health aides, and the necessary infrequent re-evaluations of the beneficiary and the program to the degree that the specialized knowledge and judgment of a physical therapist, speech-language pathologist, or occupational therapist is required.

(iv) The amount, frequency, and duration of the services must be reasonable.

[59 FR 65494, Dec. 20, 1994]
beneficiary’s eligibility for home health services has been established by virtue of a prior need for intermittent skilled nursing care, speech-language pathology services, or physical therapy in the current or prior certification period; and otherwise meets the qualifying criteria (confined to the home, under the care of a physician, in need of skilled services, and under a plan of care) specified in §409.42. Home health coverage is not available for services furnished to a beneficiary who is no longer in need of one of the qualifying skilled services specified in this paragraph. Therefore, dependent services furnished after the final qualifying skilled service are not covered, except when the dependent service was not followed by a qualifying skilled service as a result of the unexpected inpatient admission or death of the beneficiary, or due to some other unanticipated event.

(b) Home health aide services. To be covered, home health aide services must meet each of the following requirements:

(1) The reason for the visits by the home health aide must be to provide hands-on personal care to the beneficiary or services that are needed to maintain the beneficiary’s health or to facilitate treatment of the beneficiary’s illness or injury. The physician’s order must indicate the frequency of the home health aide services required by the beneficiary. These services may include but are not limited to:

(i) Personal care services such as bathing, dressing, grooming, caring for hair, nail and oral hygiene that are needed to facilitate treatment or to prevent deterioration of the beneficiary’s health, changing the bed linens of an incontinent beneficiary, shaving, deodorant application, skin care with lotions and/or powder, foot care, ear care, feeding, assistance with elimination (including enemas unless the skills of a licensed nurse are required due to the beneficiary’s condition, routine catheter care, and routine colostomy care), assistance with ambulation, changing position in bed, and assistance with transfers.

(ii) Simple dressing changes that do not require the skills of a licensed nurse.

(iii) Assistance with medications that are ordinarily self-administered and that do not require the skills of a licensed nurse to be provided safely and effectively.

(iv) Assistance with activities that are directly supportive of skilled therapy services but do not require the skills of a therapist to be safely and effectively performed, such as routine maintenance exercises and repetitive practice of functional communication skills to support speech-language pathology services.

(v) Routine care of prosthetic and orthotic devices.

(2) The services to be provided by the home health aide must be—

(i) Ordered by a physician in the plan of care; and

(ii) Provided by the home health aide on a part-time or intermittent basis.

(3) The services provided by the home health aide must be reasonable and necessary. To be considered reasonable and necessary, the services must—

(i) Meet the requirement for home health aide services in paragraph (b)(1) of this section;

(ii) Be of a type the beneficiary cannot perform for himself or herself; and

(iii) Be of a type that there is no able or willing caregiver to provide, or, if there is a potential caregiver, the beneficiary is unwilling to use the services of that individual.

(4) The home health aide also may perform services incidental to a visit that was for the provision of care as described in paragraphs (b)(3)(i) through (iii) of this section. For example, these incidental services may include changing bed linens, personal laundry, or preparing a light meal.

(c) Medical social services. Medical social services may be covered if the following requirements are met:

(1) The services are ordered by a physician and included in the plan of care.

(2)(i) The services are necessary to resolve social or emotional problems that are expected to be an impediment to the effective treatment of the beneficiary’s medical condition or to his or her rate of recovery.
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(ii) If these services are furnished to a beneficiary's family member or caregiver, they are furnished on a short-term basis and it can be demonstrated that the service is necessary to resolve a clear and direct impediment to the effective treatment of the beneficiary's medical condition or to his or her rate of recovery.

(3) The frequency and nature of the medical social services are reasonable and necessary to the treatment of the beneficiary's condition:

(4) The medical social services are furnished by a qualified social worker or qualified social work assistant under the supervision of a social worker as defined in § 484.4 of this chapter.

(5) The services needed to resolve the problems that are impeding the beneficiary's recovery require the skills of a social worker or a social work assistant under the supervision of a social worker to be performed safely and effectively.

(d) Occupational therapy. Occupational therapy services that are not qualifying services under § 409.44(c) are nevertheless covered as dependent services if the requirements of § 409.44(c)(2)(i) through (iv), as to reasonableness and necessity, are met.

(e) Durable medical equipment. Durable medical equipment in accordance with § 410.38 of this chapter, which describes the scope and conditions of payment for durable medical equipment under Part B, may be covered under the home health benefit as either a Part A or Part B service. Durable medical equipment furnished by an HHA as a home health service is always covered by Part A if the beneficiary is entitled to Part A.

(f) Medical supplies. Medical supplies (including catheters, catheter supplies, ostomy bags, and supplies relating to ostomy care but excluding drugs and biologicals) may be covered as a home health benefit. For medical supplies to be covered as a Medicare home health benefit, the medical supplies must be needed to treat the beneficiary's illness or injury that occasioned the home health care.

(g) Intern and resident services. The medical services of interns and residents in training under an approved hospital teaching program are covered if the services are ordered by the physician who is responsible for the plan of care and the HHA is affiliated with or under the common control of the hospital furnishing the medical services.

Approved means—

(1) Approved by the Accreditation Council for Graduate Medical Education;

(2) In the case of an osteopathic hospital, approved by the Committee on Hospitals of the Bureau of Professional Education of the American Osteopathic Association;

(3) In the case of an intern or resident-in-training in the field of dentistry, approved by the Council on Dental Education of the American Dental Association;

(4) In the case of an intern or resident-in-training in the field of podiatry, approved by the Council on Podiatric Medical Education of the American Podiatric Medical Association.

[59 FR 65495, Dec. 20, 1994; 60 FR 39122, 39123, Aug. 1, 1995]

§ 409.46 Allowable administrative costs.

Services that are allowable as administrative costs but are not separately billable include, but are not limited to, the following:

(a) Registered nurse initial evaluation visits. Initial evaluation visits by a registered nurse for the purpose of assessing a beneficiary's health needs, determining if the agency can meet those health needs, and formulating a plan of care for the beneficiary are allowable administrative costs. If a physician specifically orders that a particular skilled service be furnished during the evaluation in which the agency accepts the beneficiary for treatment and all other coverage criteria are met, the visit is billable as a skilled nursing visit. Otherwise it is considered to be an administrative cost.

(b) Visits by registered nurses or qualified professionals for the supervision of home health aides. Visits by registered nurses or qualified professionals for the purpose of supervising home health aides as required at § 484.36(d) of this chapter are allowable administrative costs. Only if the registered nurse or
qualified professional visits the beneficiary for the purpose of furnishing care that meets the coverage criteria at §409.44, and the supervisory visit occurs simultaneously with the provision of covered care, is the visit billable as a skilled nursing or therapist's visit.

(c) Respiratory care services. If a respiratory therapist is used to furnish overall training or consultative advice to an HHA's staff and incidentally provides respiratory therapy services to beneficiaries in their homes, the costs of the respiratory therapist's services are allowable as administrative costs. Visits by a respiratory therapist to a beneficiary's home are not separately billable. However, respiratory therapy services that are furnished as part of a plan of care by a skilled nurse or physical therapist and that constitute skilled care may be separately billed as skilled visits.

(d) Dietary and nutrition personnel. If dieticians or nutritionists are used to provide overall training or consultative advice to HHA staff and incidentally provide dietetic or nutritional services to beneficiaries in their homes, the costs of these professional services are allowable as administrative costs. Visits by a dietician or nutritionist to a beneficiary's home are not separately billable.

§409.48 Visits.

(a) Number of allowable visits under Part A. To the extent that all coverage requirements specified in this subpart are met, payment may be made on behalf of eligible beneficiaries under Part A for an unlimited number of covered home health visits. Medicare home health services are covered under hospital insurance unless there is no Part A entitlement.

(b) Number of visits under Part B. To the extent that all coverage requirements specified in this subpart are met, payment may be made on behalf of eligible beneficiaries under Part B for an unlimited number of covered home health visits. Medicare home health services are covered under Part B only when the beneficiary is not entitled to coverage under Part A.

(c) Definition of visit. A visit is an episode of personal contact with the beneficiary by staff of the HHA or others under arrangements with the HHA, for the purpose of providing a covered service.

(1) Generally, one visit may be covered each time an HHA employee or someone providing home health services in an outpatient facility under arrangements with the facility, one visit may be covered.

(2) If two individuals are needed to provide a service, two visits may be covered. If two individuals are present, but only one is needed to provide the care, only one visit may be covered.

(3) If two individuals are needed to provide a service, two visits may be covered. If two individuals are present, but only one is needed to provide the care, only one visit may be covered.

(4) A visit is initiated with the delivery of covered home health services and ends at the conclusion of delivery of covered home health services in those circumstances in which all reasonable and necessary home health...
services cannot be provided in the course of a single visit, HHA staff or others providing services under arrangements with the HHA may remain at the beneficiary’s residence between visits (for example, to provide non-covered services). However, if all covered services could be provided in the course of one visit, only one visit may be covered.

[59 FR 65497, Dec. 20, 1994]

§ 409.49 Excluded services.

(a) Drugs and biologicals. Drugs and biologicals are excluded from payment under the Medicare home health benefit.

(1) A drug is any chemical compound that may be used on or administered to humans or animals as an aid in the diagnosis, treatment or prevention of disease or other condition or for the relief of pain or suffering or to control or improve any physiological pathologic condition.

(2) A biological is any medicinal preparation made from living organisms and their products including, but not limited to, serums, vaccines, antigens, and antitoxins.

(b) Transportation. The transportation of beneficiaries, whether to receive covered care or for other purposes, is excluded from home health coverage. Costs of transportation of equipment, materials, supplies, or staff may be allowable as administrative costs, but no separate payment is made for them.

(c) Services that would not be covered as inpatient services. Services that would not be covered if furnished as inpatient hospital services are excluded from home health coverage.

(d) Housekeeping services. Services whose sole purpose is to enable the beneficiary to continue residing in his or her home (for example, cooking, shopping, Meals on Wheels, cleaning, laundry) are excluded from home health coverage.

(e) Services covered under the End Stage Renal Disease (ESRD) program. Services that are covered under the ESRD program and are contained in the composite rate reimbursement methodology, including any service furnished to a Medicare ESRD beneficiary that is directly related to that individual’s dialysis, are excluded from coverage under the Medicare home health benefit.

(f) Prosthetic devices. Items that meet the requirements of §410.36(a)(2) of this chapter for prosthetic devices covered under Part B are excluded from home health coverage. Catheters, catheter supplies, ostomy bags, and supplies relating to ostomy care are not considered prosthetic devices if furnished under a home health plan of care and are not subject to this exclusion from coverage.

(g) Medical social services provided to family members. Except as provided in §409.45(c)(2), medical social services provided solely to members of the beneficiary’s family and that are not incidental to covered medical social services being provided to the beneficiary are not covered.

[59 FR 65497, Dec. 20, 1994; 60 FR 39123, Aug. 1, 1995]

§ 409.50 Coinsurance for durable medical equipment (DME) furnished as a home health service.

The coinsurance liability of the beneficiary or other person for DME furnished as a home health service is 20 percent of the customary (insofar as reasonable) charge for the services.


Subpart F—Scope of Hospital Insurance Benefits

§ 409.60 Benefit periods.

(a) When benefit periods begin. The initial benefit period begins on the day the beneficiary receives inpatient hospital, inpatient CAH, or SNF services for the first time after becoming entitled to hospital insurance. Thereafter, a new benefit period begins whenever the beneficiary receives inpatient hospital, inpatient CAH, or SNF services after he or she has ended a benefit period as described in paragraph (b) of this section.

(b) When benefit periods end—(1) A benefit period ends when a beneficiary has, for at least 60 consecutive days not been an inpatient in any of the following:
§ 409.60

(i) A hospital that meets the requirements of section 1861(e)(1) of the Act.

(ii) A CAH that meets the requirements of section 1861(y) of the Act.

(iii) A SNF that meets the requirements of sections 1819(a)(1) or 1861(y) of the Act.

(2) For purposes of ending a benefit period, a beneficiary was an inpatient of a SNF if his or her care in the SNF met the skilled level of care requirements specified in § 409.31(b)(1) and (3).

(c) Presumptions. (1) For purposes of determining whether a beneficiary was an inpatient of a SNF under paragraph (b)(2) of this section—

(i) A beneficiary's care met the skilled level of care requirements if inpatient SNF claims were paid for those services under Medicare or Medicaid, unless:

(A) Such payments were made under § 405.330 or Medicaid administratively necessary days provisions which result in payment for care not meeting the skilled level of care requirements, or

(B) A Medicare denial and a Medicaid payment are made for the same period, in which case the presumption in paragraph (c)(2)(ii) of this section applies;

(ii) A beneficiary's care met the skilled level of care requirements if a SNF claim was paid under section 1879(e) of the Social Security Act;

(iii) A beneficiary's care did not meet the skilled level of care requirements if a SNF claim was paid under section 1879(e) of the Social Security Act;

(iv) A beneficiary's care did not meet the skilled level of care requirements if a Medicaid SNF claim was denied on the ground that the services were not at the skilled level of care and payment was not made under § 405.330; or

(iv) Not to have met the skilled level of care requirements if no Medicare or Medicaid claim was submitted by the SNF.

(3) If information upon which to base a presumption is not readily available, the intermediary may, at its discretion review the beneficiary's medical records to determine whether he or she was an inpatient of a SNF as set forth under paragraph (b)(2) of this section.

(4) When the intermediary makes a benefit period determination based upon paragraph (c)(1) of this section, the beneficiary may seek to reverse the benefit period determination by timely appealing the prior Medicare SNF claim determination under part 405, subpart G of this chapter, or the prior Medicaid SNF claim under part 431, subpart E of this chapter.

(5) When the intermediary makes a benefit period determination under paragraph (c)(2) of this section, the beneficiary will be notified of the basis for the determination, and of his or her right to present evidence to rebut the determination that the skilled level of care requirements specified in § 409.31(b)(1) and (b)(3) were or were not met on reconsideration and appeal under 42 CFR, part 405, subpart G of this chapter.

(d) Limitation on benefit period determinations. When the intermediary considers the same prior SNF stay of a particular beneficiary in making benefit period determinations for more than one inpatient Medicare claim—

(1) Medicare will recognize only the initial level of care characterization for that prior SNF stay (or if appealed under 42 CFR part 405, subpart G of this chapter, the level of care determined under appeal); or

(2) If part of a prior SNF stay has one level of care characterization and another part has another level of care.
§ 409.61 General limitations on amount of benefits.

(a) Inpatient hospital or inpatient CAH services. (1) Regular benefit days. Up to 90 days are available in each benefit period, subject to the limitations on days for psychiatric hospital services set forth in § 409.62 and 409.63.

(i) For the first 60 days (referred to in this subpart as full benefit days), Medicare pays the hospital or CAH for all covered services furnished the beneficiary, except for a deductible which is the beneficiary’s responsibility. (Section 409.82 specifies the requirements for the inpatient hospital deductible.)

(ii) For the next 30 days (referred to in this subpart as coinsurance days), Medicare pays for all covered services except for a daily coinsurance amount, which is the beneficiary’s responsibility. (Section 409.83 specifies the inpatient hospital coinsurance amounts.)

(2) Lifetime reserve days. Each beneficiary has a non-renewable lifetime reserve of 60 days of inpatient hospital or inpatient CAH services that he may draw upon whenever he is hospitalized for more than 90 days in a benefit period. Upon exhaustion of the regular benefit days, the reserve days will be used unless the beneficiary elects not to use them, as provided in § 409.65. For lifetime reserve days, Medicare pays for all covered services except for a daily coinsurance amount that is the beneficiary’s responsibility. (See § 409.83.)

(b) Posthospital SNF care furnished by a SNF, or by a hospital or a CAH with a swing-bed approval. Up to 100 days are available in each benefit period after discharge from a hospital or CAH. For the first 20 days, Medicare pays for all covered services. For the 21st through 100th day, Medicare pays for all covered services except for a daily coinsurance amount that is the beneficiary’s responsibility.

(c) Renewal of inpatient benefits. The beneficiary’s full entitlement to the 90 inpatient hospital or inpatient CAH regular benefit days, and the 100 SNF benefit days, is renewed each time he or she begins a benefit period. However, once lifetime reserve days are used, they can never be renewed.

(d) Home health services. Medicare Part A pays for all covered home health services with no deductible, and subject to the following limitations on payment for durable medical equipment (DME):

(1) For DME furnished by an HHA that is a nominal charge provider, Medicare Part A pays 80 percent of fair compensation.

(2) For DME furnished by an HHA that is not a nominal charge provider, Medicare Part A pays the lesser of the following:

(i) 80 percent of the reasonable cost of the service.

(ii) The reasonable cost of, or the customary charge for, the service, whichever is less, minus 20 percent of the customary (insofar as reasonable) charge for the service.

§ 409.62 Lifetime maximum on inpatient psychiatric care.

There is a lifetime maximum of 190 days on inpatient psychiatric hospital care.
services available to any beneficiary. Therefore, once an individual receives benefits for 190 days of care in a psychiatric hospital, no further benefits of that type are available to that individual.

§ 409.63 Reduction of inpatient psychiatric benefit days available in the initial benefit period.

(a) Reduction rule. (1) If the individual was an inpatient in a psychiatric hospital on the first day of Medicare entitlement and for any of the 150 days immediately before that first day of entitlement, those days are subtracted from the 150 days (90 regular days plus 60 lifetime reserve days) which would otherwise be available in the initial benefit period for inpatient psychiatric services in a psychiatric or general hospital.

(2) Reduction is required only if the hospital was participating in Medicare as a psychiatric hospital on the individual's first day of entitlement.

(3) The reduction applies only to the beneficiary's first benefit period. For subsequent benefit periods, the 90 benefit days, plus any remaining lifetime reserve days, subject to the 190 day lifetime limit on psychiatric hospital care, are available.

(b) Application to general hospital days.

(1) Days spent in a general hospital before entitlement are not subtracted under paragraph (a) of this section even if the stay was for diagnosis or treatment of mental illness.

(2) After entitlement, all psychiatric care days, whether in a general or a psychiatric hospital, are counted toward the number of days available in the initial benefit period.

(c) Examples:

(1) The individual was an inpatient of a participating psychiatric hospital for 20 days before the first day of entitlement and remained there for another 6 months. Therefore, 130 days of benefits (150 minus 20) are payable. Payment could be made for: 60 full benefit days, 30 coinsurance days, and 40 lifetime reserve days.

(2) During the 150-day period preceding Medicare entitlement, an individual had been a patient of a general hospital for 60 days of inpatient psychiatric care and had spent 90 days in a psychiatric hospital, ending with the first day of entitlement. During the initial benefit period, the beneficiary spent 90 days in a general hospital and received psychiatric care there. The 60 days spent in the general hospital for psychiatric treatment before entitlement do not reduce the benefits available in the first benefit period. Only the 90 days spent in the psychiatric hospital before entitlement reduce such benefits, leaving a total of 60 available psychiatric days. However, after entitlement, the reduction applies not only to days spent in a psychiatric hospital, but also to days of psychiatric treatment in a general hospital. Thus, Medicare payment could be made only for 60 of the 90 days spent in the general hospital.

(3) An individual was admitted to a general hospital for a mental condition and, after 10 days, transferred to a participating psychiatric hospital. The individual remained in the psychiatric hospital for 78 days before becoming entitled to hospital insurance benefits and for 130 days after entitlement. The beneficiary was then transferred to a general hospital and received treatment of a medical condition for 20 days. The 10 days spent in the general hospital during the 150-day pre-entitlement period have no effect on the inpatient hospital benefit days available to the individual for psychiatric care in the first benefit period, even though the general hospital stay was for a mental condition. Only the 78 days spent in the psychiatric hospital during the pre-entitlement period are subtracted from the 150 benefit days. Accordingly, the individual has 72 days of psychiatric care (150 days less 78 days) available in the first benefit period. Benefits could be paid for the individual's hospitalization during the first benefit period in the following manner. For the 130-day psychiatric hospital stay, 72 days (60 full benefit days and 12 coinsurance days), and for the general hospital stay, 20 days (18 coinsurance and 2 lifetime reserve days).

§ 409.64 Services that are counted toward allowable amounts.

(a) Except as provided in paragraph (b) of this section for lifetime reserve days, all covered inpatient days and home health visits are counted toward

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§ 409.65 Lifetime reserve days.

(a) Election not to use lifetime reserve days. (1) Whenever a beneficiary has exhausted the 90 regular benefit days, the hospital or CAH may bill Medicare for lifetime reserve days unless the beneficiary elects not to use them or, in accordance with paragraph (b) of this section, is deemed to have elected not to use them.

(2) It may be advantageous to elect not to use lifetime reserve days if the beneficiary has private insurance coverage that begins after the first 90 inpatient days in a benefit period, or if the daily charge is only slightly higher than the lifetime reserve days coinsurance amount. In such cases, the beneficiary may want to save the lifetime reserve days for future care that may be more expensive.

(3) If the beneficiary elects not to use lifetime reserve days for a particular hospital or CAH stay, they are still available for a later stay. However, once the beneficiary uses lifetime reserve days, they can never be renewed.

(4) If the beneficiary elects not to use lifetime reserve days, the hospital or CAH may require him or her to pay for any services furnished after the regular days are exhausted.

(b) Deemed election. A beneficiary will be deemed to have elected not to use lifetime reserve days if the average daily charges for such days is equal to or less than the applicable coinsurance amount specified in § 409.63. A beneficiary would get no benefit from using the days under those circumstances.

(c) Who may file an election. An election not to use reserve days may be filed by—

(1) The beneficiary; or

(2) If the beneficiary is physically or mentally unable to act, by the beneficiary's legal representative. In addition, if some other payment source is available, such as private insurance, any person authorized under § 405.1664 of this chapter to execute a request for payment for the beneficiary may file the election.

(d) Filing the election. (1) The beneficiary's election not to use lifetime reserve days must be filed in writing with the hospital or CAH.

(2) The election may be filed at the time of admission to the hospital or CAH or at any time thereafter up to 90 days after the beneficiary's discharge.

(3) A retroactive election (that is, one made after lifetime reserve days have been used because the regular days were exhausted), is not acceptable unless it is approved by the hospital or CAH.

(e) Period covered by election—(1) General rule. Except as provided in paragraph (e)(2) of this section, an election not to use lifetime reserve days may apply to an entire hospital or CAH stay or to a single period of consecutive days in a stay, but cannot apply to selected days in a stay. For example, a beneficiary may restrict the election to the period covered by private insurance but cannot use individual lifetime reserve days within that period. If an election not to use reserve days is effective after the first day on which reserve days are available, it must remain in effect until the end of the stay, unless it is revoked in accordance with § 409.66.

(2) Exception. A beneficiary election not to use lifetime reserve days for an inpatient hospital or inpatient CAH stay for which payment may be made under the prospective payment system

§ 409.66 Revocation of election not to use lifetime reserve days.

(a) Except as provided in paragraph (c) of this section, a beneficiary (or anyone authorized to execute a request for payment, if the beneficiary is incapacitated) may revoke an election not to use lifetime reserve days during hospitalization or within 90 days after discharge.

(b) The revocation must be submitted to the hospital or CAH in writing and identify the stay or stays to which it applies.

(c) Exceptions. A revocation of an election not to use lifetime reserve days may not be filed—

(1) After the beneficiary dies; or

(2) After the hospital or CAH has filed a claim under the supplementary medical insurance program (Medicare Part B), for medical and other health services furnished to the beneficiary on the days in question.

§ 409.68 Guarantee of payment for inpatient hospital or inpatient CAH services furnished before notification of exhaustion of benefits.

(a) Conditions for payment. Payment may be made for inpatient hospital or inpatient CAH services furnished to a beneficiary after he or she has exhausted the available benefit days if the following conditions are met:

(1) The services were furnished before HCFA or the intermediary notified the hospital or CAH that the beneficiary had exhausted the available benefit days and was not entitled to have payment made for those services.

(2) At the time the hospital or CAH furnished the services, it was unaware that the beneficiary had exhausted the available benefit days and could reasonably have assumed that he or she was entitled to have payment made for these services.

(3) Payment would be precluded solely because the beneficiary has no benefit days available for the particular hospital or CAH stay.

(b) Limitations on payment. (1) If all of the conditions in paragraph (a) of this section are met, Medicare payment may be made for the day of admission, and up to 6 weekdays thereafter, plus any intervening Saturdays, Sundays, and Federal holidays.

(2) Payment may not be made under this section for any day after the hospital or CAH is notified that the beneficiary has exhausted the available benefit days.

(c) Recovery from the beneficiary. Any payment made to a hospital or CAH under this section is considered an overpayment to the beneficiary and may be recovered from him or her under the provisions set forth elsewhere in this chapter.
§ 409.82 Inpatient hospital deductible.  
(a) General provisions—(1) The inpatient hospital deductible is a fixed amount chargeable to the beneficiary when he or she receives covered services in a hospital or a CAH for the first time in a benefit period.  
(2) Although the beneficiary may be hospitalized several times during a benefit period, the deductible is charged only once during that period. If the beneficiary begins more than one benefit period in the same year, a deductible is charged for each of those periods.  
(3) For services furnished before January 1, 1982, the applicable deductible is the one in effect when the benefit period began.  
(4) For services furnished after December 31, 1981, the applicable deductible is the one in effect during the calendar year in which the services were furnished.  
(b) Specific deductible amounts. The specific deductible amounts for each calendar year are published in the Federal Register no later than October 1 of the preceding year.

§ 409.83 Inpatient hospital coinsurance.  
(a) General provisions—(1) Inpatient hospital coinsurance is the amount chargeable to a beneficiary for each day after the first 60 days of inpatient hospital care or inpatient CAH care or both in a benefit period.  
(2) For each day from the 61st to the 90th day, the coinsurance amount is $\frac{1}{4}$ of the applicable deductible.  
(3) For each day from the 91st to the 150th day (lifetime reserve days), the coinsurance amount is $\frac{1}{2}$ of the applicable deductible.  
(4) For coinsurance days before January 1, 1982, the coinsurance amount is based on the deductible applicable for the calendar year in which the benefit period began. The coinsurance amounts do not change during a beneficiary’s benefit period even though the coinsurance days may fall in a subsequent year for which a higher deductible amount has been determined.  
(5) For coinsurance days after December 31, 1981, the coinsurance amount is based on the deductible applicable for the calendar year in which the services were furnished. For example, if an individual starts a benefit period by being admitted to a hospital in 1981 and remains in the hospital long enough to use coinsurance days in 1982, the coinsurance amount charged for those days is based on the 1982 inpatient hospital deductible.  
(b) Specific coinsurance amounts. The specific coinsurance amounts for each calendar year are published in the Federal Register no later than October 1 of the preceding year.  
(c) Exceptions to published amounts. (1) If the actual charge to the patient for the 61st through the 90th day of inpatient hospital or inpatient CAH services is less than the coinsurance amount applicable for the calendar year in which the services were furnished, the actual charge per day is the daily coinsurance amount.
§ 409.85 Skilled nursing facility (SNF) care coinsurance.

(a) General provisions. (1) SNF care coinsurance is the amount chargeable to a beneficiary after the first 20 days of SNF care in a benefit period.

(2) For each day from the 21st through the 100th day, the coinsurance is 1/8 of the applicable inpatient hospital deductible.

(3) For coinsurance days before January 1, 1982, the coinsurance amount is based on the deductible applicable for the year in which the benefit period began. The coinsurance amounts do not change during a beneficiary’s benefit period even though the coinsurance days may fall in a subsequent year for which a higher deductible amount has been determined.

(4) For coinsurance days after December 31, 1981, the coinsurance amount is based on the deductible applicable for the calendar year in which the services were furnished.

(b) Specific coinsurance amounts. The specific SNF coinsurance amounts for each calendar year are published in the Federal Register no later than October 1 of the preceding year.

(c) Exception to published amounts. If the actual charge to the patient is less than the coinsurance amount applicable for the calendar year in which the services were furnished, the actual charge per day is the daily coinsurance.


§ 409.87 Blood deductible.

(a) General provisions. (1) As used in this section, packed red cells means the red blood cells that remain after plasma is separated from whole blood.

(2) A unit of packed red cells is treated as the equivalent of a unit of whole blood.

(3) Medicare does not pay for the first 3 units of whole blood or units of packed red cells that a beneficiary receives, during a calendar year, as an inpatient of a hospital or CAH or SNF, or on an outpatient basis under Medicare Part B.

(4) The deductible does not apply to other blood components such as platelets, fibrinogen, plasma, gamma globulin, and serum albumin, or to the cost of processing, storing, and administering blood.

(5) The blood deductible is in addition to the inpatient hospital deductible and daily coinsurance.

(6) The Part A blood deductible is reduced to the extent that the Part B blood deductible has been applied. For example, if a beneficiary had received one unit under Medicare Part B, and later in the same benefit period received three units under Medicare Part A, Medicare Part A would pay for the third of the latter units. (As specified in §401.161 of this chapter, the Part B blood deductible is reduced to the extent a blood deductible has been applied under Medicare Part A.)

(b) Beneficiary’s responsibility for the first 3 units of whole blood or packed red cells. (1) Basic rule. Except as specified in paragraph (b)(2) of this section, the beneficiary is responsible for the first 3 units of whole blood or packed red cells. He or she has the option of paying the hospital’s or CAH’s charges for the blood or packed red cells or arranging for it to be replaced.

(2) Exception. The beneficiary is not responsible for the first 3 units of whole blood or packed red cells if the provider obtained that blood or red cells at no charge other than a processing or service charge. In that case, the blood or red cells is deemed to have been replaced.

(c) Provider’s right to charge for the first 3 units of whole blood or packed red cells—(1) Basic rule. Except as specified in paragraph (c)(2) of this section, a provider may charge a beneficiary its customary charge for any of the first 3 units of whole blood or packed red cells.
(2) Exception. A provider may not charge the beneficiary for the first 3 units of whole blood or packed red cells in any of the following circumstances:
   (i) The blood or packed red cells has been replaced.
   (ii) The provider (or its blood supplier) receives, from an individual or a blood bank, a replacement offer that meets the criteria specified in paragraph (d) of this section. The provider is precluded from charging even if it or its blood supplier rejects the replacement offer.
   (iii) The provider obtained the blood or packed red cells at no charge other than a processing or service charge and it is therefore deemed to have been replaced.

(d) Criteria for replacement of blood. A blood replacement offer made by a beneficiary, or an individual or a blood bank on behalf of a beneficiary, discharges the beneficiary’s obligation to pay for deductible blood or packed red cells if the replacement blood meets the applicable criteria specified in Food and Drug Administration regulations under 21 CFR part 640, i.e.—
   (1) The replacement blood would not endanger the health of a recipient; and
   (2) The prospective donor’s health would not be endangered by making a blood donation.

Subpart H—Payment of Hospital Insurance Benefits

§ 409.89 Exemption of kidney donors from deductible and coinsurance requirements.

The deductible and coinsurance requirements set forth in this subpart do not apply to any services furnished to an individual in connection with the donation of a kidney for transplant surgery.

§ 409.100 To whom payment is made.

(a) Basic rule. Except as provided in paragraph (b) of this section—

(1) Medicare pays hospital insurance benefits only to a participating provider.

(2) For home health services (including medical supplies described in section 1861(m)(5) of the Act, but excluding durable medical equipment to the extent provided for in such section) furnished to an individual who at the time the item or service is furnished is under a plan of care of an HHA, payment is made to the HHA (without regard to whether the item or service is furnished by the HHA directly, under arrangement with the HHA, or under any other contracting or consulting arrangement).

(b) Exceptions. Medicare may pay hospital insurance benefits as follows:

(1) For emergency services furnished by a nonparticipating hospital, to the hospital or to the beneficiary, under the conditions prescribed in subpart G of part 424 of this chapter.

(2) For services furnished by a Canadian or Mexican hospital, to the hospital or to the beneficiary, under the conditions prescribed in subpart H of part 424 of this chapter.

§ 409.102 Amounts of payment.

(a) The amounts Medicare pays for hospital insurance benefits are generally determined in accordance with part 412 or part 413 of this chapter.

(b) Except as provided in §§ 409.61(d) and 409.89, hospital insurance benefits are subject to the deductible and coinsurance requirements set forth in subpart G of this part.

PART 410—SUPPLEMENTARY MEDICAL INSURANCE (SMI) BENEFITS

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§ 410.172 Payment for partial hospitalization services in CMHCs: Conditions.

§ 410.175 Alien absent from the United States.

AUTHORITY: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh).

SOURCE: 51 FR 41339, Nov. 14, 1986, unless otherwise noted.


Subpart A—General Provisions

§ 410.2 Definitions.

As used in this part—
Community mental health center (CMHC) means an entity that—

(1) Provides outpatient services, including specialized outpatient services for children, the elderly, individuals who are chronically mentally ill, and residents of its mental health service area who have been discharged from inpatient treatment at a mental health facility;

(2) Provides 24-hour-a-day emergency care services;

(3) Provides day treatment or other partial hospitalization services, or psychosocial rehabilitation services;

(4) Provides screening for patients being considered for admission to State mental health facilities to determine the appropriateness of this admission; and

(5) Meets applicable licensing or certification requirements for CMHCs in the State in which it is located.

Encounter means a direct personal contact between a patient and a physician, or other person who is authorized by State licensure law and, if applicable, by hospital or CAH staff bylaws, to order or furnish hospital services for diagnosis or treatment of the patient.

Nominal charge provider means a provider that furnishes services free of charge or at a nominal charge, and is either a public provider or another provider that (1) demonstrates to HCFA’s satisfaction that a significant portion of its patients are low-income; and (2) requests that payment for its services be determined accordingly.

Outpatient means a person who has not been admitted as an inpatient but who is registered on the hospital or CAH records as an outpatient and receives services (rather than supplies alone) directly from the hospital or CAH.

Partial hospitalization services means a distinct and organized intensive ambulatory treatment program that offers less than 24-hour daily care and furnishes the services described in § 410.43.

Participating refers to a hospital, CAH, SNF, HHA, CORF, or hospice that has in effect an agreement to participate in Medicare; or a clinic, rehabilitation agency, or public health agency that has a provider agreement to participate in Medicare but only for purposes of providing outpatient physical therapy, occupational therapy, or speech pathology services; or a CMHC.
that has in effect a similar agreement but only for purposes of providing partial hospitalization services, and non-participating refers to a hospital, CAH, SNF, HHA, CORF, hospice, clinic, rehabilitation agency, public health agency, or CMHC that does not have in effect a provider agreement to participate in Medicare.


§ 410.3 Scope of benefits.

(a) Covered services. The SMI program helps pay for the following:

(1) Medical and other health services such as physicians' services, outpatient services furnished by a hospital or a CAH, diagnostic tests, outpatient physical therapy and speech pathology services, rural health clinic services, Federally qualified health center services, and outpatient renal dialysis services.

(2) Services furnished by ambulatory surgical centers (ASCs), home health agencies (HHAs), comprehensive outpatient rehabilitation facilities (CORFs), and partial hospitalization services provided by community mental health centers (CMHCs).

(3) Other medical services, equipment, and supplies that are not covered under Medicare Part A hospital insurance.

(b) Limitations on amount of payment. (1) Medicare Part B does not pay the full reasonable costs or charges for all covered services. The beneficiary is responsible for an annual deductible and a blood deductible and, after the annual deductible has been satisfied, for coinsurance amounts specified for most of the services.

(2) Specific rules on payment are set forth in subpart E of this part.


Subpart B—Medical and Other Health Services

§ 410.10 Medical and other health services: Included services.

Subject to the conditions and limitations specified in this subpart, “medical and other health services” includes the following services:

(a) Physicians’ services.

(b) Services and supplies furnished incident to a physician’s professional services, of kinds that are commonly furnished in physicians’ offices and are commonly either furnished without charge or included in the physicians’ bills.

(c) Services and supplies, including partial hospitalization services, that are incident to physician services and are furnished to outpatients by or under arrangements made by a hospital or a CAH.

(d) Diagnostic services furnished to outpatients by or under arrangements made by a hospital or a CAH if the services are services that the hospital or CAH ordinarily furnishes to its outpatients for diagnostic study.

(e) Diagnostic laboratory and X-ray tests (including diagnostic mammography that meets the conditions for coverage specified in §410.34(b) of this subpart) and other diagnostic tests.

(f) X-ray therapy and other radiation therapy services.

(g) Medical supplies, appliances, and devices.

(h) Durable medical equipment.

(i) Ambulance services.

(j) Rural health clinic services.

(k) Home dialysis supplies and equipment; on or after July 1, 1991, epoetin (EPO) for home dialysis patients, and, on or after January 1, 1994, for dialysis patients, competent to use the drug; self-care home dialysis support services; and institutional dialysis services and supplies.

§ 410.5 Other applicable rules.

The following other rules of this chapter set forth additional policies and procedures applicable to four of the kinds of services covered under the SMI program:

(a) Part 405, subpart U: End-Stage Renal Disease services.

(b) Part 405, Subpart X: Rural Health Clinic and Federally Qualified Health Center services.

(c) Part 416: Ambulatory Surgical Center services.

(d) Part 493: Laboratory Services.

§ 410.20 Physicians’ services.

(a) Included services. Medicare Part B pays for physicians’ services, including diagnosis, therapy, surgery, consultation, and home, office, and institutional calls.

(b) By whom services must be furnished. Medicare Part B pays for the services specified in paragraph (a) of this section if they are furnished by one of the following professionals who is legally authorized to practice by the State in which he or she performs the functions or actions, and who is acting within the scope of his or her license:

(1) A doctor of medicine or osteopathy, including an osteopathic practitioner recognized in section 1101(a)(7) of the Act.

(2) By whom the services must be furnished. The services must be furnished by a facility or other entity as specified in §§410.3 and 410.8.

(3) Physician certification and recertification requirements. If the services are subject to physician certification requirements, they must be certified as being medically necessary, and as meeting other applicable requirements, in accordance with subpart B of part 424 of this chapter.

(b) Limitations on payment. Payment for medical and other health services is subject to limitations on the amounts of payment as specified in §§410.126 and 410.155 and to the annual and blood deductibles as set forth in §§410.160 and 410.161.

§ 410.22 Limitations on services of a chiropractor.

(a) Qualifications for chiropractors. (1) A chiropractor licensed or authorized to practice before July 1, 1974, and an individual who began studies in a chiropractic college before that date, must have—
   (i) Had preliminary education equal to the requirements for graduation from an accredited high school or other secondary school;
   (ii) Graduated from a college of chiropractic approved by the State's chiropractic examiners after completing a course of study covering a period of not less than 3 school years of 6 months each year in actual continuous attendance and covering adequate courses of study in the subjects of anatomy, physiology, symptomatology and diagnosis, hygiene and sanitation, chemistry, histology, pathology, and principles and practice of chiropractic, including clinical instruction in vertebral palpation, nerve tracing and adjusting; and
   (iii) Passed an examination prescribed by the State's chiropractic examiners covering the subjects specified in paragraph (a)(1)(ii) of this section.

(b) Limitations on services. (1) Medicare Part B pays only for a chiropractor's manual manipulation of the spine to correct a subluxation if the subluxation has resulted in a neuro-musculoskeletal condition for which manual manipulation is appropriate treatment.

Medicare Part B does not pay for X-rays or other diagnostic or therapeutic services furnished or ordered by a chiropractor.

[51 FR 41339, Nov. 14, 1986, as amended at 64 FR 59439, Nov. 2, 1999]

§ 410.23 Limitations on services of an optometrist.

Medicare Part B pays for the services of a doctor of optometry, which he or she is legally authorized to perform in the State in which he or she performs them, if the services are among those described in section 1861(s) of the Act and §410.10 of this part.

[64 FR 59439, Nov. 2, 1999]

§ 410.24 Limitations on services of a doctor of dental surgery or dental medicine.

Medicare Part B pays for services furnished by a doctor of dental surgery or dental medicine within the scope of his or her license, if the services would be covered as physicians' services when performed by a doctor of medicine or osteopathy.


1 For services furnished before July 1, 1981, Medicare Part B paid only for the following services of a doctor of dental surgery or dental medicine:
   Surgery on the jaw or any adjoining structure; and
   Reduction of a fracture of the jaw or other facial bone.
§ 410.25 Limitations on services of a podiatrist.

Medicare Part B pays for the services of a doctor of podiatric medicine, acting within the scope of his or her license, if the services would be covered as physicians' services when performed by a doctor of medicine or osteopathy.

§ 410.26 Services and supplies incident to a physician's professional services: Conditions.

(a) Medicare Part B pays for services and supplies incident to a physician's professional services, including drugs and biologicals that cannot be self-administered, if the services or supplies are of the type that are commonly furnished in a physician's office or clinic, and are commonly furnished either without charge, or included in the physician's bill.

(b) Drugs and biologicals are also subject to the limitations specified in §410.29.

§ 410.27 Outpatient hospital services and supplies incident to a physician service: Conditions.

(a) Medicare Part B pays for hospital services and supplies furnished incident to a physician service to outpatients, including drugs and biologicals required in the performance of the services (even if those drugs or biologicals are self-administered), if—

(1) They are furnished—

(i) By or under arrangements made by a participating hospital, except in the case of an SNF resident as provided in §411.15(p) of this chapter;

(ii) As an integral though incidental part of a physician's services; and

(iii) In the hospital or at a location (other than an RHC or an FQHC) that HCFA designates as a department of a provider under §413.65 of this chapter; and

(2) In the case of partial hospitalization services, also meet the conditions of paragraph (d) of this section.

(b) Drugs and biologicals are also subject to the limitations specified in §410.168.

(c) Rules on emergency services furnished to outpatients by nonparticipating hospitals are specified in §410.168.

(d) Medicare Part B pays for partial hospitalization services if they are—

(1) Prescribed by a physician who certifies and recertifies the need for the services in accordance with subpart B of part 424 of this chapter; and

(2) Furnished under a plan of treatment as required under subpart B of part 424 of this chapter.

(e) Services furnished by an entity other than the hospital are subject to the limitations specified in §410.42(a).

(f) Services furnished at a location (other than an RHC or an FQHC) that HCFA designates as a department of a provider under §413.65 of this chapter must be under the direct supervision of a physician. “Direct supervision” means the physician must be present and on the premises of the location and immediately available to furnish assistance and direction throughout the performance of the procedure. It does not mean that the physician must be present in the room when the procedure is performed.


§ 410.28 Hospital or CAH diagnostic services furnished to outpatients: Conditions.

(a) Medicare Part B pays for hospital or CAH diagnostic services furnished to outpatients, including drugs and biologicals required in the performance of the services (even if those drugs or biologicals are self-administered), if those services meet the following conditions:

(1) They are furnished by or under arrangements made by a participating hospital or participating CAH, except in the case of an SNF resident as provided in §411.15(p) of this chapter.

(2) They are ordinarily furnished by, or under arrangements made by, the hospital or CAH to its outpatients for the purpose of diagnostic study.

(3) They would be covered as inpatient hospital services if furnished to an inpatient.

(b) Drugs and biologicals are also subject to the limitations specified in §410.29(b) and (c).

(c) Diagnostic services furnished by an entity other than the hospital or CAH are subject to the limitations specified in §410.42(a).
§ 410.29 Limitations on drugs and biologicals.

Medicare part B does not pay for the following:

(a) Except as provided in § 410.28(a) for outpatient diagnostic services and § 410.63(b) for blood clotting factors, and except for EPO, any drug or biological that can be self-administered.

(b) Any drug product that meets all of the following conditions:

(1) The approved labeling includes the indication for preventing or treating the rejection of a transplanted organ or tissue.

(2) The approved labeling includes the indication for use in conjunction with immunosuppressive drugs to prevent or treat rejection of a transplanted organ or tissue.

(3) Have been determined by a carrier (in accordance with part 421, subpart C of this chapter), in processing a Medicare claim, to be reasonable and necessary for the specific purpose of preventing or treating the rejection of a patient’s transplanted organ or tissue, or for use in conjunction with immunosuppressive drugs for the purpose of preventing or treating the rejection of a patient’s transplanted organ or tissue. (In making these determinations, the carriers may consider factors such as authoritative drug compendia, current medical literature, recognized standards of medical practice, and professional medical publications.)

§ 410.30 Prescription drugs used in immunosuppressive therapy.

(a) Scope. Payment may be made for prescription drugs used in immunosuppressive therapy that have been approved for marketing by the FDA and that meet one of the following conditions:

(1) The approved labeling includes the indication for preventing or treating the rejection of a transplanted organ or tissue.

(2) The approved labeling includes the indication for use in conjunction with immunosuppressive drugs to prevent or treat rejection of a transplanted organ or tissue.

(3) Have been determined by a carrier (in accordance with part 421, subpart C of this chapter), in processing a Medicare claim, to be reasonable and necessary for the specific purpose of preventing or treating the rejection of a patient’s transplanted organ or tissue, or for use in conjunction with immunosuppressive drugs for the purpose of preventing or treating the rejection of a patient’s transplanted organ or tissue. (In making these determinations, the carriers may consider factors such as authoritative drug compendia, current medical literature, recognized standards of medical practice, and professional medical publications.)

(b) Period of eligibility. Coverage is available only for prescription drugs used in immunosuppressive therapy, furnished to an individual who receives an organ or tissue transplant for which Medicare payment is made, for the following periods:

(1) For drugs furnished before 1995, for a period of up to 1 year beginning with the date of discharge from the hospital during which the covered transplant was performed.

(2) For drugs furnished during 1995, within 18 months after the date of discharge from the hospital during which the covered transplant was performed.
(3) For drugs furnished during 1996, within 24 months after the date of discharge from the hospital during which the covered transplant was performed.

(4) For drugs furnished during 1997, within 30 months after the date of discharge from the hospital during which the covered transplant was performed.

(5) For drugs furnished after 1997, within 36 months after the date of discharge from the hospital during which the covered transplant was performed.

(c) Coverage. Drugs are covered under this provision irrespective of whether they can be self-administered.

§ 410.31 Bone mass measurement: Conditions for coverage and frequency standards.

(a) Definition. As used in this section unless specified otherwise, the following definition applies:

Bone mass measurement means a radiologic, radioisotopic, or other procedure that meets the following conditions:

(1) Is performed for the purpose of identifying bone mass, detecting bone loss, or determining bone quality.

(2) Is performed with either a bone densitometer (other than dual-photon absorptiometry) or with a bone sonometer system that has been cleared for marketing for this use by the FDA under 21 CFR part 807, or approved for marketing by the FDA for this use under 21 CFR part 814.

(3) Includes a physician's interpretation of the results of the procedure.

(b) Conditions for coverage. Medicare covers a medically necessary bone mass measurement if the following conditions are met:

(1) Following an evaluation of the beneficiary's need for the measurement, including a determination as to the medically appropriate procedure to be used for the beneficiary, it is ordered by the physician or a qualified nonphysician practitioner (as these terms are defined in §410.32(a)) treating the beneficiary.

(2) It is performed under the appropriate level of supervision of a physician (as set forth in §410.32(b)).

(3) It is reasonable and necessary for diagnosing, treating, or monitoring the condition of a beneficiary who meets the conditions described in paragraph (d) of this section.

(c) Standards on frequency of coverage—(1) General rule. Except as allowed under paragraph (c)(2) of this section, Medicare may cover a bone mass measurement for a beneficiary if at least 23 months have passed since the month the last bone mass measurement was performed.

(2) Exception. If medically necessary, Medicare may cover a bone mass measurement for a beneficiary more frequently than allowed under paragraph (c)(1) of this section. Examples of situations where more frequent bone mass measurement procedures may be medically necessary include, but are not limited to, the following medical circumstances:

(i) Monitoring beneficiaries on long-term glucocorticoid (steroid) therapy of more than 3 months.

(ii) Allowing for a confirmatory baseline bone mass measurement (either central or peripheral) to permit monitoring of beneficiaries in the future if the initial test was performed with a technique that is different from the proposed monitoring method.

(d) Beneficiaries who may be covered. The following categories of beneficiaries may receive Medicare coverage for a medically necessary bone mass measurement:

(1) A woman who has been determined by the physician (or a qualified nonphysician practitioner) treating her to be estrogen-deficient and at clinical risk for osteoporosis, based on her medical history and other findings.

(2) An individual with vertebral abnormalities as demonstrated by an x-ray to be indicative of osteoporosis, osteopenia, or vertebral fracture.

(3) An individual receiving (or expecting to receive) glucocorticoid (steroid) therapy equivalent to 7.5 mg of prednisone, or greater, per day for more than 3 months.

(4) An individual with primary hyperparathyroidism.

(5) An individual being monitored to assess the response to or efficacy of an FDA-approved osteoporosis drug therapy.

(e) Denial as not reasonable and necessary. If HCFA determines that a bone
§ 410.32 Diagnostic x-ray tests, diagnostic laboratory tests, and other diagnostic tests: Conditions.

(a) Ordering diagnostic tests. All diagnostic x-ray tests, diagnostic laboratory tests, and other diagnostic tests must be ordered by the physician who is treating the beneficiary, that is, the physician who furnishes a consultation or treats a beneficiary for a specific medical problem and who uses the results in the management of the beneficiary's specific medical problem. Tests not ordered by the physician who is treating the beneficiary are not reasonable and necessary (see § 411.15(k)(1) of this chapter).

(1) Chiropractic exception. A physician may order an x-ray to be used by a chiropractor to demonstrate the subluxation of the spine that is the basis for a beneficiary to receive manual manipulation treatments even though the physician does not treat the beneficiary.

(2) Mammography exception. A physician who meets the qualification requirements for an interpreting physician under section 354 of the Public Health Service Act as provided in § 410.34(a)(7) may order a diagnostic mammogram based on the findings of a screening mammogram even though the physician does not treat the beneficiary.

(3) Application to nonphysician practitioners. Nonphysician practitioners (that is, clinical nurse specialists, clinical psychologists, clinical social workers, nurse practitioners, and physician assistants) who furnish services that would be physician services if furnished by a physician, and who are operating within the scope of their authority under State law and within the scope of their Medicare statutory benefit, may be treated the same as physicians treating beneficiaries for the purpose of this paragraph.

(b) Diagnostic x-ray and other diagnostic tests—(1) Basic rule. Except as indicated in paragraphs (b)(2) of this section, all diagnostic x-ray and other diagnostic tests covered under section 1861(s)(3) of the Act and payable under the physician fee schedule must be furnished under the appropriate level of supervision by a physician as defined in section 1861(r) of the Act. Services furnished without the required level of supervision are not reasonable and necessary (see § 411.15(k)(1) of this chapter).

(2) Exceptions. The following diagnostic tests payable under the physician fee schedule are excluded from the basic rule set forth in paragraph (b)(1) of this section:
   (i) Diagnostic mammography procedures, which are regulated by the Food and Drug Administration.
   (ii) Diagnostic tests personally furnished by a qualified audiologist as defined in section 1861(ll)(3) of the Act.
   (iii) Diagnostic psychological testing services personally furnished by a clinical psychologist or a qualified independent psychologist as defined in program instructions.
   (iv) Diagnostic tests (as established through program instructions) personally performed by a physical therapist who is certified by the American Board of Physical Therapy Specialties as a qualified electrophysiologic clinical specialist and permitted to provide the service under State law.
   (v) Diagnostic tests performed by a nurse practitioner or clinical nurse specialist authorized to perform the tests under applicable State laws.

(3) Levels of supervision. Except where otherwise indicated, all diagnostic x-ray and other diagnostic tests subject to this provision and payable under the physician fee schedule must be furnished under at least a general level of physician supervision as defined in paragraph (b)(3)(i) of this section.

[63 FR 34327, June 24, 1998]
addition, some of these tests also require either direct or personal supervision as defined in paragraphs (b)(3)(ii) or (b)(3)(iii) of this section, respectively. (However, diagnostic tests performed by a physician assistant (PA) that the PA is legally authorized to perform under State law require only a general level of physician supervision.) When direct or personal supervision is required, physician supervision at the specified level is required throughout the performance of the test.

(i) General supervision means the procedure is furnished under the physician's overall direction and control, but the physician's presence is not required during the performance of the procedure. Under general supervision, the training of the nonphysician personnel who actually perform the diagnostic procedure and the maintenance of the necessary equipment and supplies are the continuing responsibility of the physician.

(ii) Direct supervision in the office setting means the physician must be present in the office suite and immediately available to furnish assistance and direction throughout the performance of the procedure. It does not mean that the physician must be present in the room when the procedure is performed.

(iii) Personal supervision means a physician must be in attendance in the room during the performance of the procedure.

(c) Portable x-ray services. Portable x-ray services furnished in a place of residence used as the patient's home are covered if the following conditions are met:

(1) These services are furnished under the general supervision of a physician, as defined in paragraph (b)(3)(i) of this section.

(2) The supplier of these services meets the requirements set forth in part 486, subpart C of this chapter, concerning conditions for coverage for portable x-ray services.

(3) The procedures are limited to—

(i) Skeletal films involving the extremities, pelvis, vertebral column, or skull;

(ii) Chest or abdominal films that do not involve the use of contrast media; and

(iii) Diagnostic mammograms if the approved portable x-ray supplier, as defined in subpart C of part 486 of this chapter, meets the certification requirements of section 354 of the Public Health Service Act, as implemented by 21 CFR part 900, subpart B.

(d) Diagnostic laboratory tests. Medicare Part B pays for covered diagnostic laboratory tests that are furnished by any of the following:

(1) A participating hospital or participating RPCH.

(2) A nonparticipating hospital that meets the requirements for emergency outpatient services specified in subpart G of part 424 of this chapter and the laboratory requirements specified in part 493 of this chapter.

(3) The office of the patient's attending or consulting physician if that physician is a doctor of medicine, osteopathy, pediatric medicine, dental surgery, or dental medicine.

(4) An RHC.

(5) A laboratory, if it meets the applicable requirements for laboratories of part 493 of this chapter, including the laboratory of a nonparticipating hospital that does not meet the requirements for emergency outpatient services in subpart G of part 424 of this chapter.

(6) An FQHC.

(7) An SNF to its resident under §411.15(p) of this chapter, either directly (in accordance with §483.75(k)(1)(i) of this chapter) or under an arrangement (as defined in §409.3 of this chapter) with another entity described in this paragraph.


§410.33 Independent diagnostic testing facility.

(a) General rule. (1) Effective for diagnostic procedures performed on or after March 15, 1999, carriers will pay for diagnostic procedures under the physician fee schedule only when performed by a physician, a group practice of physicians, an approved supplier of portable x-ray services, a nurse practitioner, or a clinical nurse specialist when he or she performs a test he or
§ 410.33 Supervising physicians and technicians

† she is authorized by the State to perform, or an independent diagnostic testing facility (IDTF). An IDTF may be a fixed location, a mobile entity, or an individual nonphysician practitioner. It is independent of a physician’s office or hospital; however, these rules apply when an IDTF furnishes diagnostic procedures in a physician’s office.

(2) Exceptions. The following diagnostic tests that are payable under the physician fee schedule and furnished by a nonhospital testing entity are not required to be furnished in accordance with the criteria set forth in paragraphs (b) through (e) of this section:

(i) Diagnostic mammography procedures, which are regulated by the Food and Drug Administration.

(ii) Diagnostic tests personally furnished by a qualified audiologist as defined in section 1861(ll)(3) of the Act.

(iii) Diagnostic psychological testing services personally furnished by a clinical psychologist or a qualified independent psychologist as defined in program instructions.

(iv) Diagnostic tests (as established through program instructions) personally performed by a physical therapist who is certified by the American Board of Physical Therapy Specialties as a qualified electrophysiologic clinical specialist and permitted to provide the service under State law.

(b) Supervising physician. (1) An IDTF must have one or more supervising physicians who are responsible for the direct and ongoing oversight of the quality of the testing performed, the proper operation and calibration of the equipment used to perform tests, and the qualification of nonphysician personnel who use the equipment. This level of supervision is that required for general supervision set forth in § 410.32(b)(3)(i).

(2) The supervising physician must evidence proficiency in the performance and interpretation of each type of diagnostic procedure performed by the IDTF. The proficiency may be documented by certification in specific medical specialties or subspecialties or by criteria established by the carrier for the service area in which the IDTF is located. In the case of a procedure requiring the direct or personal supervision of a physician as set forth in § 410.32(b)(3)(ii) or (b)(3)(iii), the IDTF’s supervising physician must personally furnish this level of supervision whether the procedure is performed in the IDTF or, in the case of mobile services, at the remote location. The IDTF must maintain documentation of sufficient physician resources during all hours of operations to assure that the required physician supervision is furnished. In the case of procedures requiring direct supervision, the supervising physician may oversee concurrent procedures.

(c) Nonphysician personnel. Any nonphysician personnel used by the IDTF to perform tests must demonstrate the basic qualifications to perform the tests in question and have training and proficiency as evidenced by licensure or certification by the appropriate State health or education department. In the absence of a State licensing board, the technician must be certified by an appropriate national credentialing body. The IDTF must maintain documentation available for review that these requirements are met.

(d) Ordering of tests. All procedures performed by the IDTF must be specifically ordered in writing by the physician who is treating the beneficiary, that is, the physician who is furnishing a consultation or treating a beneficiary for a specific medical problem and who uses the results in the management of the beneficiary’s specific medical problem. (Nonphysician practitioners may order tests as set forth in § 410.32(a)(3).) The order must specify the diagnosis or other basis for the testing. The supervising physician for the IDTF may not order tests to be performed by the IDTF, unless the IDTF’s supervising physician is in fact the beneficiary’s treating physician. That is, the physician in question had a relationship with the beneficiary prior to the performance of the testing and is treating the beneficiary for a specific medical problem. The IDTF may not add any procedures based on internal protocols without a written order from the treating physician.

(e) Multi-State entities. An IDTF that operates across State boundaries must maintain documentation that its supervising physicians and technicians
are licensed and certified in each of the States in which it is furnishing services.

(f) Applicability of State law. An IDTF must comply with the applicable laws of any State in which it operates.

§ 410.34 Mammography services: Conditions for and limitations on coverage.

(a) Definitions. As used in this section, the following definitions apply:

(1) Diagnostic mammography means a radiologic procedure furnished to a man or woman with signs or symptoms of breast disease, or a personal history of breast cancer, or a personal history of biopsy-proven benign breast disease, and includes a physician’s interpretation of the results of the procedure.

(2) Screening mammography means a radiologic procedure furnished to a woman without signs or symptoms of breast disease, for the purpose of early detection of breast cancer, and includes a physician’s interpretation of the results of the procedure.

(3) Supplier of diagnostic mammography means a facility that is certified and responsible for ensuring that all diagnostic mammography services furnished to Medicare beneficiaries meet the conditions for coverage of diagnostic mammography services as specified in paragraph (b) of this section.

(4) Supplier of screening mammography means a facility that is certified and responsible for ensuring that all screening mammography services furnished to Medicare beneficiaries meet the conditions and limitations for coverage of screening mammography services as specified in paragraphs (c) and (d) of this section.

(5) Certificate means the certificate described in 21 CFR 900.2(b) that may be issued to, or renewed for, a facility that meets the requirements for conducting an examination or procedure involving mammography.

(6) Provisional certificate means the provisional certificate described in 21 CFR 900.2(m) that may be issued to a facility to enable the facility to qualify to meet the requirements for conducting an examination or procedure involving mammography.

(7) The term meets the certification requirements of section 354 of the Public Health Service (PHS) Act means that in order to qualify for coverage of its services under the Medicare program, a supplier of diagnostic or screening mammography services must meet the following requirements:

(i) Must have a valid provisional certificate, or a valid certificate, that has been issued by FDA indicating that the supplier meets the certification requirements of section 354 of the PHS Act, as implemented by 21 CFR part 900, subpart B.

(ii) Has not been issued a written notification by FDA that states that the supplier must cease conducting mammography examinations because the supplier is not in compliance with certain critical certification requirements of section 354 of the PHS Act, implemented by 21 CFR part 900, subpart B.

(iii) Must not employ for provision of the professional component of mammography services a physician or physicians for whom the facility has received written notification by FDA that the physician (or physicians) is (or are) in violation of the certification requirements set forth in section 354 of the PHS Act, as implemented by 21 CFR 900.12(a)(1)(i).

(b) Conditions for coverage of diagnostic mammography services. Medicare Part B pays for diagnostic mammography services if they meet the following conditions:

(1) They are ordered by a doctor of medicine or osteopathy (as defined in section 1861(r)(1) of the Act).

(2) They are furnished by a supplier of diagnostic mammography services that meets the certification requirements of section 354 of the PHS Act, as implemented by 21 CFR part 900, subpart B.

(c) Conditions for coverage of screening mammography services. Medicare Part B pays for screening mammography services if they are furnished by a supplier of screening mammography services that meets the certification requirements of section 354 of the PHS Act, as implemented by 21 CFR part 900, subpart B.

(d) Limitations on coverage of screening mammography services. The following limitations apply to coverage of
screening mammography services as described in paragraphs (c) and (d) of this section:

(1) The service must be, at a minimum, a two-view exposure (that is, a cranio-caudal and a medial lateral oblique view) of each breast.

(2) Payment may not be made for screening mammography performed on a woman under age 35.

(3) Payment may be made for only 1 screening mammography performed on a woman over age 34, but under age 40.

(4) For an asymptomatic woman over 39 years of age, payment may be made for a screening mammography performed after at least 11 months have passed following the month in which the last screening mammography was performed.


§ 410.35 X-ray therapy and other radiation therapy services. Scope.

Medicare Part B pays for X-ray therapy and other radiation therapy services, including radium therapy and radioactive isotope therapy, and materials and the services of technicians administering the treatment.


§ 410.36 Medical supplies, appliances, and devices. Scope.

(a) Medicare Part B pays for the following medical supplies, appliances and devices:

(1) Surgical dressings, and splints, casts, and other devices used for reduction of fractures and dislocations.

(2) Prosthetic devices, other than dental, that replace all or part of an internal body organ, including colostomy bags and supplies directly related to colostomy care, including—

   (i) Replacement of prosthetic devices; and

   (ii) One pair of conventional eyeglasses or conventional contact lenses furnished after each cataract surgery during which an intraocular lens is inserted.

(3) Leg, arm, back, and neck braces and artificial legs, arms, and eyes, including replacements if required because of a change in the individual’s physical condition.

(b) As a requirement for payment, HCFA may determine through carrier instructions, or carriers may determine, that an item listed in paragraph (a) of this section requires a written physician order before delivery of the item.


§ 410.37 Colorectal cancer screening tests: Conditions for and limitations on coverage.

(a) Definitions. As used in this section, the following definitions apply:

(1) Colorectal cancer screening tests means any of the following procedures furnished to an individual for the purpose of early detection of colorectal cancer:

(i) Screening fecal-occult blood tests.

(ii) Screening flexible sigmoidoscopies.

(iii) In the case of an individual at high risk for colorectal cancer, screening colonoscopies.

(iv) Screening barium enemas.

(v) Other tests or procedures, and modifications to tests under this paragraph, with such frequency and payment limits as HCFA determines appropriate, in consultation with appropriate organizations.

(2) Screening fecal-occult blood test means a guaiac-based test for peroxidase activity, testing two samples from each of three consecutive stools.
§410.38 Durable medical equipment: Scope and conditions.

(a) Medicare Part B pays for the rental or purchase of durable medical equipment, including iron lungs, oxygen tents, hospital beds, and wheelchairs, if the equipment is used in the patient’s home or in an institution that is used as a home.
§ 410.39  Prostate cancer screening tests: Conditions for and limitations on coverage.

(a) Definitions. As used in this section, the following definitions apply:

(1) Prostate cancer screening tests means any of the following procedures furnished to an individual for the purpose of early detection of prostate cancer:

(i) A screening digital rectal examination.

(ii) A screening prostate-specific antigen blood test.

(iii) For years beginning after 2002, other procedures HCFA finds appropriate for the purpose of early detection of prostate cancer, taking into account changes in technology and standards of medical practice, availability, effectiveness, costs, and other factors HCFA considers appropriate.

(2) Screening digital rectal examination means a clinical examination of an individual’s prostate for nodules or other abnormalities of the prostate.

(3) A screening prostate-specific antigen blood test means a test that measures the level of prostate-specific antigen in an individual’s blood.
(4) A physician for purposes of this provision means a doctor of medicine or osteopathy (as defined in section 1861(r)(1) of the Act) who is fully knowledgeable about the beneficiary, and who would be responsible for explaining the results of the screening examination or test.

(5) A physician assistant, nurse practitioner, clinical nurse specialist, or certified nurse midwife for purposes of this provision means a physician assistant, nurse practitioner, clinical nurse specialist, or certified nurse midwife (as defined in sections 1861(aa) and 1861(gg) of the Act) who is fully knowledgeable about the beneficiary, and who would be responsible for explaining the results of the screening examination or test.

(b) Condition for coverage of screening digital rectal examinations. Medicare Part B pays for a screening digital rectal examination if it is performed by the beneficiary’s physician, or by the beneficiary’s physician assistant, nurse practitioner, clinical nurse specialist, or certified nurse midwife as defined in paragraphs (a)(4) or (a)(5) of this section who is authorized to perform this service under State law.

(c) Limitation on coverage of screening digital rectal examinations. (1) Payment may not be made for a screening digital rectal examination performed for a man age 50 or younger.

(2) For an individual over 50 years of age, payment may be made for a screening digital rectal examination only if the man has not had such an examination paid for by Medicare during the preceding 11 months following the month in which his last Medicare-covered screening digital rectal examination was performed.

(d) Condition for coverage of screening prostate-specific antigen blood tests. Medicare Part B pays for a screening prostate-specific antigen blood test performed for a man age 50 or younger.

(e) Limitation on coverage of screening prostate-specific antigen blood test. (1) Payment may not be made for a screening prostate-specific antigen blood test performed for a man age 50 or younger.

(2) For an individual over 50 years of age, payment may be made for a screening prostate-specific antigen blood test only if the man has not had such an examination paid for by Medicare during the preceding 11 months following the month in which his last Medicare-covered screening prostate-specific antigen blood test was performed.

§ 410.40 Coverage of ambulance services.

(a) Basic rules. Medicare Part B covers ambulance services if the following conditions are met:

(1) The supplier meets the applicable vehicle, staff, and billing and reporting requirements of § 410.41 and the service meets the medical necessity and origin and destination requirements of paragraphs (d) and (e) of this section.

(2) Medicare Part A payment is not made directly or indirectly for the services.

(b) Levels of services. Medicare covers ambulance services within the United States at the following levels of services:

(1) Basic life support (BLS) services.

(2) Advanced life support (ALS) services.

(3) Paramedic ALS intercept services described in paragraph (c) of this section.

(c) Paramedic ALS intercept services. Paramedic ALS intercept services must meet the following requirements:

(1) Be furnished in an area that is designated as a rural area by any law or regulation of the State or that is located in a rural census tract of a metropolitan statistical area (as determined under the most recent Goldsmith Modification). (The Goldsmith Modification is a methodology to identify small towns and rural areas within large metropolitan counties that are isolated from central areas by distance or other features.)

(2) Be furnished under contract with one or more volunteer ambulance services that meet the following conditions:
§ 410.41 Requirements for ambulance suppliers.

(a) Vehicle. A vehicle used as an ambulance must meet the following requirements:

(1) Be specially designed to respond to medical emergencies or provide acute medical care to transport the sick and injured and comply with all State and local laws governing an emergency transportation vehicle.

(2) Be equipped with emergency warning lights and sirens, as required by State or local laws.

(3) Be equipped with telecommunications equipment as required by State or local law to include, at a minimum, one two-way voice radio or wireless telephone.

(i) Are certified to furnish ambulance services as required under § 410.41.

(ii) Furnish services only at the BLS level.

(iii) Be prohibited by State law from billing for any service.

(3) Be furnished by a paramedic ALS intercept supplier that meets the following conditions:

(i) Is certified to furnish ALS services as required in § 410.41(b)(2).

(ii) Bills all the recipients who receive ALS intercept services from the entity, regardless of whether or not those recipients are Medicare beneficiaries.

(d) Medical necessity requirements—(1) General rule. Medicare covers ambulance services only if they are furnished to a beneficiary whose medical condition is such that other means of transportation would be contraindicated. For nonemergency ambulance transportation, the following criteria must be met to ensure that ambulance transportation is medically necessary:

(i) The beneficiary is unable to get up from bed without assistance.

(ii) The beneficiary is unable to ambulate.

(iii) The beneficiary is unable to sit in a chair or wheelchair.

(2) Special rule for nonemergency, scheduled ambulance services. Medicare covers nonemergency, scheduled ambulance services if the ambulance supplier, before furnishing the service to the beneficiary, obtains a written order from the beneficiary's attending physician certifying that the medical necessity requirements of paragraph (d)(1) of this section are met. The physician's order must be dated no earlier than 60 days before the date the service is furnished.

(3) Special rule for nonemergency, unscheduled ambulance services. Medicare covers nonemergency, unscheduled ambulance services under the following circumstances:

(i) For a resident of a facility who is under the care of a physician if the ambulance supplier obtains a written order from the beneficiary’s attending physician, within 48 hours after the transport, certifying that the medical necessity requirements of paragraph (d)(1) of this section are met.

(iii) For a beneficiary residing at home or in a facility who is not under the direct care of a physician. A physician certification is not required.

(e) Origin and destination requirements. Medicare covers the following ambulance transportation:

(1) From any point of origin to the nearest hospital, CAH, or SNF that is capable of furnishing the required level and type of care for the beneficiary's illness or injury. The hospital or CAH must have available the type of physician or physician specialist needed to treat the beneficiary's condition.

(2) From a hospital, CAH, or SNF to the beneficiary's home.

(3) From a SNF to the nearest supplier of medically necessary services not available at the SNF where the beneficiary is a resident, including the return trip.

(4) For a beneficiary who is receiving renal dialysis for treatment of ESRD, from the beneficiary's home to the nearest facility that furnishes renal dialysis, including the return trip.

(f) Specific limits on coverage of ambulance services outside the United States. If services are furnished outside the United States, Medicare Part B covers ambulance transportation to a foreign hospital only in conjunction with the beneficiary's admission for medically necessary inpatient services as specified in subpart H of part 424 of this chapter.

[64 FR 3648, Jan. 25, 1999, as amended at 65 FR 13914, Mar. 15, 2000]
§ 410.34-lined medical equipment as required by State or local laws.

(b) Vehicle staff—(1) BLS vehicles. A vehicle furnishing ambulance services must be staffed by at least two people, one of whom must meet the following requirements:
   (i) Be certified as an emergency medical technician by the State or local authority where the services are furnished.
   (ii) Be legally authorized to operate all lifesaving and life-sustaining equipment on board the vehicle.

(2) ALS vehicles. In addition to meeting the vehicle staff requirements of paragraph (b)(1) of this section, one of the two staff members must be certified as a paramedic or an emergency medical technician, by the State or local authority where the services are being furnished, to perform one or more ALS services.

(c) Billing and reporting requirements. An ambulance supplier must comply with the following requirements:

(1) Bill for ambulance services using HCFA-designated procedure codes to describe origin and destination and indicate on claims form that the physician certification is on file.

(2) Upon a carrier’s request, complete and return the ambulance supplier form designated by HCFA and provide the Medicare carrier with documentation of compliance with emergency vehicle and staff licensure and certification requirements in accordance with State and local laws.

(3) Upon a carrier’s request, provide additional information and documentation as required.

§ 410.43 Partial hospitalization services: Conditions and exclusions.

(a) Partial hospitalization services are services that—

(1) Are reasonable and necessary for the diagnosis or active treatment of the individual’s condition;

(2) Are reasonably expected to improve or maintain the individual’s condition and functional level and to prevent relapse or hospitalization; and

(3) Include any of the following:
   (i) Individual and group therapy with physicians or psychologists or other mental health professionals to the extent authorized under State law.
   (ii) Occupational therapy requiring the skills of a qualified occupational therapist.
   (iii) Services of social workers, trained psychiatric nurses, and other staff trained to work with psychiatric patients.
   (iv) Drugs and biologicals furnished for therapeutic purposes, subject to the limitations specified in §410.29.
   (v) Individualized activity therapies that are not primarily recreational or diversionary.

§ 410.42 Limitations on coverage of certain services furnished to hospital outpatients.

(a) General rule. Except as provided in paragraph (b) of this section, Medicare Part B does not pay for any item or service that is furnished to a hospital outpatient (as defined in §410.2) during an encounter (as defined in §410.2) by an entity other than the hospital unless the hospital has an arrangement (as defined in §409.3 of this chapter) with that entity to furnish that particular service to its patients. As used in this paragraph, the term “hospital” includes a CAH.

(b) Exception. The limitations stated in paragraph (a) of this section do not apply to the following services:

(1) Physician services that meet the requirements of §415.102(a) of this chapter for payment on a fee schedule basis.

(2) Physician assistant services, as defined in section 1861(s)(2)(K)(i) of the Act.

(3) Nurse practitioner and clinical nurse specialist services, as defined in section 1861(s)(2)(K)(ii) of the Act.

(4) Certified nurse mid-wife services, as defined in section 1861(gg) of the Act.

(5) Qualified psychologist services, as defined in section 1861(ii) of the Act.

(6) Services of an anesthetist, as defined in §410.69.

(7) Services furnished to SNF residents as defined in §411.15(p) of this chapter.

[65 FR 18536, Apr. 7, 2000]
§ 410.45 Rural health clinic services: Scope and conditions.

(a) Medicare Part B pays for the following rural health clinic services, if they are furnished in accordance with the requirements and conditions specified in part 405, subpart X, and part 491 of this chapter:

(1) Physicians' services.
(2) Services and supplies furnished as an incident to physicians' professional services.
(3) Nurse practitioner and physician assistant services.
(4) Services and supplies furnished as an incident to nurse practitioners' or physician assistants' services.
(5) Visiting nurse services.
(b) Medicare pays for rural health clinic services when they are furnished at the clinic, at a hospital or other medical facility, or at the beneficiary's place of residence.

§ 410.50 Institutional dialysis services and supplies: Scope and conditions.

Medicare Part B pays for the following institutional dialysis services and supplies if they are furnished in approved ESRD facilities:

(a) All services, items, supplies, and equipment necessary to perform dialysis and drugs medically necessary in the treatment of the patient for ESRD.
(b) Routine dialysis monitoring tests (i.e., hematocrit and clotting time) used by the facility to monitor the patient's fluids incident to each dialysis treatment, when performed by qualified staff of the facility under the direction of a physician, as provided in § 405.2163(b) of this chapter, even if the facility does not meet the conditions for coverage of services of independent laboratories in subpart M of part 405 of this chapter.
(c) Routine diagnostic tests.
(d) Epoetin (EPO) and its administration.

§ 410.52 Home dialysis services, supplies, and equipment: Scope and conditions.

(a) Medicare Part B pays for the following services, supplies, and equipment furnished to an ESRD patient in his or her home:

(1) Purchase or rental, installation, and maintenance of all dialysis equipment necessary for home dialysis, and reconditioning of this equipment. Dialysis equipment includes, but is not limited to, artificial kidney and automated peritoneal dialysis machines, and support equipment such as blood pumps, bubble detectors, and other alarm systems.
(2) Items and supplies required for dialysis, including (but not limited to) dialyzers, syringes and needles, forceps, scissors, scales, sphygmomanometer with cuff and stethoscope, alcohol wipes, sterile drapes, and rubber gloves.
(3) Home dialysis support services furnished by an approved ESRD facility, including periodic monitoring of the patient's home adaptation, emergency visits by qualified provider or facility personnel, any of the tests specified in paragraphs (b) through (d) of § 410.50, personnel costs associated with the installation and maintenance of dialysis equipment, testing and appropriate treatment of water, and ordering of supplies on an ongoing basis.
§ 410.56 Screening pelvic examinations.

(a) Conditions for screening pelvic examinations. Medicare Part B pays for a screening pelvic examination (including a clinical breast examination) if it is performed by a doctor of medicine or osteopathy (as defined in section 1861(r)(1) of the Act), or by a certified nurse midwife (as defined in section 1861(gg) of the Act), or a physician assistant, nurse practitioner, or clinic nurse specialist (as defined in section 1861(aa) of the Act) who is authorized under State law to perform the examination.

(b) Limits on coverage of screening pelvic examinations. The following limitations apply to coverage of screening pelvic examination services:

(1) General rule. Except as specified in paragraphs (b)(2) and (b)(3) of this section, payment may be made for a pelvic examination performed on an asymptomatic woman only if the individual has not had a pelvic examination paid for by Medicare during the preceding 35 months following the month in which her last Medicare-covered screening pelvic examination was performed.

(2) More frequent screening based on high-risk factors. Subject to the limitation as specified in paragraph (b)(4) of this section, payment may be made for a screening pelvic examination performed more frequently than once every 36 months if the test is performed by a physician or other practitioner specified in paragraph (a) of this section, and there is evidence that the woman is at high risk (on the basis of her medical history or other findings) of developing cervical cancer, or vaginal cancer, as determined in accordance with the following risk factors:

(i) High risk factors for cervical cancer:

(A) Early onset of sexual activity (under 16 years of age).
(B) Multiple sexual partners (five or more in a lifetime).
(C) History of a sexually transmitted disease (including HIV infection).
(D) Absence of three negative or any Pap smears within the previous 7 years.

(ii) High risk factor for vaginal cancer: DES (diethylstilbestrol)-exposed daughters of women who took DES during pregnancy.

(3) More frequent screening for women of childbearing age. Subject to the limitation as specified in paragraph (b)(4) of this section, payment may be made for a screening pelvic examination performed more frequently than once every 36 months if the test is performed by a physician or other practitioner as specified in paragraph (a) of this section for a woman of childbearing age who has had such an examination that indicated the presence of cervical or vaginal cancer or other abnormality during any of the preceding 3 years. The term ‘woman of childbearing age’ means a woman who is premenopausal, and has been determined by a physician, or qualified practitioner as specified in paragraph (a) of this section, to be of childbearing age, based on her medical history or other findings.

(4) Limitation applicable to women at high risk and those of childbearing age.
§ 410.57 Pneumococcal vaccine and flu vaccine.

(a) Medicare Part B pays for pneumococcal vaccine and its administration when reasonable and necessary for the prevention of disease, if the vaccine is ordered by a doctor of medicine or osteopathy.

(b) Medicare Part B pays for the influenza virus vaccine and its administration.

§ 410.58 Additional services to HMO and CMP enrollees.

Services not usually covered under Medicare Part B may be covered as medical and other health services if they are furnished to an enrollee of an HMO or a CMP and the following conditions are met:

(a) The services are—

(1) Furnished by a physician assistant or nurse practitioner as defined in § 491.2 of this chapter, or are incident to services furnished by such a practitioner; or

(2) Furnished by a clinical psychologist as defined in § 417.416 of this chapter to an enrollee of an HMO or CMP that participates in Medicare under a risk-sharing contract, or are incident to those services.

(b) The services are services that would be covered under Medicare Part B if they were furnished by a physician or as incident to a physician’s professional services.

§ 410.59 Outpatient occupational therapy services: Conditions.

(a) Basic rule. Medicare Part B pays for outpatient occupational therapy services if they meet the following conditions:

(1) They are furnished to a beneficiary while he or she is under the care of a physician who is a doctor of medicine, osteopathy, or podiatric medicine.

(2) They are furnished under a written plan of treatment that meets the requirements of § 410.61.

(3) They are furnished—

(i) By a provider as defined in § 491.2 of this chapter, or by others under arrangements with, and under the supervision of, a provider; or

(ii) By or under the personal supervision of an occupational therapist in private practice as described in paragraph (c) of this section.

(b) Outpatient occupational therapy services furnished to certain inpatients of a hospital or a CAH or SNF. Medicare Part B pays for outpatient occupational therapy services furnished to an inpatient of a hospital, CAH, or SNF who requires them but who has exhausted or is otherwise ineligible for benefit days under Medicare Part A.

(c) Special provisions for services furnished by occupational therapists in private practice.

(1) Basic qualifications. In order to qualify under Medicare as a supplier of outpatient occupational therapy services, each individual occupational therapist in private practice must meet the following requirements:

(i) Be legally authorized (if applicable, licensed, certified, or registered) to engage in the private practice of occupational therapy by the State in which he or she practices, and practice only within the scope of his or her license, certification, or registration.

(ii) Engage in the private practice of occupational therapy on a regular basis as an individual, in one of the following practice types:

(A) An unincorporated solo practice.

(B) A partnership or unincorporated group practice.

(C) An unincorporated solo practice, partnership, or group practice, a professional corporation or other incorporated occupational therapy practice. Private practice does not include any individual during the time he or she is working as an employee of a provider.
(iii) Bill Medicare only for services furnished in his or her private practice office space, or in the patient's home. A therapist's private practice office space refers to the location(s) where the practice is operated, in the State(s) where the therapist (and practice, if applicable) is legally authorized to furnish services, during the hours that the therapist engages in practice at that location. When services are furnished in private practice office space, that space must be owned, leased, or rented by the practice and used for the exclusive purpose of operating the practice. A patient's home does not include any institution that is a hospital, an CAH, or a SNF.

(iv) Treat individuals who are patients of the practice and for whom the practice collects fees for the services furnished.

(2) Supervision of occupational therapy services. Occupational therapy services are performed by, or under the personal supervision of, the occupational therapist in private practice. All services not performed personally by the therapist must be performed by employees of the practice, personally supervised by the therapist, and included in the fee for the therapist's services.

(d) Excluded services. No service is included as an outpatient occupational therapy service if it would not be included as an inpatient hospital service if furnished to a hospital or CAH inpatient.

(e) Annual limitation on incurred expenses. (1) Amount of limitation. (i) In 1999, 2000, and 2001, no more than $1,500 of allowable charges incurred in a calendar year for outpatient occupational therapy services are recognized incurred expenses.

(ii) In 2002 and thereafter, the limitation is determined by increasing the limitation in effect in the previous calendar year by the increase in the Medicare Economic Index for the current year.

(2) For purposes of applying the limitation, outpatient occupational therapy includes:

(i) Except as provided in paragraph (e)(3) of this section, outpatient occupational therapy services furnished under this section;

(ii) Outpatient occupational therapy services furnished by a comprehensive outpatient rehabilitation facility;

(iii) Outpatient occupational therapy services furnished by a physician or incident to a physician's service;

(iv) Outpatient occupational therapy services furnished by a nurse practitioner, clinical nurse specialist, or physician assistant or incident to their services.

(3) For purposes of applying the limitation, outpatient occupational therapy services excludes services furnished by a hospital directly or under arrangements.

[63 FR 58906, Nov. 2, 1998]

§ 410.60 Outpatient physical therapy services: Conditions.

(a) Basic rule. Medicare Part B pays for outpatient physical therapy services if they meet the following conditions:

(1) They are furnished to a beneficiary while he or she is under the care of a physician who is a doctor of medicine, osteopathy, or podiatric medicine.

(2) They are furnished under a written plan of treatment that meets the requirements of § 410.61.

(3) They are furnished—

(i) By a provider as defined in § 489.2 of this chapter, or by others under arrangements with, and under the supervision of, a provider; or

(ii) By or under the personal supervision of a physical therapist in private practice as described in paragraph (c) of this section.

(b) Outpatient physical therapy services furnished to certain inpatients of a hospital or a CAH or SNF. Medicare Part B pays for outpatient physical therapy services furnished to an inpatient of a hospital, CAH, or SNF who requires them but who has exhausted or is otherwise ineligible for benefit days under Medicare Part A.

(c) Special provisions for services furnished by physical therapists in private practice. (1) Basic qualifications. In order to qualify under Medicare as a supplier of outpatient physical therapy services, each individual physical therapist in private practice must meet the following requirements:
§ 410.61 Plan of treatment requirements for outpatient rehabilitation services.

(a) Basic requirement. Outpatient rehabilitation services (including services furnished by a qualified physical or occupational therapist in private practice), must be furnished under a written plan of treatment that meets the requirements of paragraphs (b) through (e) of this section.

(b) Establishment of the plan. The plan is established before treatment is begun by one of the following:

(1) A physician.

(e) Annual limitation on incurred expenses. (1) Amount of limitation. (i) In 1999, 2000, and 2001, no more than $1,500 of allowable charges incurred in a calendar year for outpatient physical therapy services are recognized incurred expenses.

(ii) In 2002 and thereafter, the limitation shall be determined by increasing the limitation in effect in the previous calendar year by the increase in the Medicare Economic Index for the current year.

(2) For purposes of applying the limitation, outpatient physical therapy includes:

(i) Except as provided in paragraph (e)(3) of this section, outpatient physical therapy services furnished under this section;

(ii) Except as provided in paragraph (e)(3) of this section outpatient speech-language pathology services furnished under § 410.62;

(iii) Outpatient physical therapy and speech-language pathology services furnished by a comprehensive outpatient rehabilitation facility;

(iv) Outpatient physical therapy and speech-language pathology services furnished by a physician or incident to a physician’s service;

(v) Outpatient physical therapy and speech-language pathology services furnished by a nurse practitioner, clinical nurse specialist, or physician assistant or incident to their services.

(3) For purposes of applying the limitation, outpatient physical therapy excludes services furnished by a hospital or CAH directly or under arrangements.

[63 FR 58906, Nov. 2, 1998]
§ 410.62 Outpatient speech-language pathology services: Conditions and exclusions.

(a) Basic rule. Medicare Part B pays for outpatient speech pathology services if they meet the following conditions:

(1) They are furnished to a beneficiary while he or she is under the care of a physician who is a doctor of medicine or osteopathy.

(2) They are furnished under a written plan of treatment that—

(i) Is established by a physician or, effective January 1, 1982, by either a physician or the speech pathologist who will provide the services to the particular individual;

(ii) Is periodically reviewed by a physician; and

(iii) Meets the requirements of § 410.63.

(3) They are furnished by a provider as defined in § 489.2 of this chapter or by others under arrangements with, or under the supervision of, a provider.

(b) Outpatient speech pathology services to certain inpatients of a hospital, CAH, or SNF. Medicare Part B pays for outpatient speech pathology services furnished to an inpatient of a hospital, CAH, or SNF who requires them but has exhausted or is otherwise ineligible for benefit days under Medicare Part A.

(c) Excluded services. No service is included as an outpatient speech pathology service if it would not be included as an inpatient hospital service if furnished to a hospital or CAH inpatient.

(d) Limitation. After 1998, outpatient speech-language pathology services are subject to the limitation in § 410.60(e).


§ 410.63 Hepatitis B vaccine and blood clotting factors: Conditions.

Notwithstanding the exclusion from coverage of vaccines (see § 405.310 of this chapter) and self-administered drugs (see § 410.29), the following services are included as medical and other health services covered under § 410.10, subject to the specified conditions:

(a) Hepatitis B vaccine: Conditions. Effective September 1, 1984, hepatitis B vaccinations that are reasonable and necessary for the prevention of illness for those individuals who are at high or intermediate risk of contracting hepatitis B as listed below:
(1) High risk groups. (i) End-Stage Renal Disease (ESRD) patients; (ii) Hemophiliacs who receive Factor VIII or IX concentrates; (iii) Clients of institutions for the mentally retarded; (iv) Persons who live in the same household as a hepatitis B carrier; (v) Homosexual men; (vi) Illicit injectable drug abusers; and (vii) Pacific Islanders (that is, those Medicare beneficiaries who reside on Pacific islands under U.S. jurisdiction, other than residents of Hawaii).

(2) Intermediate risk groups. (i) Staff in institutions for the mentally retarded and classroom employees who work with mentally retarded persons; (ii) Workers in health care professions who have frequent contact with blood or blood-derived body fluids during routine work (including workers who work outside of a hospital and have frequent contact with blood or other infectious secretions); and (iii) Heterosexually active persons with multiple sexual partners (that is, those Medicare beneficiaries who have had at least two documented episodes of sexually transmitted diseases within the preceding 5 years).

(3) Exception. Individuals described in paragraphs (a) (1) and (2) of this section are not considered at high or intermediate risk of contracting hepatitis B if they have undergone a prevaccination screening and have been found to be currently positive for antibodies to hepatitis B.

(b) Blood clotting factors. Effective July 18, 1984, blood clotting factors to control bleeding for hemophilia patients competent to use these factors without medical or other supervision, and items related to the administration of those factors. The amount of clotting factors covered under this provision is determined by the carrier based on the historical utilization pattern or profile developed by the carrier for each patient, and based on consideration of the need for a reasonable reserve supply to be kept in the home in the event of emergency or unforeseen circumstance.

§ 410.64 Services related to cardiac pacemakers and pacemaker leads.

(a) Requirement. (1) Physicians or providers that request or receive payment for the implantation, removal, or replacement of permanent cardiac pacemakers and pacemaker leads, must submit to HCFA the information required for the pacemaker registry.

(2) The required information is set forth under 21 CFR part 805 of the FDA regulations and must be submitted in accordance with general instructions issued by HCFA.

(b) Denial of payment. If HCFA finds that a physician or provider has failed to comply with paragraph (a) of this section, HCFA will deny payment for the implantation, removal, or replacement of any permanent cardiac pacemaker or pacemaker lead, effective 45 days after sending the physician or provider written notice in accordance with paragraph (c) of this section.

(c) Notice of denial of payment. The notice of denial of payment— (1) States the reasons for the determination; (2) Grants the physician or provider 45 days from the date of the notice to submit the information or evidence showing that the determination is in error; and (3) Informs the physician or provider of its right to hearing.

(d) Right to hearing. If the denial of payment goes into effect at the expiration of the 45-day period, it constitutes an “initial determination” subject to administrative and judicial review under part 498 of this chapter.

§ 410.66 Emergency outpatient services furnished by a nonparticipating hospital and services furnished in Mexico or Canada.

Conditions for payment of emergency outpatient services furnished by a nonparticipating U.S. hospital and for services furnished in Mexico or Canada are set forth in subparts G and H of part 424 of this chapter.
§ 410.68 Antigens: Scope and conditions.

Medicare Part B pays for—
(a) Antigens that are furnished as services incident to a physician's professional services; or
(b) A supply of antigen sufficient for not more than 12 weeks that is—
(1) Prepared for a patient by a doctor of medicine or osteopathy who has examined the patient and developed a plan of treatment including dosage levels; and
(2) Administered—
(i) In accord with the plan of treatment developed by the doctor of medicine or osteopathy who prepared the antigen; and
(ii) By a doctor of medicine or osteopathy or by a properly instructed person under the supervision of a doctor of medicine or osteopathy.

[54 FR 4026, Jan. 27, 1989]

§ 410.69 Services of a certified registered nurse anesthetist or an anesthesiologist's assistant: Basic rule and definitions.

(a) Basic rule. Medicare Part B pays for anesthesia services and related care furnished by a certified registered nurse anesthetist or an anesthesiologist's assistant who is legally authorized to perform the services by the State in which the services are furnished.

(b) Definitions. For purposes of this part—
Anesthesiologist's assistant means a person who—
(1) Works under the direction of an anesthesiologist;
(2) Is in compliance with all applicable requirements of State law, including any licensure requirements the State imposes on nonphysician anesthetists; and
(3) Is a graduate of a medical school-based anesthesiologist's assistant educational program that—
(A) Is accredited by the Committee on Allied Health Education and Accreditation; and
(B) Includes approximately two years of specialized basic science and clinical education in anesthesia at a level that builds on a premedical undergraduate science background.

Anesthetist includes both an anesthesiologist's assistant and a certified registered nurse anesthetist.

Certified registered nurse anesthetist means a registered nurse who:
(1) Is licensed as a registered professional nurse by the State in which the nurse practices;
(2) Meets any licensure requirements the State imposes with respect to nonphysician anesthetists;
(3) Has graduated from a nurse anesthesia educational program that meets the standards of the Council on Accreditation of Nurse Anesthesia Programs, or such other accreditation organization as may be designated by the Secretary; and
(4) Meets the following criteria:
(i) Has passed a certification examination of the Council on Certification of Nurse Anesthetists, the Council on Recertification of Nurse Anesthetists, or any other certification organization that may be designated by the Secretary; or
(ii) Is a graduate of a program described in paragraph (3) of this definition and within 24 months after that graduation meets the requirements of paragraph (4)(i) of this definition.

[57 FR 33896, July 31, 1992]

§ 410.71 Clinical psychologist services and services and supplies incident to clinical psychologist services.

(a) Included services. (1) Medicare Part B covers services furnished by a clinical psychologist, who meets the requirements specified in paragraph (d) of this section, that are within the scope of his or her State license, if the services would be covered if furnished by a physician or as an incident to a physician's services.

(2) Medicare Part B covers services and supplies furnished as an incident to the services of a clinical psychologist if the following requirements are met:
(i) The services and supplies would be covered if furnished by a physician or as an incident to a physician's services.
(ii) The services or supplies are of the type that are commonly furnished in a physician's or clinical psychologist's office and are either furnished without charge or are included in the physician's or clinical psychologist's bill.
(iii) The services are an integral, although incidental, part of the professional services performed by the clinical psychologist.

(iv) The services are performed under the direct supervision of the clinical psychologist. For example, when services are performed in the clinical psychologist’s office, the clinical psychologist must be present in the office suite and immediately available to provide assistance and direction throughout the time the service is being performed.

(v) The individual performing the service must be an employee of either the clinical psychologist or the legal entity that employs the supervising clinical psychologist, under the common law control test of the Act as more fully set forth in 20 CFR 404.1007.

(b) Application of mental health treatment limitation. The treatment services of a clinical psychologist and services and supplies furnished as an incident to those services are subject to the limitation on payment for outpatient mental health treatment services set forth in §410.155.

(c) Payment for consultations. A clinical psychologist or an attending or primary care physician may not bill Medicare or the beneficiary for the consultation that is required under paragraph (e) of this section.

(d) Qualifications. For purposes of this subpart, a clinical psychologist is an individual who—

(1) Holds a doctoral degree in psychology; and

(2) Is licensed or certified, on the basis of the doctoral degree in psychology, by the State in which he or she practices, at the independent practice level of psychology to furnish diagnostic, assessment, preventive, and therapeutic services directly to individuals.

(e) Agreement to consult. A clinical psychologist who bills Medicare Part B must agree to meet the requirements of paragraphs (e)(1) through (e)(3) of this section. The clinical psychologist’s signature on a Medicare provider/supplier enrollment form indicates his or her agreement.

(i) Unless the beneficiary’s primary care or attending physician has referred the beneficiary to the clinical psychologist, to inform the beneficiary that it is desirable for the clinical psychologist to consult with the beneficiary’s attending or primary care physician (if the beneficiary has such a physician) to consider any conditions contributing to the beneficiary’s symptoms.

(2) If the beneficiary assents to the consultation, in accordance with accepted professional ethical norms and taking into consideration patient confidentiality—

(i) To attempt, within a reasonable time after receiving the consent, to consult with the physician; and

(ii) If attempts to consult directly with the physician are not successful, to notify the physician, within a reasonable time, that he or she is furnishing services to the beneficiary.

(3) Unless the primary care or attending physician referred the beneficiary to the clinical psychologist, to document, in the beneficiary’s medical record, the date the patient consented or declined consent to consultation, the date of consultation, or, if attempts to consult did not succeed, the date and manner of notification to the physician.

[63 FR 20128, Apr. 23, 1998]

§410.73 Clinical social worker services.

(a) Definition: clinical social worker. For purposes of this part, a clinical social worker is defined as an individual who—

(1) Possesses a master’s or doctor’s degree in social work;

(2) After obtaining the degree, has performed at least 2 years of supervised clinical social work; and

(3) Either is licensed or certified as a clinical social worker by the State in which the services are performed or, in the case of an individual in a State that does not provide for licensure or certification as a clinical social worker—

(i) Is licensed or certified at the highest level of practice provided by the laws of the State in which the services are performed; and

(ii) Has completed at least 2 years or 3,000 hours of post master’s degree supervised clinical social work practice
§ 410.74 Physician assistants’ services.

(a) Basic rule. Medicare Part B covers physician assistants’ services only if the following conditions are met:

(1) The services would be covered as physicians’ services if furnished by a physician (a doctor of medicine or osteopathy, as set forth in section 1861(r)(1) of the Act).

(2) The physician assistant—
   (i) Meets the qualifications set forth in paragraph (c) of this section;
   (ii) Is legally authorized to perform the services in the State in which they are performed;
   (iii) Performs services that are not otherwise precluded from coverage because of a statutory exclusion;
   (iv) Performs the services under the general supervision of a physician (the supervising physician need not be physically present when the physician assistant is performing the services unless required by State law; however, the supervising physician must be immediately available to the physician assistant for consultation);
   (v) Furnishes services that are billed by the employer of a physician assistant; and
   (vi) Performs the services—
      (A) In all settings in either rural and urban areas; or
      (B) As an assistant at surgery.

(b) Services and supplies furnished incident to a physician assistant’s services. Medicare covers services and supplies (including drugs and biologicals that cannot be self-administered) that are furnished incident to the physician assistant’s services described in paragraph (a) of this section. These services and supplies are covered only if they—

(1) Would be covered if furnished by a physician or as incident to the professional services of a physician;

(2) Are the type that are commonly furnished in a physician’s office and are either furnished without charge or are included in the bill for the physician assistants’ services;

(3) Are, although incidental, an integral part of the professional service performed by the physician;

(4) Are performed under the direct supervision of the physician assistant (that is, the physician assistant is physically present and immediately available); and

(5) Are performed by the employee of a physician assistant or an entity that employs both the physician assistant and the person providing the services.
§ 410.75 Nurse practitioners’ services.

(a) Definition. As used in this section, the term “physician” means a doctor of medicine or osteopathy, as set forth in section 1861(r)(1) of the Act.

(b) Qualifications. For Medicare Part B coverage of his or her services, a nurse practitioner must—
(1) Be legally authorized to perform them in the State in which they are performed;
(2) Be authorized by the State in which the services are furnished to practice as a nurse practitioner in accordance with State law; and
(3) Be certified as a nurse practitioner by a recognized national certifying body that has established standards for nurse practitioners; or
(4) Be a registered professional nurse who is authorized by the State in which the services are furnished to practice as a nurse practitioner in accordance with State law and have been granted a Medicare billing number as a nurse practitioner by December 31, 2000, or
(3) Be a nurse practitioner who on or after January 1, 2001, applies for a Medicare billing number after the first time and meets the standards for nurse practitioners in paragraphs (b)(1)(i) and (b)(1)(ii) of this section; or
(4) Be a nurse practitioner who on or after January 1, 2003, applies for a Medicare billing number for the first time and possesses a master’s degree in nursing and meets the standards for nurse practitioners in paragraphs (b)(1)(i) and (b)(1)(ii) of this section.

(c) Services. Medicare Part B covers nurse practitioners’ services in all settings in both rural and urban areas, only if the services would be covered if furnished by a physician and the nurse practitioner—
(1) Is legally authorized to perform them in the State in which they are performed;
(2) Is not performing services that are otherwise excluded from coverage because of one of the statutory exclusions; and
(3) Performs them while working in collaboration with a physician.

(i) Collaboration is a process in which a nurse practitioner works with one or more physicians to deliver health care services within the scope of the practitioner’s expertise, with medical direction and appropriate supervision as provided for in jointly developed guidelines or other mechanisms as provided by the law of the State in which the services are performed.

(ii) In the absence of State law governing collaboration, collaboration is a process in which a nurse practitioner has a relationship with one or more physicians to deliver health care services. Such collaboration is to be evidenced by nurse practitioners documenting the nurse practitioners’ scope of practice and indicating the relationships that they have with physicians to deal with issues outside their scope of practice. Nurse practitioners must document this collaborative process with physicians. 

[63 FR 58907, Nov. 2, 1998; 64 FR 25457, May 12, 1999]
(iii) The collaborating physician does not need to be present with the nurse practitioner when the services are furnished or to make an independent evaluation of each patient who is seen by the nurse practitioner.

(d) Services and supplies incident to a nurse practitioners' services. Medicare Part B covers services and supplies (including drugs and biologicals that cannot be self-administered) incident to a nurse practitioner's services that meet the requirements in paragraph (c) of this section. These services and supplies are covered only if they—

(1) Would be covered if furnished by a physician or as incident to the professional services of a physician;
(2) Are of the type that are commonly furnished in a physician's office and are either furnished without charge or are included in the bill for the nurse practitioner's services;
(3) Although incidental, are an integral part of the professional service performed by the nurse practitioner; and
(4) Are performed under the direct supervision of the nurse practitioner (that is, the nurse practitioner must be physically present and immediately available).

(e) Professional services. Nurse practitioners can be paid for professional services only when the services have been personally performed by them and no facility or other provider charges, or is paid, any amount for the furnishing of the professional services.

(1) Supervision of other nonphysician staff by a nurse practitioner does not constitute personal performance of a professional service by a nurse practitioner.

(2) The services are provided on an assignment-related basis, and a nurse practitioner may not charge a beneficiary for a service not payable under this provision. If a beneficiary has made payment for a service, the nurse practitioner must make the appropriate refund to the beneficiary.


§ 410.76 Clinical nurse specialists' services.

(a) Definition. As used in this section, the term "physician" means a doctor of medicine or osteopathy, as set forth in section 1861(r)(1) of the Act.

(b) Qualifications. For Medicare Part B coverage of his or her services, a clinical nurse specialist must—

(1) Be a registered nurse who is currently licensed to practice in the State where he or she practices and be authorized to perform the services of a clinical nurse specialist in accordance with State law;
(2) Have a master's degree in a defined clinical area of nursing from an accredited educational institution; and
(3) Be certified as a clinical nurse specialist by the American Nurses Credentialing Center.

(c) Services. Medicare Part B covers clinical nurse specialists' services in all settings in both rural and urban areas only if the services would be covered if furnished by a physician and the clinical nurse specialist—

(1) Is legally authorized to perform them in the State in which they are performed;
(2) Is not performing services that are otherwise excluded from coverage by one of the statutory exclusions; and
(3) Performs them while working in collaboration with a physician.

(i) Collaboration is a process in which a clinical nurse specialist works with one or more physicians to deliver health care services within the scope of the practitioner's expertise, with medical direction and appropriate supervision as provided for in jointly developed guidelines or other mechanisms as provided by the law of the State in which the services are performed.

(ii) In the absence of State law governing collaboration, collaboration is a process in which a clinical nurse specialist has a relationship with one or more physicians to deliver health care services. Such collaboration is to be evidenced by clinical nurse specialists documenting the clinical nurse specialists' scope of practice and indicating the relationships that they have with physicians to deal with issues outside their scope of practice. Clinical nurse specialists must document this collaborative process with physicians.

(iii) The collaborating physician does not need to be present with the clinical nurse specialist when the services are furnished, or to make an independent
evaluation of each patient who is seen by the clinical nurse specialist.

(d) Services and supplies furnished incident to clinical nurse specialists’ services. Medicare Part B covers services and supplies (including drugs and biologicals that cannot be self-administered) incident to a clinical nurse specialist’s services that meet the requirements in paragraph (c) of this section. These services and supplies are covered only if they—

(1) Would be covered if furnished by a physician or as incident to the professional services of a physician;

(2) Are of the type that are commonly furnished in a physician's office and are either furnished without charge or are included in the bill for the clinical nurse specialist’s services;

(3) Although incidental, are an integral part of the professional service performed by the clinical nurse specialist; and

(4) Are performed under the direct supervision of the clinical nurse specialist (that is, the clinical nurse specialist must be physically present and immediately available).

(e) Professional services. Clinical nurse specialists can be paid for professional services only when the services have been personally performed by them and no facility or other provider charges, or is paid, any amount for the furnishing of the professional services.

(1) Supervision of other nonphysician staff by clinical nurse specialists does not constitute personal performance of a professional service by clinical nurse specialists.

(2) The services are provided on an assignment-related basis, and a clinical nurse specialist may not charge a beneficiary for a service not payable under this provision. If a beneficiary has made payment for a service, the clinical nurse specialist must make the appropriate refund to the beneficiary.

[63 FR 58908, Nov. 2, 1998]

§ 410.77 Certified nurse-midwives’ services: Qualifications and conditions.

(a) Qualifications. For Medicare coverage of his or her services, a certified nurse-midwife must:

(1) Be a registered nurse who is legally authorized to practice as a nurse-midwife in the State where services are performed;

(2) Have successfully completed a program of study and clinical experience for nurse-midwives that is accredited by an accrediting body approved by the U.S. Department of Education; and

(3) Be certified as a nurse-midwife by the American College of Nurse-Midwives or the American College of Nurse-Midwives Certification Council.

(b) Services. A certified nurse-midwife’s services are services furnished by a certified nurse-midwife and services and supplies furnished as an incident to the certified nurse-midwife’s services that—

(1) Are within the scope of practice authorized by the law of the State in which they are furnished and would otherwise be covered if furnished by a physician or as an incident to a physician’s service; and

(2) Unless required by State law, are provided without regard to whether the certified nurse-midwife is under the supervision of, or associated with, a physician or other health care provider.

(c) Incident to services: Basic rule. Medicare covers services and supplies furnished incident to the services of a certified nurse-midwife, including drugs and biologicals that cannot be self-administered, if the services and supplies meet the following conditions:

(1) They would be covered if furnished by a physician or as incident to the professional services of a physician.

(2) They are of the type that are commonly furnished in a physician’s office and are either furnished without charge or are included in the bill for the certified nurse-midwife’s services.

(3) Although incidental, they are an integral part of the professional service performed by the certified nurse-midwife.

(4) They are furnished under the direct supervision of a certified nurse-midwife (that is, the midwife is physically present and immediately available).

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(1) Supervision of other nonphysician staff by a nurse-midwife does not constitute personal performance of a professional service by the nurse-midwife.

(2) The service is provided on an assignment-related basis, and a nurse-midwife may not charge a beneficiary for a service not payable under this provision. If the beneficiary has made payment for a service, the nurse-midwife must make the appropriate refund to the beneficiary.

(3) A nurse-midwife may provide services that he or she is legally authorized to perform under State law as a nurse-midwife, if the services would otherwise be covered by the Medicare program when furnished by a physician or incident to a physicians' professional services.

[63 FR 58909, Nov. 2, 1998]

§ 410.78 Consultations via telecommunications systems.

(a) General rule. Medicare Part B pays for professional consultations furnished by means of interactive telecommunications systems if the following conditions are met:

(1) The consulting practitioner is any of the following:
   (i) A physician as described in § 410.20.
   (ii) A physician assistant as defined in § 410.74.
   (iii) A nurse practitioner as defined in § 410.75.
   (iv) A clinical nurse specialist as described in § 410.76.
   (v) A nurse-midwife as defined in § 410.77.
   (vi) A clinical psychologist as described in § 410.71.
   (vii) A clinical social worker as defined in § 410.73.

(2) The referring practitioner is any of the following:
   (i) A physician as described in § 410.20.
   (ii) A physician assistant as defined in § 410.74.
   (iii) A nurse practitioner as defined in § 410.75.
   (iv) A clinical nurse specialist as described in § 410.76.
   (v) A nurse-midwife as defined in § 410.77.

(b) Definition. For purposes of this section, interactive telecommunications systems means multimedia communications equipment that includes, at a minimum, audio and video equipment permitting real-time consultation among the patient, consultant, and referring practitioner, or a practitioner described in section 1842(b)(18)(C) of the Act (other than a certified registered nurse anesthetist or anesthesiologist assistant) who is an employee of the referring practitioner, as appropriate to the medical needs of the patient and as needed to provide information to and at the direction of the consultant.

(6) The consultation results in a written report that is furnished to the referring practitioner.

Subpart C—Home Health Services Under SMI

§ 410.80 Applicable rules.

Home health services furnished under Medicare Part B are subject to the rules set forth in subpart E of part 409 of this chapter.
§ 410.100  Included services.

Subject to the conditions and limitations set forth in §§ 410.102 and 410.105, CORF services means the following services furnished to an outpatient of the CORF by personnel that meet the qualifications set forth in § 485.70 of this chapter.

(a) Physicians' services. The following services of the facility physician constitute CORF services: consultation with and medical supervision of non-physician staff, establishment and review of the plan of treatment, and other medical and facility administration activities. Those services are reimbursed on a reasonable cost basis under part 413 of this chapter. Diagnostic and therapeutic services furnished to an individual patient are not CORF physician's services. If covered, payment for these services would be made by the carrier on a reasonable charge basis subject to the provisions of subpart E of part 405 of this chapter.

(b) Physical therapy services.

(1) These services include—

(i) Testing and measurement of the function or dysfunction of the neuromuscular, musculoskeletal, cardiovascular and respiratory systems; and.

(ii) Assessment and treatment related to dysfunction caused by illness or injury, and aimed at preventing or reducing disability or pain and restoring lost function.

(2) The establishment of a maintenance therapy program for an individual whose restoration potential has been reached is a physical therapy service; however, maintenance therapy itself is not covered as part of these services.

(c) Occupational therapy services. These services include—

(1) Teaching of compensatory techniques to permit an individual with a physical impairment or limitation to engage in daily activities.

(2) Evaluation of an individual's level of independent functioning.

(3) Selection and teaching of task-oriented therapeutic activities to restore sensory-integrative function; and

(4) Assessment of an individual's vocational potential, except when the assessment is related solely to vocational rehabilitation.

(d) Speech-language pathology services. These are services for the diagnosis and treatment of speech and language disorders that create difficulties in communication.

(e) Respiratory therapy services. (1) These are services for the assessment, diagnostic evaluation, treatment, management, and monitoring of patients with deficiencies or abnormalities of cardiopulmonary function.

(2) These services include—

(i) Application of techniques for support of oxygenation and ventilation of the patient and for pulmonary rehabilitation.

(ii) Therapeutic use and monitoring of gases, mists, and aerosols and related equipment;

(iii) Bronchial hygiene therapy;

(iv) Pulmonary rehabilitation techniques such as exercise conditioning, breathing retraining and patient education in the management of respiratory problems.

(v) Diagnostic tests to be evaluated by a physician, such as pulmonary function tests, spirometry and blood gas analysis; and

(vi) Periodic assessment of chronically ill patients and their need for respiratory therapy.

(f) Prosthetic device services. These services include—

(1) Prosthetic devices (excluding dental devices and renal dialysis machines), that replace all or part of an internal body organ or external body member (including contiguous tissue) or replace all or part of the function of a permanently inoperative or malfunctioning external body member or internal body organ; and

(2) Services necessary to design the device, select materials and components, measure, fit, and align the device, and instruct the patient in its use.

(g) Orthotic device services. These services include—

(1) Orthopedic devices that support or align movable parts of the body, prevent or correct deformities, or improve functioning; and
(2) Services necessary to design the device, select the materials and components, measure, fit, and align the device, and instruct the patient in its use.

(h) Social services. These services include—

(1) Assessment of the social and emotional factors related to the individual's illness, need for care, response to treatment, and adjustment to care furnished by the facility;

(2) Casework services to assist in resolving social or emotional problems that may have an adverse effect on the beneficiary's ability to respond to treatment; and

(3) Assessment of the relationship of the individual's medical and nursing requirements to his or her home situation, financial resources, and the community resources available upon discharge from facility care.

(i) Psychological services. These services include—

(1) Assessment, diagnosis and treatment of an individual's mental and emotional functioning as it relates to the individual's rehabilitation;

(2) Psychological evaluations of the individual's response to and rate of progress under the treatment plan; and

(3) Assessment of those aspects of an individual's family and home situation that affect the individual's rehabilitation treatment.

(j) Nursing care services. These services include nursing services specified in the plan of treatment and any other nursing services necessary for the attainment of the rehabilitation goals.

(k) Drugs and biologicals. These are drugs and biologicals that are—

(1) Prescribed by a physician and administered by or under the supervision of a physician or a registered professional nurse; and

(2) Not excluded from Medicare Part B payment for reasons specified in §410.29.

(l) Supplies, appliances, and equipment. These include—

(1) Non-reusable supplies such as oxygen and bandages;

(2) Medical equipment and appliances; and

(3) Durable medical equipment of the type specified in §410.38, (except renal dialysis systems) for use outside the CORF, whether purchased or rented.

(m) Home environment evaluation. This is a single home visit to evaluate the potential impact of the home situation on the rehabilitation goals.


§ 410.102 Excluded services.

None of the services specified in §410.100 is covered as a CORF service if the service—

(a) Would not be covered as an inpatient hospital service if furnished to a hospital inpatient;

(b) Is not reasonable and necessary for the diagnosis or treatment of illness or injury or to improve the functioning of a malformed body member. An example would be services furnished as part of a maintenance program involving repetitive activities that do not require the skilled services of nurses or therapists.

§ 410.105 Requirements for coverage of CORF services.

Services specified in §410.100 and not excluded under §410.102 are covered as CORF services if they are furnished by a participating CORF (that is, a CORF that meets the conditions of subpart B of part 485 of this chapter, and has in effect a provider agreement under part 489 of this chapter) and if the following requirements are met:

(a) Referral and medical history. The services must be furnished to an individual who is referred by a physician who certifies that the individual needs skilled rehabilitation services, and makes the following information available to the CORF before or at the time treatment is begun:

(1) The individual's significant medical history.

(2) Current medical findings.

(3) Diagnosis(es) and contraindications to any treatment modality.

(4) Rehabilitation goals, if determined.

(b) When and where services are furnished. (1) All services must be furnished while the individual is under the care of a physician.

(2) Except as provided in paragraph (b)(3) of this section, the services must
§ 410.110

be furnished on the premises of the CORF.

(3) Exceptions. (i) Physical therapy, occupational therapy, and speech pathology services may be furnished away from the premises of the CORF.

(ii) The single home visit specified in §410.100(m) is also covered.

(c) Plan of treatment. (1) The services must be furnished under a written plan of treatment that—

(i) Is established and signed by a physician before treatment is begun; and

(ii) Prescribes the type, amount, frequency, and duration of the services to be furnished, and indicates the diagnosis and anticipated rehabilitation goals.

(2) The plan must be reviewed at least every 60 days by a facility physician who, when appropriate, consults with the professional personnel providing the services.

(3) The reviewing physician must certify or recertify that the plan is being followed, the patient is making progress in attaining the rehabilitation goals, and the treatment is having no harmful effects on the patient.


Subpart E—Community Mental Health Centers (CMHCs) Providing Partial Hospitalization Services

§ 410.110 Requirements for coverage of partial hospitalization services by CMHCs.

Medicare part B covers partial hospitalization services furnished by or under arrangements made by a CMHC if they are provided by a CMHC as defined in §410.2 that has in effect a provider agreement under part 409 of this chapter and if the services are—

(a) Prescribed by a physician and furnished under the general supervision of a physician;

(b) Subject to certification by a physician in accordance with §424.24(e)(1) of this subchapter; and

(c) Furnished under a plan of treatment that meets the requirements of §424.24(e)(2) of this subchapter.

[59 FR 6577, Feb. 11, 1994]

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Subparts F–H—[Reserved]

Subpart I—Payment of SMI Benefits


§ 410.150 To whom payment is made.

(a) General rules. (1) Any SMI enrollee is subject to the conditions, limitations, and exclusions set forth in this part and in parts 405, 416 and 424 of this chapter, entitled to have payment made as specified in paragraph (b) of this section.

(2) The services specified in paragraphs (b)(5) through (b)(14) of this section must be furnished by a facility that has in effect a provider agreement or other appropriate agreement to participate in Medicare.

(b) Specific rules. Subject to the conditions set forth in paragraph (a) of this section, Medicare Part B pays as follows:

(1) To the individual, or to a physician or other supplier on the individual's behalf, for medical and other health services furnished by the physician or other supplier.

(2) To a nonparticipating hospital on the individual's behalf for emergency outpatient services furnished by the hospital, in accordance with subpart G of part 424 of this chapter.

(3) To the individual, for emergency outpatient services furnished by a nonparticipating hospital, in accordance with §424.53 of this chapter.

(4) To the individual, for physicians' services and ambulance services furnished outside the United States in accordance with §424.53 of this chapter.

(5) To a provider on the individual's behalf for medical and other health services furnished by the provider (or by others under arrangements made with them by the provider).

(6) To a home health agency on the individual's behalf for home health services furnished by the home health agency.

(7) To a clinic, rehabilitation agency, or public health agency on the individual's behalf for outpatient physical therapy or speech pathology services furnished by the clinic or agency (or by...
§ 410.152 Amounts of payment.

(a) General provisions—(1) Exclusion from incurred expenses. As used in this section, “incurred expenses” are expenses incurred by an individual, during his or her coverage period, for covered Part B services, excluding the following:

(i) Expenses incurred for services for which the beneficiary is entitled to have payment made under Medicare Part A or would be so entitled except for the application of the Part A deductible and coinsurance requirements.

(ii) Expenses incurred in meeting the Part B blood deductible (§ 410.161).

(iii) In the case of services payable under a formula that takes into account reasonable charges, reasonable costs, customary charges, customary (insofar as reasonable) charges, charges related to reasonable costs, fair compensation, a pre-treatment prospective payment rate, or a standard overhead amount, or any combination of two or more of these factors, expenses in excess of any factor taken into account under that formula.

(iv) Expenses in excess of the outpatient mental health treatment limitation described in § 410.155.

(v) In the case of expenses incurred for outpatient physical therapy services including speech-language pathology services, the expenses excluded are from the incurred expenses under § 410.60(e). In the case of expenses incurred for outpatient occupational therapy including speech-language pathology services, the expenses excluded are from the incurred expenses under § 410.59(e).

(ii) Other applicable provisions. Medicare Part B pays for incurred expenses the amounts specified in paragraphs (b) through (k) of this section, subject to the following:

(i) The principles and procedures for determining reasonable costs and reasonable charges and the conditions for Medicare payment, as set forth in parts 405 (subparts E and X), 413, and 424 of this chapter.


(iii) The special rules for payment to health maintenance organizations (HMOs), health care prepayment plans (HCPPs), and competitive medical plans (CMPs) that are set forth in part 417 of this chapter. (A prepayment organization that does not qualify as an HMO, CMP, or HCPP is paid in accordance with paragraph (b)(4) of this section.)

(b) Basic rules for payment. Except as specified in paragraphs (c) through (h) of this section, Medicare Part B pays the following amounts:

(1) For services furnished by, or under arrangements made by, a provider other than a nominal charge provider, whichever of the following is less:

(i) 80 percent of the reasonable cost of the services.

(ii) The reasonable cost of, or the customary charges for, the services, whichever is less, minus 20 percent of the customary (insofar as reasonable) charges for the services.

(2) For services furnished by, or under arrangements made by, a nominal charge provider, 80 percent of fair compensation.

(3) For emergency outpatient hospital services furnished by a non-participating hospital that is eligible to receive payment for those services under subpart G of part 424 of this chapter, the amount specified in paragraph (b)(1) of this section.

(4) For services furnished by a person or an entity other than those specified in paragraphs (b)(1) through (b)(3) of this section, 80 percent of the reasonable charges or 80 percent of the payment amount computed on any other payment basis for the services.

(c) Amount of payment: Home health services other than durable medical equipment (DME). For home health services other than DME furnished by, or under arrangements made by, a participating HHA, Medicare Part B pays the following amounts:

(1) For services furnished by an HHA that is a nominal charge provider, 100 percent of fair compensation.

(2) For services furnished by an HHA that is not a nominal charge provider, the lesser of the reasonable cost of the services and the customary charges for the services.

(d) Amount of payment: DME furnished as a home health service.

(1) Basic rule. Except as specified in paragraph (d)(2) of this section—

(i) For DME furnished by an HHA that is a nominal charge provider, Medicare Part B pays 80 percent of fair compensation.

(ii) For DME furnished by an HHA that is not a nominal charge provider, Medicare Part B pays the lesser of the following:

(A) 80 percent of the reasonable cost of the service.

(B) The reasonable cost of, or the customary charge for, the service, whichever is less, minus 20 percent of
For services furnished before July 1, 1987, Medicare Part B paid 100 percent of the standard amount.

(2) Exception. If the DME is used DME purchased by or on behalf of the beneficiary at a price at least 25 percent less than the reasonable charge for new equipment—

(i) For used DME furnished by an HHA that is a nominal charge provider, Medicare Part B pays 100 percent of fair compensation.

(ii) For used DME furnished by an HHA that is not a nominal charge provider, Medicare Part B pays 100 percent of the reasonable cost of, or the customary charge for, the services, whichever is less.

(e) Amount of payment: Renal dialysis services, supplies, and equipment. Effective for services furnished on or after August 1, 1983, Medicare Part B pays for the institutional dialysis services specified in §409.250 and the home dialysis services, supplies, and equipment specified in §409.252, as follows:

(1) Except as provided in paragraph (d)(2) of this section, 80 percent of the per treatment prospective reimbursement rate established under §413.170 of this chapter, for outpatient maintenance dialysis furnished by ESRD facilities approved in accordance with subpart U of part 405 of this chapter.

(2) Exception. If a home dialysis patient elects to obtain home dialysis supplies or equipment (or both) from a party other than an approved ESRD facility, payment is in accordance with paragraph (b)(4) of this section.

(f) Amount of payment: Rural health clinic and Federally qualified health center services. Medicare Part B pays, for services by a participating independent rural health clinic or Federally qualified health center, 80 percent of the costs determined under subpart X of part 405 of this chapter, to the extent those costs are reasonable and related to the cost of furnishing rural health clinic or Federally qualified health center services or reasonable on the basis of other tests specified by HCFA.

(g) Amount of payment: Used durable medical equipment furnished by other than an HHA. Medicare Part B pays the following amounts for used DME purchased by or on behalf of the beneficiary at a price at least 25 percent less than the reasonable charge for comparable new equipment:

(1) For used DME furnished by, or under arrangements made by, a nominal charge provider, 100 percent of fair compensation.

(2) For used DME furnished by, or under arrangements made by a provider that is not a nominal charge provider, 100 percent of the reasonable cost of the service or the customary charge for the service, whichever is less.

(3) For used DME furnished by other than a provider, 100 percent of the reasonable charge.

(h) Amount of payment: Pneumococcal vaccine. Medicare Part B pays for pneumococcal vaccine and its administration as follows:

(1) For services furnished by a nominal charge provider, 100 percent of fair compensation.

(2) For services furnished by a provider that is not a nominal charge provider, the reasonable cost of the services or the customary charge for the service, whichever is less.

(3) For services furnished by other than a provider, a rural health clinic or a Federally qualified health center, 100 percent of the reasonable charge.

(4) For services furnished by a rural health clinic or a Federally qualified health center, 100 percent of the reasonable cost.

(i) Amount of payment: ASC facility services. For ASC facility services that are furnished in connection with the surgical procedures specified in part 416 of this chapter, Medicare Part B pays 80 percent of a standard overhead amount, as specified in §416.120(c) of this chapter.

(j) Amount of payment: services of Federally funded health facilities prior to October 1, 1991. Medicare Part B pays 80 percent of charges related to the reasonable costs that a Federally funded health facility incurs in furnishing the services. See §411.8(b)(6) of this chapter.

(k) Amount of payment: Outpatient CAH services. (1) Payment for CAH outpatient services is the reasonable cost.
of the CAH in providing these services, as determined in accordance with section 1861(v)(1)(A) of the Act, with §413.70(b) and (c) of this chapter, and with the applicable principles of cost reimbursement in part 413 and in part 415 of this chapter.

(2) Payment for CAH outpatient services is subject to the applicable Medicare Part B deductible and coinsurance amounts, except as described in §413.70(b)(2)(iii)(B) of this chapter, with Part B coinsurance being calculated as 20 percent of the customary (if reasonable) charges of the CAH for the services.

(l) Amount of payment: Flu vaccine. Medicare Part B pays 100 percent of the Medicare allowed charge.

§410.155 Outpatient mental health treatment limitation.

(a) Limitation. Only 62 1/2 percent of the expenses incurred for services subject to the limit as specified in paragraph (b) of this section are considered incurred expenses under Medicare Part B when determining the amount of payment and deductible under §§410.152 and 410.160, respectively.

(b) Application of the limitation—(1) Services subject to the limitation. Except as specified in paragraph (b)(2) of this section, the following services are subject to the limitation if they are furnished in connection with the treatment of a mental, psychoneurotic, or personality disorder (that is, any condition identified by a diagnosis code within the range of 290 through 319) and are furnished to an individual who is not an inpatient of a hospital:

(i) Services furnished by physicians and other practitioners, whether furnished directly or as an incident to those practitioners’ services.

(ii) Services provided by a CORF.

(2) Services not subject to the limitation. Services not subject to the limitation include the following:

(i) Services furnished to a hospital inpatient.

(ii) Brief office visits for the sole purpose of monitoring or changing drug prescriptions used in the treatment of mental, psychoneurotic, or personality disorders.

(iii) Partial hospitalization services not directly provided by a physician.

(iv) Diagnostic services, such as psychological testing, that are performed to establish a diagnosis.

(v) Medical management, as opposed to psychotherapy, furnished to a patient diagnosed with Alzheimer’s disease or a related disorder.

(c) Examples. (1) A clinical psychologist submitted a claim for $200 for outpatient treatment of a beneficiary’s mental disorder. The Medicare approved amount was $180. Since clinical psychologists must accept assignment, the beneficiary is not liable for the $20 in excess charges. The beneficiary previously satisfied the $100 annual Part B deductible. The limitation reduces the amount of incurred expenses to 62 1/2 percent of the approved amount. After subtracting any unmet deductible, Medicare pays 80 percent of the remaining incurred expenses. Medicare payment and beneficiary liability are computed as follows:

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<th>Description</th>
<th>Amount</th>
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<td>2. Medicare approved amount</td>
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<td>3. Medicare incurred expenses (0.625 x line 2)</td>
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<td>4. Unmet deductible</td>
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<td>5. Remainder after subtracting deductible (line 3 minus line 4)</td>
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<td>6. Medicare payment (0.80 x line 5)</td>
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<td>7. Beneficiary liability (line 2 minus line 6)</td>
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(2) A clinical social worker submitted a claim for $135 for outpatient treatment of a beneficiary’s mental disorder. The Medicare approved amount was $120. Since clinical social workers must accept assignment, the beneficiary is not liable for the $15 in excess charges. The beneficiary previously satisfied $70 of the $100 annual Part B deductible, leaving $30 unmet.

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<td>4. Unmet deductible</td>
<td>30.00</td>
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§ 410.160 Part B annual deductible.

(a) Basic rule. Except as provided in paragraph (b) of this section, incurred expenses (as defined in §410.152) are subject to, and count toward meeting the annual deductible.

(b) Exceptions. Expenses incurred for the following services are not subject to the Part B annual deductible and do not count toward meeting that deductible:

(1) Home health services.
(2) Pneumococcal vaccines and their administration.
(3) Federally qualified health center services.

(4) A beneficiary's only Part B expenses during 1995 were for a physician's services in connection with the treatment of a mental disorder that initially required inpatient hospitalization. The remaining services were furnished on an outpatient basis. The beneficiary had not satisfied any of the $100 Part B annual deductible.

(4) ASC facility services furnished before July 1987 and physician services furnished before April 1988 that met the requirements for payment of 100 percent of the reasonable charges.

(c) Application of the Part B annual deductible. (1) Before payment is made under §410.152, an individual's incurred expenses for the calendar year are reduced by the Part B annual deductible.

(2) The Part B annual deductible is applied to incurred expenses in the order in which claims for those expenses are processed by the Medicare program.

(3) Only one Part B annual deductible may be imposed for any calendar year and it may be met by any combination of expenses incurred in that year.

(d) Special rule for services reimbursable on a formula basis. (1) In applying the formula that takes into account reasonable costs, customary charges, and customary (insofar as reasonable) charges, and is used to determine payment for services furnished by a provider that is not a nominal charge provider, the Medicare intermediary takes the following steps:

(i) Reduces the customary charges for the services by an amount equal to any unmet portion of the deductible for the calendar year, in accordance with paragraph (b) of this section. (The amount of this reduction is considered to be the amount of the deductible that
§ 410.161 Part B blood deductible.

(a) General rules. (1) As used in this section, packed red cells means the red blood cells that remain after plasma is separated from whole blood.

(ii) A unit of packed red cells is treated as the equivalent of a pint of whole blood, which in this section is referred to as a unit of whole blood.

(3) Medicare does not pay for the first 3 units of whole blood or units of packed red cells that are furnished under Part A or Part B in a calendar year. The Part B blood deductible is reduced to the extent that a blood deductible has been applied under Part A.

(4) The blood deductible does not apply to other blood components such as plasma.
as platelets, fibrinogen, plasma, gamma globulin and serum albumin, or to the costs of processing, storing, and administering blood.

(5) The blood deductible is in addition to the Part B annual deductible specified in §410.160.

(b) Beneficiary's responsibility for the first 3 units of blood.

(1) The beneficiary is responsible for the first three units of whole blood or packed red cells received during a calendar year.

(2) If the blood is furnished by a hospital or CAH, the rules set forth in §409.87 (b), (c), and (d) of this chapter apply.

(3) If the blood is furnished by a physician, clinic, or other supplier that has accepted assignment of Medicare benefits, or claims payment under §424.64 of this chapter because the beneficiary died without assigning benefits, the supplier may charge the beneficiary the reasonable charge for the first 3 units, to the extent that those units are not replaced.

§410.163 Payment for services furnished to kidney donors.

Notwithstanding any other provisions of this chapter, there are no deductible or coinsurance requirements with respect to services furnished to an individual who donates a kidney for transplant surgery.

§410.165 Payment for rural health clinic services and ambulatory surgical center services: Conditions.

(a) Medicare Part B pays for covered rural health clinic and Federally qualified health center services if—

(1) The services are furnished in accordance with the requirements of subpart X of part 405 of this chapter and subpart A of part 491 of this chapter; and

(2) The clinic or center files a written request for payment on the form and in the manner prescribed by HCFA.

(b) Medicare Part B pays for covered ambulatory surgical center (ASC) services if—

(1) The services are furnished in accordance with the requirements of part 416 of this chapter; and

(2) The ASC files a written request for payment on the form and in the manner prescribed by HCFA.

§410.172 Payment for partial hospitalization services in CMHCs: Conditions.

Medicare Part B pays for partial hospitalization services in a CMHC on behalf of an individual only if the following conditions are met:

(a) The CMHC files a written request for payment on the HCFA form 1450 and in the manner prescribed by HCFA; and
§ 410.175

(b) The services are furnished in accordance with the requirements described in §410.110.

[59 FR 6578, Feb. 11, 1994]

§ 410.175 Alien absent from the United States.

(a) Medicare does not pay Part B benefits for services furnished to an individual who is not a citizen or a national of the United States if those services are furnished in any month for which the individual is not paid monthly social security cash benefits (or would not be paid if he or she were entitled to those benefits) because he or she has been outside the United States continuously for 6 full calendar months.

(b) Payment of benefits resumes with services furnished during the first full calendar month the alien is back in the United States.

[53 FR 6634, Mar. 2, 1988]

PART 411—EXCLUSIONS FROM MEDICARE AND LIMITATIONS ON MEDICARE PAYMENT

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Authority: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh).

Source: 54 FR 41734, Oct. 11, 1989, unless otherwise noted.

Editorial Note: Nomenclature changes affecting part 411 appear at 60 FR 45370, Aug. 31, 1995.
Subpart A—General Exclusions and Exclusion of Particular Services

§ 411.1 Basis and scope.

(a) Statutory basis. Sections 1814(c), 1835(d), and 1862 of the Act exclude from Medicare payment certain specified services. The Act provides special rules for payment of services furnished by Federal providers or agencies (sections 1814(c) and 1835(d)), by hospitals and physicians outside the United States (sections 1814(f) and 1862(a)(4)), and by hospitals and SNFs of the Indian Health Service (section 1880). Section 1877 sets forth limitations on referrals and payment for clinical laboratory services furnished by entities with which the referring physician (or an immediate family member of the referring physician) has a financial relationship. Sections 1842(l) and 1879 of the Act provide for refund to, or indemnification of, a beneficiary who has paid a provider or supplier for certain services that the provider or supplier knew were excluded from Medicare coverage.

(b) Scope. This subpart identifies:

1. The particular types of services that are excluded;
2. The circumstances under which Medicare denies payment for certain services that are usually covered; and
3. The circumstances under which Medicare pays for services usually excluded from payment.

§ 411.2 Conclusive effect of PRO determinations on payment of claims.

If a utilization and quality control peer review organization (PRO) has assumed review responsibility, in accordance with part 466 of this chapter, for services furnished to Medicare beneficiaries, Medicare payment is not made for those services unless the conditions of subpart C of part 466 of this chapter are met.

§ 411.4 Services for which neither the beneficiary nor any other person is legally obligated to pay.

(a) General rule. Except as provided in §411.8(b) (for services paid by a governmental entity), Medicare does not pay for a service if—

1. The beneficiary has no legal obligation to pay for the service; and
2. No other person or organization (such as a prepayment plan of which the beneficiary is a member) has a legal obligation to provide or pay for that service.

(b) Special conditions for services furnished to individuals in custody of penal authorities. Payment may be made for services furnished to individuals or groups of individuals who are in the custody of the police or other penal authorities or in the custody of a government agency under a penal statute only if the following conditions are met:

1. State or local law requires those individuals or groups of individuals to repay the cost of medical services they receive while in custody.
2. The State or local government entity enforces the requirement to pay by billing all such individuals, whether or not covered by Medicare or any other health insurance, and by pursuing collection of the amounts they owe in the same way and with the same vigor that it pursues the collection of other debts.

§ 411.6 Services furnished by a Federal provider of services or other Federal agency.

(a) Basic rule. Except as provided in paragraph (b) of this section, Medicare does not pay for services furnished by a Federal provider of services or other Federal agency.

(b) Exceptions. Payment may be made—

1. For emergency hospital services, if the conditions of §424.103 of this chapter are met;
2. For services furnished by a participating Federal provider which HCFA has determined is providing services to the public generally as a community institution or agency;
3. For services furnished by participating hospitals and SNFs of the Indian Health Service; and
4. For services furnished under arrangements (as defined in §409.3 of this chapter) made by a participating hospital.

§ 411.7 Services that must be furnished at public expense under a Federal law or Federal Government contract.

(a) Basic rule. Except as provided in paragraph (b) of this section, payment may not be made for services that any provider or supplier is obligated to furnish at public expense, in accordance with a law of, or a contract with, the United States.

(b) Exception. Payment may be made for services that a hospital or SNF of the Indian Health Service is obligated to furnish at public expense.

§ 411.8 Services paid for by a Government entity.

(a) Basic rule. Except as provided in paragraph (b) of this section, Medicare does not pay for services that are paid for directly or indirectly by a government entity.

(b) Exceptions. Payment may be made for the following:

(1) Services furnished under a health insurance plan established for employees of the government entity.

(2) Services furnished under a title of the Social Security Act other than title XVIII.

(3) Services furnished in or by a participating general or special hospital that—

(i) Is operated by a State or local government agency; and

(ii) Serves the general community.

(4) Services furnished in a hospital or elsewhere, as a means of controlling infectious diseases or because the individual is medically indigent.

(5) Services furnished by a participating hospital or SNF of the Indian Health Service.

(b) Services furnished by a public or private health facility that—

(i) Is not a Federal provider or other facility operated by a Federal agency;

(ii) Receives U.S. government funds under a Federal program that provides support to facilities that furnish health care services;

(iii) Customarily seeks payment for services not covered under Medicare from all available sources, including private insurance and patients' cash resources; and

(iv) Limits the amounts it collects or seeks to collect from a Medicare Part B beneficiary and others on the beneficiary's behalf to:

(A) Any unmet deductible applied to the charges related to the reasonable costs that the facility incurs in providing the covered services;

(B) Twenty percent of the remainder of those charges;

(C) The charges for noncovered services.

(7) Rural health clinic services that meet the requirements set forth in part 491 of this chapter.


§ 411.9 Services furnished outside the United States.

(a) Basic rule. Except as specified in paragraph (b) of this section, Medicare does not pay for services furnished outside the United States. For purposes of this paragraph (a), the following rules apply:

(1) The United States includes the 50 States, the District of Columbia, Puerto Rico, the Virgin Islands, Guam, American Samoa, The Northern Mariana Islands, and for purposes of services rendered on board ship, the territorial waters adjoining the land areas of the United States.

(2) Services furnished on board ship are considered to have been furnished in United States territorial waters if they were furnished while the ship was in a port of one of the jurisdictions listed in paragraph (a)(1) of this section, or within 6 hours before arrival at, or 6 hours after departure from, such a port.

(3) A hospital that is not physically situated in one of the jurisdictions listed in paragraph (a)(1) of this section is considered to be outside the United States, even if it is owned or operated by the United States Government.

(b) Exception. Under the circumstances specified in subpart H of part 424 of this chapter, payment may be made for covered inpatient services furnished in a foreign hospital and, on the basis of an itemized bill, for covered physicians' services and ambulance service furnished in connection with those inpatient services, but only for the period during which the inpatient hospital services are furnished.
§ 411.10 Services required as a result of war.

Medicare does not pay for services that are required as a result of war, or an act of war, that occurs after the effective date of a beneficiary's current coverage for hospital insurance benefits or supplementary medical insurance benefits.

§ 411.12 Charges imposed by an immediate relative or member of the beneficiary's household.

(a) Basic rule. Medicare does not pay for services usually covered under Medicare if the charges for those services are imposed by—
(1) An immediate relative of the beneficiary; or
(2) A member of the beneficiary's household.

(b) Definitions. As used in this section—

Immediate relative means any of the following:
(1) Husband or wife.
(2) Natural or adoptive parent, child, or sibling.
(3) Stepparent, stepchild, stepbrother, or stepsister.
(5) Grandparent or grandchild.
(6) Spouse of grandparent or grandchild.

Member of the household means any person sharing a common abode as part of a single family unit, including domestic employees and others who live together as part of a family unit, but not including a mere roomer or boarder.

Professional corporation means a corporation that is completely owned by one or more physicians and is operated for the purpose of conducting the practice of medicine, osteopathy, dentistry, podiatry, optometry, or chiropractic, or is owned by other health care professionals as authorized by State law.

(c) Applicability of the exclusion. The exclusion applies to the following charges in the specified circumstances:

(i) Charges for physicians' services furnished by an immediate relative of the beneficiary or member of the beneficiary's household, even if the bill or claim is submitted by another individual or by an entity such as a partnership or a professional corporation.

(ii) Charges for services furnished incident to a physician's professional services (for example by the physician's nurse or technician), only if the physician who ordered or supervised the services has an excluded relationship to the beneficiary.

(2) Services other than physicians' services. (i) Charges imposed by an individually owned provider or supplier if the owner has an excluded relationship to the beneficiary; and

(ii) Charges imposed by a partnership if any of the partners has an excluded relationship to the beneficiary.

(d) Exception to the exclusion. The exclusion does not apply to charges imposed by a corporation other than a professional corporation.

§ 411.15 Particular services excluded from coverage.

The following services are excluded from coverage:

(a) Routine physical checkups such as:

(1) Examinations performed for a purpose other than treatment or diagnosis of a specific illness, symptoms, complaint, or injury, except for screening mammography, colorectal cancer screening tests, screening pelvic examinations, or prostate cancer screening tests that meet the criteria specified in paragraphs (k)(6) through (k)(9) of this section.

(2) Examinations required by insurance companies, business establishments, government agencies, or other third parties.

(b) Eyeglasses or contact lenses, except for:

(1) Post-surgical prosthetic lenses customarily used during convalescence for eye surgery in which the lens of the eye was removed (e.g., cataract surgery);

(2) Prosthetic lenses for patients who lack the lens of the eye because of congenital absence or surgical removal; and

(3) One pair of conventional eyeglasses or conventional contact lenses furnished after each cataract surgery during which an intraocular lens is inserted.
Before July 1981, inpatient hospital care in connection with dental procedures was covered only when required by the patient’s underlying medical condition and clinical status.
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the conditions and limitations specified in §410.39 of this chapter.

(l) Foot care. (1) Basic rule. Except as provided in paragraph (l)(2) of this section, any services furnished in connection with the following:

(i) Routine foot care, such as the cutting or removal of corns, or calluses, the trimming of nails, routine hygiene care (preventive maintenance care ordinarily within the realm of self care), and any service performed in the absence of localized illness, injury, or symptoms involving the feet.

(ii) The evaluation or treatment of subluxations of the feet regardless of underlying pathology. (Subluxations are structural misalignments of the joints, other than fractures or complete dislocations, that require treatment only by nonsurgical methods.

(iii) The evaluation or treatment of flattened arches (including the prescription of supportive devices) regardless of the underlying pathology.

(2) Exceptions. (i) Treatment of warts is not excluded.

(ii) Treatment of mycotic toenails may be covered if it is furnished no more often than every 60 days or the billing physician documents the need for more frequent treatment.

(iii) The services listed in paragraph (l)(1) of this section are not excluded if they are furnished—

(A) As an incident to, at the same time as, or as a necessary integral part of a primary covered procedure performed on the foot; or

(B) As initial diagnostic services (regardless of the resulting diagnosis) in connection with a specific symptom or complaint that might arise from a condition whose treatment would be covered.

(m) Services to hospital patients—(1) Basic rule. Except as provided in paragraph (m)(3) of this section, any service furnished to an inpatient of a hospital or to a hospital outpatient (as defined in §410.2 of this chapter) during an encounter (as defined in §410.2 of this chapter) by an entity other than the hospital unless the hospital has an arrangement (as defined in §409.3 of this chapter) with that entity to furnish that particular service to the hospital’s patients. As used in this paragraph (m)(1), the term “hospital” includes a CAH.

(2) Scope of exclusion. Services subject to exclusion from coverage under the provisions of this paragraph (m) include, but are not limited to, clinical laboratory services; pacemakers and other prostheses and prosthetic devices (other than dental) that replace all or part of an internal body organ (for example, intraocular lenses); artificial limbs, knees, and hips; equipment and supplies covered under the prosthetic device benefits; and services incident to a physician service.

(3) Exceptions. The following services are not excluded from coverage:

(i) Physicians’ services that meet the criteria of §415.102(a) of this chapter for payment on a reasonable charge or fee schedule basis.

(ii) Physician assistant services, as defined in section 1861(s)(2)(K)(i) of the Act, that are furnished after December 31, 1990.

(iii) Nurse practitioner and clinical nurse specialist services, as defined in section 1861(s)(2)(K)(ii) of the Act.

(iv) Certified nurse-midwife services, as defined in section 1861(ff) of the Act, that are furnished after December 31, 1990.

(v) Qualified psychologist services, as defined in section 1861(iii) of the Act, that are furnished after December 31, 1990.

(vi) Services of an anesthetist, as defined in §410.69 of this chapter.

(n) Certain services of an assistant-at-surgery.

(1) Services of an assistant-at-surgery in a cataract operation (including subsequent insertion of an intraocular lens) unless, before the surgery is performed, the appropriate PRO or a carrier has approved the use of such an assistant in the surgical procedure based on the existence of a complicating medical condition.

(2) Services on an assistant-at-surgery in a surgical procedure (or class of surgical procedures) for which assistants-at-surgery on average are used in fewer than 5 percent of such procedures nationally.

(o) Experimental or investigational devices, except for certain devices—

(1) Categorized by the FDA as a non-experimental/Investigational (Category
B) device defined in §405.201(b) of this chapter; and
(2) Furnished in accordance with the FDA-approved protocols governing clinical trials.

(p) Services furnished to SNF residents—(1) Basic rule. Except as provided in paragraph (p)(2) of this section, any service furnished to a resident of an SNF by an entity other than the SNF, unless the SNF has an arrangement (as defined in §409.3 of this chapter) with that entity to furnish that particular service to the SNF’s residents. Services subject to exclusion under this paragraph include, but are not limited to—
(i) Any physical, occupational, or speech-language therapy services regardless of whether or not the services are furnished by, or under the supervision of, a physician or other health care professional; and
(ii) Services furnished as an incident to the professional services of a physician or other health care professional specified in paragraph (p)(2) of this section.
(2) Exceptions. The following services are not excluded from coverage:
(i) Physicians’ services that meet the criteria of §415.102(a) of this chapter for payment on a fee schedule basis, provided that the claim for payment includes the SNF’s Medicare provider number in accordance with §424.32(a)(2) of this chapter.
(ii) Services performed under a physician’s supervision by a physician assistant who meets the applicable definition in section 1861(aa)(5) of the Act and is working in collaboration (as defined in section 1861(aa)(6) of the Act) with a physician.
(iii) Services performed by a nurse practitioner or clinical nurse specialist who meets the applicable definition in section 1861(aa)(5) of the Act and is working in collaboration (as defined in section 1861(aa)(6) of the Act) with a physician.
(iv) Services performed by a certified nurse-midwife, as defined in section 1861(gg) of the Act.
(v) Services performed by a qualified psychologist, as defined in section 1861(ii) of the Act.
(vi) Services performed by a certified registered nurse anesthetist, as defined in section 1861(bb) of the Act.
(vii) Dialysis services and supplies, as defined in section 1861(s)(2)(F) of the Act, and those ambulance services that are furnished in conjunction with them.
(viii) Erythropoietin (EPO) for dialysis patients, as defined in section 1861(s)(2)(I) of the Act.
(ix) Hospice care, as defined in section 1861(dd) of the Act.
(x) An ambulance trip that initially conveys an individual to the SNF to be admitted as a resident, or that conveys an individual from the SNF in connection with one of the circumstances specified in paragraphs (p)(3)(i) through (p)(3)(iv) of this section as ending the individual’s status as an SNF resident.
(xi) The transportation costs of electrocardiogram equipment (HCPCS code R0076), but only with respect to those electrocardiogram test services furnished during 1998.
(xii) Those chemotherapy items identified, as of July 1, 1999, by HCPCS codes J9000-J9020; J9040-J9151; J9170-J9185; J9200-J9208; J9211; J9230-J9245; and J9265-J9600.
(xiii) Those chemotherapy administration services identified, as of July 1, 1999, by HCPCS codes 36260-36262; 36489; 36530-36535; 36640; 36823; and 96405-96542.
(xiv) Those radioisotope services identified, as of July 1, 1999, by HCPCS codes 79030-79440.
(xv) Those customized prosthetic devices (including artificial limbs and their components) identified, as of July 1, 1999, by HCPCS codes L5050-L5343; L5500-L5611; L5613-L5986; L5988; L6050-L6370; L6400-6880; L6920-L7274; and L7362-L7366, which are delivered for a resident’s use during a stay in the SNF and intended to be used by the resident after discharge from the SNF.
(3) SNF resident defined. For purposes of this paragraph, a beneficiary who is admitted to a Medicare-participating SNF (or to the nonparticipating portion of a nursing home of which a distinct part is a Medicare-participating SNF) is considered to be a resident of the SNF, regardless of whether Part A covers the stay. Whenever such a beneficiary leaves the facility, the beneficiary’s status as an SNF resident for purposes of this paragraph (along with the SNF’s responsibility to furnish or
§411.20 Basis and scope.

(a) Statutory basis. (1) Section 1862(b)(2)(A)(i) of the Act precludes Medicare payment for services to the extent that payment has been made or can reasonably be expected to be made promptly under any of the following:

(i) Workers' compensation.

(ii) Liability insurance.

(iii) No-fault insurance.

(b) Scope. This subpart sets forth general rules that apply to the types of insurance specified in paragraph (a) of this section. Other general rules that apply to group health plans are set forth in subpart E of this part.

[60 FR 45361, Aug. 31, 1995]
or assume legal liability for injury or illness.

Prompt or promptly, when used in connection with third party payments, except as provided in §411.50, for payments by liability insurers, means payment within 120 days after receipt of the claim.

Proper claim means a claim that is filed timely and meets all other claim filing requirements specified by the plan, program, or insurer.

Secondary, when used to characterize Medicare benefits, means that those benefits are payable only to the extent that payment has not been made and cannot reasonably be expected to be made under other coverage that is primary to Medicare.

Secondary payments means payments made for Medicare covered services or portions of services that are not payable under other coverage that is primary to Medicare.

Third party payer means an insurance policy, plan, or program that is primary to Medicare.

Third party payment means payment by a third party payer for services that are also covered under Medicare.

§ 411.24 Recovery of conditional payments.

If a Medicare conditional payment is made, the following rules apply:

(a) Release of information. The filing of a Medicare claim by on or behalf of the beneficiary constitutes an express authorization for any entity, including State Medicaid and workers' compensation agencies, and data depositaries, that possesses information pertinent to the Medicare claim to release that information to HCFA. This information will be used only for Medicare claims processing and for coordination of benefits purposes. (b) Right to initiate recovery. HCFA may initiate recovery as soon as it learns that payment has been made or could be made under workers' compensation, any liability or no-fault insurance, or an employer group health plan.

(c) Amount of recovery. (1) If it is not necessary for HCFA to take legal action to recover, HCFA recovers the lesser of the following:

(i) The amount of the Medicare primary payment.

(ii) The full primary payment amount that the primary payer is obligated to pay under this part without regard to any payment, other than a full primary payment that the primary payer has paid or will make, or, in the case of a third party payment recipient, the amount of the third party payment.

(2) If it is necessary for HCFA to take legal action to recover from the primary payer, HCFA may recover twice the amount specified in paragraph (c)(1)(i) of this section.

(d) Methods of recovery. HCFA may recover by direct collection or by offset against any monies HCFA owes the entity responsible for refunding the conditional payment.

(e) Recovery from third parties. HCFA has a direct right of action to recover from any entity responsible for making primary payment. This includes an employer, an insurance carrier, plan, or program, and a third party administrator.

(f) Claims filing requirements. (1) HCFA may recover without regard to any claims filing requirements that the insurance program or plan imposes on the beneficiary or other claimant such as a time limit for filing a claim or a time limit for notifying the plan or program about the need for or receipt of services.

(2) However, HCFA will not recover its payment for particular services in the face of a claims filing requirement unless it has filed a claim for recovery by the end of the year following the year in which the Medicare intermediary or carrier that paid the claim has notice that the third party is primary to Medicare for those particular services. (A notice received during the
§ 411.24

last three months of a year is consid-
ered received during the following
year.)

(g) Recovery from parties that receive
third party payments. HCFA has a right
of action to recover its payments from
any entity, including a beneficiary,
provider, supplier, physician, attorney,
State agency or private insurer that
has received a third party payment.

(h) Reimbursement to Medicare. If the
beneficiary or other party receives a
third party payment, the beneficiary or
other party must reimburse Medicare
within 60 days.

(i) Special rules. (1) In the case of li-
ability insurance settlements and dis-
puted claims under employer group
health plans and no-fault insurance,
the following rule applies: If Medicare
is not reimbursed as required by para-
graph (h) of this section, the third
party payer must reimburse Medicare
even though it has already reimbursed
the beneficiary or other party.

(2) The provisions of paragraph (i)(1)
of this section also apply if a third
party payer makes its payment to an
entity other than Medicare when it is,
or should be, aware that Medicare has
made a conditional primary payment.

(3) In situations that involve procure-
ment costs, the rule of § 411.37(b) ap-
plies.

(j) Recovery against Medicaid agency.
If a third party payment is made to a
State Medicaid agency and that agency
does not reimburse Medicare, HCFA
may reduce any Federal funds due the
Medicaid agency (under title XIX of
the Act) by an amount equal to the
Medicare payment or the third party
payment, whichever is less.

(k) Recovery against Medicare con-
tactor. If a Medicare contractor, in-
cluding an intermediary or carrier,
also insure, underwrites, or admin-
isters as a third party administrator, a
program or plan that is primary to
Medicare, and does not reimburse Me-
dicare, HCFA may offset the Federal funds due the
intermediary or carrier under title
XVIII of the Act or due the contractor
under the contract.

(l) Recovery when there is failure to file
a proper claim. (1) Basic rule. If Medicare
makes a conditional payment with re-
spect to services for which the bene-

ficiary or provider or supplier has not
filed a proper claim with a third party
payer, and Medicare is unable to re-
cover from the third party payer, Medi-
care may recover from the beneficiary
or provider or supplier that was respon-
sible for the failure to file a proper
claim.

(2) Exceptions: (i) This rule does not
apply in the case of liability insurance
nor when failure to file a proper claim
is due to mental or physical incapacity
of the beneficiary.

(ii) HCFA will not recover from pro-
viders or suppliers that are in compli-
ance with the requirements of § 489.20
of this chapter and can show that
the reason they failed to file a proper
claim is that the beneficiary, or some-
one acting on his or her behalf, failed
to give, or gave erroneous, information
regarding coverage that is primary to
Medicare.

(m) Interest charges. (1) With respect
to recovery of payments for items and
services furnished before October 31,
1994, HCFA charges interest, exercising
common law authority in accordance
with 45 CFR 30.13, consistent with the
Federal Claims Collection Act (31

(2) In addition to its common law au-
thority with respect to recovery of
payments for items and services fur-
nished on or after October 31, 1994,
HCFA charges interest in accordance
with section 1862(b)(2)(B)(i) of the Act.
Under that provision—

(i) HCFA may charge interest if reim-
bursement is not made to the appro-
priate trust fund before the expiration
of the 60-day period that begins on the
date on which notice or other informa-
tion is received by HCFA that payment
has been or could be made under a pri-
mary plan;

(ii) Interest may accrue from the
date when that notice or other infor-
mation is received by HCFA and is
charged until reimbursement is made;
and

(iii) The rate of interest is that pro-
vided at 42 CFR 405.376(d).

[54 FR 41794, Oct. 11, 1989, as amended at 55
31, 1995]
§ 411.25 Third party payer's notice of mistaken Medicare primary payment.

(a) If a third party payer learns that HCFA has made a Medicare primary payment for services for which the third party payer has made or should have made primary payment, it must give notice to that effect to the Medicare intermediary or carrier that paid the claim.

(b) The notice must describe the specific situation and the circumstances (including the particular type of insurance coverage as specified in §411.20(a)) and, if appropriate, the time period during which the insurer is primary to Medicare.

(c) If a plan is self-insured and self-administered, the employer must give the notice to HCFA. Otherwise, the insurer, underwriter, or third party administrator must give the notice.


§ 411.26 Subrogation and right to intervene.

(a) Subrogation. With respect to services for which Medicare paid, HCFA is subrogated to any individual, provider, supplier, physician, private insurer, State agency, attorney, or any other entity entitled to payment by a third party payer.

(b) Right to intervene. HCFA may join or intervene in any action related to the events that gave rise to the need for services for which Medicare paid.


§ 411.28 Waiver of recovery and compromise of claims.

(a) HCFA may waive recovery, in whole or in part, if the probability of recovery, or the amount involved, does not warrant pursuit of the claim.

(b) General rules applicable to compromise of claims are set forth in subpart F of part 401 and §405.376 of this chapter.

(c) Other rules pertinent to recovery are contained in subpart C of part 405 of this chapter.


§ 411.30 Effect of third party payment on benefit utilization and deductibles.

(a) Benefit utilization. Inpatient psychiatric hospital and SNF care that is paid for by a third party payer is not counted against the number of inpatient care days available to the beneficiary under Medicare Part A.

(b) Deductibles. Expenses for Medicare covered services that are paid for by third party payers are credited toward the Medicare Part A and Part B deductibles.

§ 411.31 Authority to bill third party payers for full charges.

(a) The fact that Medicare payments are limited to the DRG amount, or the reasonable charge, reasonable cost, capitation or fee schedule rate, does not affect the amount that a third party payer may pay.

(b) With respect to workers' compensation plans, no-fault insurers, and employer group health plans, a provider or supplier may bill its full charges and expect those charges to be paid unless there are limits imposed by laws other than title XVIII of the Act or by agreements with the third party payer.

§ 411.32 Basis for Medicare secondary payments.

(a) Basic rules. (1) Medicare benefits are secondary to benefits payable by a third party payer even if State law or the third party payer states that its benefits are secondary to Medicare benefits or otherwise limits its payments to Medicare beneficiaries.

(2) Except as provided in paragraph (b) of this section, Medicare makes secondary payments, within the limits specified in paragraph (c) of this section and in §411.33, to supplement the third party payment if that payment is less than the charges for the services and, in the case of services paid on other than a reasonable charge basis, less than the gross amount payable by Medicare under §411.33(e).

(b) Exception. Medicare does not make a secondary payment if the provider or supplier is either obligated to accept, or voluntarily accepts, as full payment, a third party payment that is less than its charges.
§ 411.33 Amount of Medicare secondary payment.

(a) Services for which HCFA pays on a Medicare fee schedule or reasonable charge basis. The Medicare secondary payment is the lowest of the following:

(1) The actual charge by the supplier (or the amount the supplier is obligated to accept as payment in full if that is less than the charges) minus the amount paid by the third party payer.

(2) The amount that Medicare would pay if the services were not covered by a third party payer.

(3) The higher of the Medicare fee schedule, Medicare reasonable charge, or other amount which would be payable under Medicare (without regard to any applicable Medicare deductible or coinsurance amounts) or the third party payer’s allowable charge (without regard to any deductible or co-insurance imposed by the policy or plan) minus the amount actually paid by the third party payer.

(b) Example: An individual received treatment from a physician for which the physician charged $175. The third party payer allowed $150 of the charge and paid 80 percent of this amount or $120. The Medicare fee schedule for this treatment is $125. The individual’s Part B deductible had been met. As secondary payer, Medicare pays the lowest of the following amounts:

(1) Excess of actual charge minus the third party payment: $175 – $120 = $55.

(2) Amount Medicare would pay if the services were not covered by a third party payer: .80 × $125 = $100.

(3) The third party payer’s allowable charge without regard to its coinsurance (since that amount is higher than the Medicare fee schedule in this case) minus amount paid by the third party payer: $150 – $120 = $30.

The Medicare payment is $30.

(c)-(d) [Reserved]

(e) Services reimbursed on a basis other than fee schedule, reasonable charge, or monthly capitation rate. The Medicare secondary payment is the lowest of the following:

(1) The gross amount payable by Medicare (that is, the amount payable without considering the effect of the Medicare deductible and coinsurance or the payment by the third party payer), minus the applicable Medicare deductible and coinsurance amounts.

(2) The gross amount payable by Medicare, minus the amount paid by the third party payer.

(3) The provider’s charges (or the amount the provider is obligated to accept as payment in full if that is less than the charges), minus the amount payable by the third party payer.

(4) The provider’s charges (or the amount the provider is obligated to accept as payment in full if that is less than the charges), minus the applicable Medicare deductible and coinsurance amounts.

(f) Examples: (1) A hospital furnished 7 days of inpatient hospital care in 1987 to a Medicare beneficiary. The provider’s charges for Medicare-covered services totaled $2,000. The third party payer paid $2,360. No part of the Medicare inpatient hospital deductible of $520 had been met. If the gross amount payable by Medicare in this case is $2,700, then as secondary payer, Medicare pays the lowest of the following amounts:

(i) The gross amount payable by Medicare minus the Medicare inpatient hospital deductible: $2,700 – $520 = $2,180.

(ii) The gross amount payable by Medicare minus the third party payment: $2,700 – $2,360 = $340.

(iii) The provider’s charges minus the third party payment: $2,800 – $2,360 = $440.

(iv) The provider’s charges minus the Medicare deductible: $2,800 – $520 = $2,280.

Medicare’s secondary payment is $340 and the combined payment made by
Health Care Financing Administration, HHS

§ 411.35 Limitations on charges to a beneficiary or other party when a workers’ compensation plan, a no-fault insurer, or an employer group health plan is primary payer.

(a) Definition. As used in this section Medicare-covered services means services for which Medicare benefits are payable or would be payable except for the Medicare deductible and coinsurance.

(iv) The provider’s charge minus the Medicare deductible and coinsurance: $1,280 – $75 – $194.60 = $990.40. Medicare pays $24. The beneficiary’s Medicare deductible and coinsurance were met by the third party payment.

(4) A hospital furnished 5 days of inpatient care in 1987 to a Medicare beneficiary. The provider’s charges for Medicare-covered services were $4,000 and the gross amount payable was $3,500. The provider agreed to accept $3,000 from the third party as payment in full. The third party payer paid $2,900 due to a deductible requirement under the third party plan. Medicare considers the amount the provider is obligated to accept as full payment ($3,000) to be the provider charges. The Medicare secondary payment is the lowest of the following:

(i) The gross amount payable by Medicare minus the applicable Medicare inpatient deductible: $3,000 – $520 = $2,480. Medicare pays $24. The beneficiary’s Medicare deductible and coinsurance were met by the third party payment.

(ii) The provider’s charges minus the Medicare deductible and coinsurance: $3,000 – $75 – $194.60 = $2,930.40. Medicare pays $24. The beneficiary’s Medicare deductible and coinsurance were met by the third party payment.

(iii) The provider’s charges minus the applicable Medicare inpatient deductible: $3,000 – $520 = $2,480. Medicare pays $24. The beneficiary’s Medicare deductible and coinsurance were met by the third party payment.

(i) The gross amount payable by Medicare minus the third party payment: $1,280 – $1,024 = $256.

(ii) The provider’s charges minus the Medicare deductible and coinsurance: $1,280 – $75 – $194.60 = $1010.40. Medicare pays $24. The beneficiary’s Medicare deductible and coinsurance were met by the third party payment.

(iii) The provider’s charges minus the third party payment: $1,280 – $1,024 = $256.

(iv) The provider’s charge minus the Medicare deductible and coinsurance: $1,280 – $75 – $194.60 = $1010.40. Medicare pays $24. The beneficiary’s Medicare deductible and coinsurance were met by the third party payment.

(i) The gross amount payable by Medicare minus the applicable Medicare inpatient deductible: $3,000 – $520 = $2,480. Medicare pays $24. The beneficiary’s Medicare deductible and coinsurance were met by the third party payment.

(ii) The provider’s charges minus the Medicare deductible and coinsurance: $3,000 – $75 – $194.60 = $2,930.40. Medicare pays $24. The beneficiary’s Medicare deductible and coinsurance were met by the third party payment.

(iii) The provider’s charges minus the applicable Medicare inpatient deductible: $3,000 – $520 = $2,480. Medicare pays $24. The beneficiary’s Medicare deductible and coinsurance were met by the third party payment.

(i) The gross amount payable by Medicare minus the third party payment: $1,280 – $1,024 = $256.

(ii) The provider’s charges minus the Medicare deductible and coinsurance: $1,280 – $75 – $194.60 = $1010.40. Medicare pays $24. The beneficiary’s Medicare deductible and coinsurance were met by the third party payment.

(iii) The provider’s charges minus the third party payment: $1,280 – $1,024 = $256.
§ 411.37 Amount of Medicare recovery when a third party payment is made as a result of a judgment or settlement.

(a) Recovery against the party that received payment—(1) General rule. Medicare reduces its recovery to take account of the cost of procuring the judgment or settlement, as provided in this section, if—

(i) Procurement costs are incurred because the claim is disputed; and

(ii) Those costs are borne by the party against which HCFA seeks to recover.

(2) Special rule. If HCFA must file suit because the party that received payment opposes HCFA's recovery, the recovery amount is as set forth in paragraph (e) of this section.

(b) Recovery against the third party payer. If HCFA seeks recovery from the third party payer, in accordance with §411.24(i), the recovery amount will be no greater than the amount determined under paragraph (c) or (d) or (e) of this section.

(c) Medicare payments are less than the judgment or settlement amount. If Medicare payments are less than the judgment or settlement amount, the recovery is computed as follows:

(1) Determine the ratio of the procurement costs to the total judgment or settlement payment.

(2) Apply the ratio to the Medicare payment. The product is the Medicare share of procurement costs.

(3) Subtract the Medicare share of procurement costs from the Medicare payments. The remainder is the Medicare recovery amount.

(d) Medicare payments equal or exceed the judgment or settlement amount. If Medicare payments equal or exceed the judgment or settlement amount, the recovery amount is the total judgment or settlement payment minus the total procurement costs.

(e) HCFA incurs procurement costs because of opposition to its recovery. If HCFA must bring suit against the party that received payment because that party opposes HCFA's recovery, the recovery amount is the lower of the following:

(1) Medicare payment.

(2) The total judgment or settlement amount, minus the party's total procurement cost.

Subpart C—Limitations on Medicare Payment for Services Covered Under Workers' Compensation

§ 411.40 General provisions.

(a) Definition. "Workers' compensation plan of the United States" includes the workers' compensation plans of the 50
Health Care Financing Administration, HHS

§ 411.46 Lump-sum payments.

(a) Lump-sum commutation of future benefits. If a lump-sum compensation award stipulates that the amount paid is intended to compensate the individual for all future medical expenses required because of the work-related injury or disease, Medicare payments for such services are excluded until medical expenses related to the injury or disease equal the amount of the lump-sum payment.

(b) Lump-sum compromise settlement. (1) A lump-sum compromise settlement is deemed to be a workers' compensation payment for Medicare purposes, even if the settlement agreement stipulates that there is no liability under the workers' compensation law or plan. (2) If a settlement appears to represent an attempt to shift to Medicare the responsibility for payment of medical expenses for the treatment of a work-related condition, the settlement will not be recognized. For example, if the parties to a settlement attempt to maximize the amount of disability benefits paid under workers' compensation by releasing the workers' compensation carrier from liability for medical expenses for a particular condition even though the facts show that the condition is work-related, Medicare will not pay for treatment of that condition.

(c) Lump-sum compromise settlement: Effect on services furnished before the

§ 411.45 Basis for conditional Medicare payment in workers' compensation cases.

A conditional Medicare payment may be made under either of the following circumstances:

(a) The beneficiary has filed a proper claim for workers' compensation benefits, but the intermediary or carrier determines that the workers' compensation carrier will not pay promptly. This includes cases in which a workers' compensation carrier has denied a claim.

(b) The beneficiary, because of physical or mental incapacity, failed to file a proper claim.

§ 411.43 Beneficiary's responsibility with respect to workers' compensation.

(a) The beneficiary is responsible for taking whatever action is necessary to obtain any payment that can reasonably be expected under workers' compensation.

(b) Except as specified in §411.45(a), Medicare does not pay until the beneficiary has exhausted his or her remedies under workers' compensation.

(c) Except as specified in §411.45(b), Medicare does not pay for services that would have been covered under workers' compensation if the beneficiary had filed a proper claim.

(d) However, if a claim is denied for reasons other than not being a proper claim, Medicare pays for the services if they are covered under Medicare.

§ 411.44 Limitations on Medicare payment.

(1) Medicare does not pay for any services for which—

(i) Payment has been made, or can reasonably be expected to be made promptly under a workers' compensation law or plan of the United States or a state; or

(ii) Payment could be made under the Federal Black Lung Program, but is precluded solely because the provider of the services has failed to secure, from the Department of Labor, a provider number to include in the claim.

(2) If the payment for a service may not be made under workers' compensation because the service is furnished by a source not authorized to provide that service under the particular workers' compensation program, Medicare pays for the service if it is a covered service.

(3) Medicare makes secondary payments in accordance with §§411.32 and 411.33.

§ 411.42 Basis for conditional Medicare payment in workers' compensation cases.

A conditional Medicare payment may be made under either of the following circumstances:

(a) The beneficiary has filed a proper claim for workers' compensation benefits, but the intermediary or carrier determines that the workers' compensation carrier will not pay promptly. This includes cases in which a workers' compensation carrier has denied a claim.

(b) The beneficiary, because of physical or mental incapacity, failed to file a proper claim.

§ 411.41 Basis for conditional Medicare payment in workers' compensation cases.

A conditional Medicare payment may be made under either of the following circumstances:

(a) The beneficiary has filed a proper claim for workers' compensation benefits, but the intermediary or carrier determines that the workers' compensation carrier will not pay promptly. This includes cases in which a workers' compensation carrier has denied a claim.

(b) The beneficiary, because of physical or mental incapacity, failed to file a proper claim.
§ 411.47 Apportionment of a lump-sum compromise settlement of a workers' compensation claim.

(a) Determining amount of compromise settlement considered as a payment for medical expenses. (1) If a compromise settlement allocates a portion of the payment for medical expenses and also gives reasonable recognition to the income replacement element, that apportionment may be accepted as a basis for determining Medicare payments.

(2) If the settlement does not give reasonable recognition to both elements of a workers' compensation award or does not apportion the sum granted, the portion to be considered as payment for medical expenses is computed as follows:

(i) Determine the ratio of the amount awarded (less the reasonable and necessary costs incurred in procuring the settlement) to the total amount that would have been payable under workers' compensation if the claim had not been compromised.

(ii) Multiply that ratio by the total medical expenses incurred as a result of the injury or disease up to the date of the settlement. The product is the amount of the workers' compensation settlement to be considered as payment for medical expenses.

Example: As the result of a work injury, an individual suffered loss of income and incurred medical expenses for which the total workers' compensation payment would have been $24,000 if the case had not been compromised. The medical expenses amounted to $18,000. The workers' compensation carrier made a settlement with the beneficiary under which it paid $8,000 in total. A separate award was made for legal fees. Since the workers' compensation compromise settlement was for one-third of the amount which would have been payable under workers' compensation had the case not been compromised ($8,000/$24,000 = 1/3), the workers' compensation compromise settlement is considered to have paid for one-third of the total medical expenses (1/3 × $18,000 = $6,000).

(b) Determining the amount of the Medicare overpayment. When conditional Medicare payments have been made, and the beneficiary receives a compromise settlement payment, the Medicare overpayment is determined as set forth in this paragraph (b). The amount of the workers' compensation payment that is considered to be for medical expenses (as determined under paragraph (a) of this section) is applied, at the workers' compensation rate of payment prevailing in the particular jurisdiction, in the following order:

(1) First to any beneficiary payments for services payable under workers' compensation but not covered under Medicare.

(2) Then to any beneficiary payments for services payable under workers' compensation and also covered under Medicare Part B. (These include deductible and coinsurance amounts and, in unassigned cases, the charge in excess of the reasonable charge.)

(3) Last to any beneficiary payments for services payable under workers' compensation and also covered under Medicare Part A. (These include Part A deductible and coinsurance amounts and charges for services furnished after benefits are exhausted.)

The difference between the amount of the workers' compensation payment for medical expenses and any beneficiary payments constitutes the Medicare overpayment. The beneficiary is liable for that amount.

Example: In the example in paragraph (a) of this section, it was determined that the workers' compensation settlement paid for $6,000 of the total medical expenses. The $18,000 in medical expenses included $1,500 in
charges for services not covered under Medicare, $7,500 in charges for services covered under Medicare Part B, and $9,000 in hospital charges for services covered under Medicare Part A. All charges were at the workers' compensation payment rate, that is, in amounts the provider or supplier must accept as payment in full.

The Medicare reasonable charge for physicians’ services was $7,000 and Medicare paid $5,600 (80 percent of the reasonable charge). The Part B deductible had been met. The Medicare payment rate for the hospital services was $8,000. Medicare paid the hospital $7,480 ($8,000—the Part A deductible of $520).

In this situation, the beneficiary’s payments totalled $3,920:

<table>
<thead>
<tr>
<th>Services not covered under Medicare</th>
<th>$1,500</th>
</tr>
</thead>
<tbody>
<tr>
<td>Excess of physicians’ charges over reasonable charges</td>
<td>$500</td>
</tr>
<tr>
<td>Medicare Part B coinsurance</td>
<td>$1,400</td>
</tr>
<tr>
<td>Part A deductible</td>
<td>$520</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>$3,920</strong></td>
</tr>
</tbody>
</table>

The Medicare overpayment, for which the beneficiary is liable, would be $2,080 ($6,000−$3,920).

Subpart D—Limitations on Medicare Payment for Services Covered Under Liability or No-Fault Insurance

§ 411.50 General provisions.

(a) Limits on applicability. The provisions of this subpart C do not apply to any services required because of accidents that occurred before December 5, 1980.

(b) Definitions.

Automobile means any self-propelled land vehicle of a type that must be registered and licensed in the State in which it is owned.

Liability insurance means insurance (including a self-insured plan) that provides payment based on legal liability for injury or illness or damage to property. It includes, but is not limited to, automobile liability insurance, uninsured motorist insurance, underinsured motorist insurance, homeowners’ liability insurance, malpractice insurance, product liability insurance, and general casualty insurance.

Liability insurance payment means a payment by a liability insurer, or an out-of-pocket payment, including a payment to cover a deductible required by a liability insurance policy, by any individual or other entity that carries liability insurance or is covered by a self-insured plan.

No-fault insurance means insurance that pays for medical expenses for injuries sustained on the property or premises of the insured, or in the use, occupancy, or operation of an automobile, regardless of who may have been responsible for causing the accident. This insurance includes but is not limited to automobile, homeowners, and commercial plans. It is sometimes called “medical payments coverage”, “personal injury protection”, or “medical expense coverage”.

Prompt or promptly, when used in connection with payment by a liability insurer means payment within 120 days after the earlier of the following:

(1) The date a claim is filed with an insurer or a lien is filed against a potential liability settlement.

(2) The date the service was furnished or, in the case of inpatient hospital services, the date of discharge.

Self-insured plan means a plan under which an individual, or a private or governmental entity, carries its own risk instead of taking out insurance with a carrier. The term includes a plan of an individual or other entity engaged in a business, trade, or profession, a plan of a non-profit organization such as a social, fraternal, labor, educational, religious, or professional organization, and the plan established by the Federal government to pay liability claims under the Federal Tort Claims Act.

Underinsured motorist insurance means insurance under which the policyholder’s level of protection against losses caused by another is extended to compensate for inadequate coverage in the other party’s policy or plan.

Uninsured motorist insurance means insurance under which the policyholder’s insurer will pay for damages caused by a motorist who has no automobile liability insurance or who carries less than the amount of insurance required by law, or is underinsured.

(c) Limitation on payment for services covered under no-fault insurance. Except as provided under §§ 411.52 and 411.53 with respect to conditional payments, Medicare does not pay for the following:
§ 411.51 Beneficiary's responsibility with respect to no-fault insurance.

(a) The beneficiary is responsible for taking whatever action is necessary to obtain any payment that can reasonably be expected under no-fault insurance.

(b) Except as specified in § 411.53, Medicare does not pay until the beneficiary has exhausted his or her remedies under no-fault insurance.

(c) Except as specified in § 411.53, Medicare does not pay for services that would have been covered by the no-fault insurance if the beneficiary had filed a proper claim.

(d) However, if a claim is denied for reasons other than not being a proper claim, Medicare pays for the services if they are covered under Medicare.

§ 411.52 Basis for conditional Medicare payment in liability cases.

If HCFA has information that services for which Medicare benefits have been claimed are for treatment of an injury or illness that was allegedly caused by another party, a conditional Medicare payment may be made.

§ 411.53 Basis for conditional Medicare payment in no-fault cases.

A conditional Medicare payment may be made in no-fault cases under either of the following circumstances:

(a) The beneficiary, or the provider or supplier, has filed a proper claim for no-fault insurance benefits but the intermediary or carrier determines that the no-fault insurer will not pay promptly for any reason other than the circumstances described in § 411.32(a)(1). This includes cases in which the no-fault insurance carrier has denied the claim.

(b) The beneficiary, because of physical or mental incapacity, failed to meet a claim-filing requirement stipulated in the policy.

§ 411.54 Limitation on charges when a beneficiary has received a liability insurance payment or has a claim pending against a liability insurer.

(a) Definition. As used in this section, Medicare-covered services means services for which Medicare benefits are payable or would be payable except for applicable Medicare deductible and coinsurance provisions. Medicare benefits are payable notwithstanding potential liability insurance payments, but are recoverable in accordance with § 411.24.

(b) Applicability. This section applies when a beneficiary has received a liability insurance payment or has a claim pending against a liability insurer for injuries or illness allegedly caused by another party.

(c) Basic rules—(1) Itemized bill. A hospital must, upon request, furnish to the beneficiary or his or her representative an itemized bill of the hospital's charges.

(2) Specific limitations. Except as provided in paragraph (d) of this section, the provider or supplier—

(i) May not bill the liability insurer nor place a lien against the beneficiary's liability insurance settlement for Medicare covered services.

(ii) May only bill Medicare for Medicare-covered services; and

(iii) May bill the beneficiary only for applicable Medicare deductible and coinsurance amounts plus the amount of any charges that may be made to a beneficiary under § 413.35 of this chapter (when cost limits are applied to the services) or under § 489.32 of this chapter (when services are partially covered).

(d) Exceptions—(1) Nonparticipating suppliers. The limitations of paragraph (c)(2) of this section do not apply if the services were furnished by a supplier that is not a participating supplier and has not accepted assignment for the services or has not claimed payment for them under § 424.64 of this chapter.

(2) Prepaid health plans. If the services were furnished through an organization that has a contract under section 1876 of the Act (that is, through an
§ 411.100 Basis and scope.

(a) Statutory basis. (1) Section 1862(b) of the Act provides in part that Medicare is secondary payer, under specified conditions, for services covered under any of the following:
   (i) Group health plans of employers that employ at least 20 employees and that cover Medicare beneficiaries who are under age 65, entitled to Medicare on the basis of age or disability, or eligible for, or entitled to Medicare on the basis of ESRD.
   (2) Sections 1862(b)(1)(A), (B), and (C) of the Act provide that group health plans and large group health plans may not take into account that the individual is entitled to Medicare on the basis of age or disability, or eligible for, or entitled to Medicare on the basis of ESRD.
   (3) Section 1862(b)(1)(A)(i)(II) of the Act provides that group health plans of employers of 20 or more employees must provide to any employee or spouse age 65 or older the same benefits, under the same conditions, that it provides to employees and spouses under 65. The requirement applies regardless of whether the individual or spouse 65 or older is entitled to Medicare.
   (4) Section 1862(b)(1)(C)(ii) of the Act provides that group health plans may not differentiate in the benefits they provide between individuals who have ESRD and other individuals covered under the plan on the basis of the existence of ESRD, the need for renal dialysis, or in any other manner. Actions that constitute “differentiating” are listed in §411.163(b).

(b) Scope. This subpart sets forth general rules pertinent to—
   (1) Medicare payment for services that are covered under a group health plan and are furnished to certain beneficiaries who are entitled on the basis of ESRD, age, or disability.
   (2) The prohibition against taking into account Medicare entitlement based on age or disability, or Medicare eligibility or entitlement based on ESRD.
   (3) The prohibition against differentiation in benefits between individuals...
who have ESRD and other individuals covered under the plan.

(4) The requirement to provide to those 65 or over the same benefits under the same conditions as are provided to those under 65.

(5) The appeals procedures for group health plans that HCFA determines are nonconforming plans.

§ 411.101 Definitions.

As used in this subpart and in parts F through H of this part—


Days means calendar days.

Employee (subject to the special rules in §411.104) means an individual who—

(1) Is working for an employer; or

(2) Is not working for an employer but is receiving payments that are subject to FICA taxes, or would be subject to FICA taxes except that the employer is exempt from those taxes under the Internal Revenue Code.

Employer means, in addition to individuals (including self-employed persons) and organizations engaged in a trade or business, other entities exempt from income tax such as religious, charitable, and educational institutions, the governments of the United States, the individual States, Puerto Rico, the Virgin Islands, Guam, American Samoa, the Northern Mariana Islands, and the District of Columbia, and the agencies, instrumentalities, and political subdivisions of these governments.

FICA stands for the Federal Insurance Contributions Act, the law that imposes social security taxes on employers and employees under section 21 of the Internal Revenue Code.

Group health plan (GHP) means any arrangement made by one or more employers or employee organizations to provide health care directly or through other methods such as insurance or reimbursement, to current or former employees, the employer, others associated or formerly associated with the employer in a business relationship, or their families, that—

(1) Is of, or contributed to by, one or more employers or employee organizations.

(2) If it involves more than one employer or employee organization, provides for common administration.

(3) Provides substantially the same benefits or the same benefit options to all those enrolled under the arrangement.

The term includes self-insured plans, plans of governmental entities (Federal, State and local), and employee organization plans; that is, union plans, employee health and welfare funds or other employee organization plans. The term also includes employee-pay-all plans, which are plans under the auspices of one or more employers or employee organizations but which receive no financial contributions from them. The term does not include a plan that is unavailable to employees; for example, a plan only for self-employed persons.

IRC stands for Internal Revenue Code.

IRS stands for Internal Revenue Service.

Large group health plan (LGHP) means

(1) A single employer or employee organization that employed at least 100 full-time or part-time employees on 50 percent or more of its regular business days during the previous calendar year; or

(2) Two or more employers, or employee organizations, at least one of which employed at least 100 full-time or part-time employees on 50 percent or more of its regular business days during the previous calendar year.

MSP stands for Medicare secondary payer.

Multi-employer plan means a plan that is sponsored jointly by two or more employers (sometimes called a multiple-employer plan) or by employers and unions (sometimes under the Taft-Hartley law).

Self-employed person encompasses consultants, owners of businesses, and directors of corporations, and members of the clergy and religious orders who are paid for their services by a religious body or other entity.

Similarly situated individual means—

(1) In the case of employees, other employees enrolled or seeking to enroll in the plan; and
(2) In the case of other categories of individuals, other persons in any of those categories who are enrolled or seeking to enroll in the plan.

§ 411.102 Basic prohibitions and requirements.

(a) ESRD. (1) A group health plan of any size—(i) May not take into account the ESRD-based Medicare eligibility or entitlement of any individual who is covered or seeks to be covered under the plan; and

(ii) May not differentiate in the benefits it provides between individuals with ESRD and other individuals covered under the plan, on the basis of the existence of ESRD, or the need for dialysis, or in any other manner.

(2) The prohibitions of paragraph (a) of this section do not prohibit a plan from paying benefits secondary to Medicare after the first 18 months of ESRD-based eligibility or entitlement.

(b) Age. A GHP of an employer or employee organization of at least 20 employees—

(1) May not take into account the age-based Medicare entitlement of an individual or spouse age 65 or older who is covered (or seeks to be covered) under the plan by virtue of current employment status; and

(2) Must provide, to employees age 65 or older and to spouses age 65 or older of employees of any age, the same benefits under the same conditions as it provides to employees and spouses under age 65.

(c) Disability. A GHP of an employer or employee organization of at least 100 employees may not take into account the disability-based Medicare entitlement of any individual who is covered (or seeks to be covered) under the plan by virtue of current employment status.

§ 411.103 Prohibition against financial and other incentives.

(a) General rule. An employer or other entity (for example, an insurer) is prohibited from offering Medicare beneficiaries financial or other benefits as incentives not to enroll in, or to terminate enrollment in, a GHP that is, or would be, primary to Medicare. This prohibition precludes offering to Medicare beneficiaries an alternative to the employer primary plan (for example, coverage of prescription drugs) unless the beneficiary has primary coverage other than Medicare. An example would be primary coverage through his own or a spouse's employer.

(b) Penalty for violation. (1) Any entity that violates the prohibition of paragraph (a) of this section is subject to a civil money penalty of up to $5,000 for each violation; and

(2) The provisions of section 1128A of the Act (other than subsections (a) and (b)) apply to the civil money penalty of up to $5,000 in the same manner as the provisions apply to a penalty or proceeding under section 1128A(a).

§ 411.104 Current employment status.

(a) General rule. An individual has current employment status if—

(1) The individual is actively working as an employee, is the employer (including a self-employed person), or is associated with the employer in a business relationship; or

(2) The individual is not actively working and—

(i) Is receiving disability benefits from an employer for up to 6 months (the first 6 months of employer disability benefits are subject to FICA taxes); or

(ii) Retains employment rights in the industry and has not had his employment terminated by the employer, if the employer provides the coverage (or has not had his membership in the employee organization terminated, if the employee organization provides the coverage), is not receiving disability benefits from an employer for more than 6 months, is not receiving disability benefits from Social Security, and has GHP coverage that is not pursuant to COBRA continuation coverage (26 U.S.C. 4980B; 29 U.S.C. 1161-1168; 42 U.S.C. 300bb-1 et seq.). Whether or not the individual is receiving pay during the period of nonwork is not a factor.

(b) Persons who retain employment rights. For purposes of paragraph (a)(2) of this section, persons who retain employment rights include but are not limited to—

(1) Persons who are furloughed, temporarily laid off, or who are on sick leave;
Teachers and seasonal workers who normally do not work throughout the year; and

Persons who have health coverage that extends beyond or between active employment periods; for example, based on an hours bank arrangement. (Active union members often have hours bank coverage.)

Coverage by virtue of current employment status. An individual has coverage by virtue of current employment status with an employer if—

(1) the individual has GHP or LGHP coverage based on employment, including coverage based on a certain number of hours worked for that employer or a certain level of commissions earned from work for that employer at any time and

(2) the individual has current employment status with that employer, as defined in paragraph (a) of this section.

Special rule: Self-employed person. A self-employed individual is considered to have GHP or LGHP coverage by virtue of current employment status during a particular tax year only if, during the preceding tax year, the individual's net earnings, from work in that year related to the employer that offers the group health coverage, are at least equal to the amount specified in section 211(b)(2) of the Act, which defines "self-employment income" for social security purposes.

Special Rule: members of religious orders and members of clergy. (1) Members of religious orders who have not taken a vow of poverty. A member of a religious order who has not taken a vow of poverty is considered to have current employment status with the religious order if—

(i) The religious order pays FICA taxes on behalf of that member; or

(ii) The individual is receiving cash remuneration from the religious order.

(2) Members of religious orders who have taken a vow of poverty. A member of a religious order whose members are required to take a vow of poverty is not considered to be employed by the order if the services he or she performs as a member of the order are considered employment only because the order extends social security coverage under section 3121(r) of the IRC. This exemption applies retroactively to services performed as a member of the order, beginning with the effective dates of the MSP provisions for the aged and the disabled, respectively. The exemption does not apply to services performed for employers outside of the order.

(3) Members of the clergy. A member of the clergy is considered to have current employment status with a church or other religious organization if the individual is receiving cash remuneration from the church or other religious organization for services rendered.

Special rule: Delayed compensation subject to FICA taxes. An individual who is not working is not considered an employee solely on the basis of receiving delayed compensation payments for previous periods of work even if those payments are subject to FICA taxes (or would be subject to FICA taxes if the employer were not exempt from paying those taxes). For example, an individual who is not working in 1993 and receives payments subject to FICA taxes for work performed in 1992 is not considered to be an employee in 1993 solely on the basis of receiving those payments.

Aggregation rules.

The following rules apply in determining the number and size of employers, as required by the MSP provisions for the aged and disabled:

(a) All employers that are treated as a single employer under subsection (a) or (b) of section 52 of the Internal Revenue Code (IRC) of 1986 (26 U.S.C. 52 (a) and (b)) are treated as a single employer.

(b) All employees of the members of an affiliated service group (as defined in section 414(m) of the IRC (26 U.S.C. 414m)) are treated as employed by a single employer.

(c) Leased employees (as defined in section 414(n)(2) of the IRC (26 U.S.C. 414n)(2)) are treated as employees of the person for whom they perform services to the same extent as they are treated under section 414(n) of the IRC.

(d) In applying the IRC provisions identified in this section, HCFA relies upon regulations and decisions of the Secretary of the Treasury respecting those provisions.
§ 411.108 Taking into account entitlement to Medicare.

(a) Examples of actions that constitute "taking into account": Actions by GHPs or LGHPs that constitute taking into account that an individual is entitled to Medicare on the basis of ESRD, age, or disability (or eligible on the basis of ESRD) include, but are not limited to, the following:

1. Failure to pay primary benefits as required by subparts F, G, and H of this part 411.
2. Offering coverage that is secondary to Medicare to individuals entitled to Medicare.
3. Terminating coverage because the individual has become entitled to Medicare, except as permitted under COBRA continuation coverage provisions (26 U.S.C. 4980B(f)(2)(B)(iv); 29 U.S.C. 1162(2)(D); and 42 U.S.C. 300bb-2(2)(D)).
4. In the case of a LGHP, denying or terminating coverage because an individual is entitled to Medicare on the basis of disability without denying or terminating coverage for similarly situated individuals who are not entitled to Medicare on the basis of disability.
5. Imposing limitations on benefits for a Medicare entitled individual that do not apply to others enrolled in the plan, such as providing less comprehensive health care coverage, excluding benefits, reducing benefits, charging higher deductibles or coinsurance, providing for lower annual or lifetime benefit limits, or more restrictive pre-existing illness limitations.
6. Charging a Medicare entitled individual higher premiums.
7. Requiring a Medicare entitled individual to wait longer for coverage to begin.
8. Paying providers and suppliers less for services furnished to a Medicare beneficiary than for the same services furnished to an enrollee who is not entitled to Medicare.
9. Providing misleading or incomplete information that would have the effect of inducing a Medicare entitled individual to reject the employer plan, thereby making Medicare the primary payer. An example of this would be informing the beneficiary of the right to accept or reject the employer plan but failing to inform the individual that, if he or she rejects the plan, the plan will not be permitted to provide or pay for secondary benefits.
10. Including in its health insurance cards, claims forms, or brochures distributed to beneficiaries, providers, and suppliers, instructions to bill Medicare first for services furnished to Medicare beneficiaries without stipulating that such action may be taken only when Medicare is the primary payer.
11. Refusing to enroll an individual for whom Medicare would be secondary payer, when enrollment is available to similarly situated individuals for whom Medicare would not be secondary payer.

(b) Permissible actions.
1. If a GHP or LGHP makes benefit distinctions among various categories of individuals (distinctions unrelated to the fact that the individual is disabled, based, for instance, on length of time employed, occupation, or marital status), the GHP or LGHP may make the same distinctions among the same categories of individuals entitled to Medicare whose plan coverage is based on current employment status. For example, if a GHP or LGHP does not offer coverage to employees who have worked less than one year and who are entitled to Medicare on the basis of disability or age, the GHP or LGHP is not required to offer coverage to employees who have worked less than one year and who are not entitled to Medicare on the basis of disability or age.
2. A GHP or LGHP may pay benefits secondary to Medicare for an aged or disabled beneficiary who has current employment status if the plan coverage is COBRA continuation coverage because of reduced hours of work. Medicare is primary payer for this beneficiary because, although he or she has current employment status, the GHP coverage is by virtue of the COBRA law rather than by virtue of the current employment status.
3. A GHP may terminate COBRA continuation coverage of an individual who becomes entitled to Medicare on the basis of ESRD, when permitted under the COBRA provisions.

[60 FR 45962, Aug. 31, 1995; 60 FR 53876, Oct. 18, 1995]
§ 411.110 Basis for determination of nonconformance.

(a) A "determination of nonconformance" is a HCFA determination that a GHP or LGHP is a nonconforming plan as provided in this section.

(b) HCFA makes a determination of nonconformance for a GHP or LGHP that, at any time during a calendar year, fails to comply with any of the following statutory provisions:

1. The prohibition against taking into account that a beneficiary who is covered or seeks to be covered under the plan is entitled to Medicare on the basis of ESRD, age, or disability, or eligible on the basis of ESRD.
2. The nondifferentiation clause for individuals with ESRD.
3. The equal benefits clause for the working aged.
4. The obligation to refund conditional Medicare primary payments.

(c) HCFA may make a determination of nonconformance for a GHP or LGHP that fails to respond to a request for information, or to provide correct information, either voluntarily or in response to a HCFA request, on the plan’s primary payment obligation with respect to a given beneficiary, if that failure contributes to either or both of the following:

1. Medicare erroneously making a primary payment.
2. A delay or foreclosure of HCFA’s ability to recover an erroneous primary payment.

§ 411.112 Documentation of conformance.

(a) Acceptable documentation. HCFA may require a GHP or LGHP to demonstrate that it has complied with the Medicare secondary payer provisions and to submit supporting documentation by an official authorized to act on behalf of the entity, under penalty of perjury. The following are examples of documentation that may be acceptable:

1. A copy of the employer’s plan or policy that specifies the services covered, conditions of coverage, benefit levels and limitations with respect to persons entitled to Medicare on the basis of ESRD, age, or disability as compared to the provisions applicable to other enrollees and potential enrollees.

2. An explanation of the plan’s allegation that it does not owe HCFA any amount HCFA claims the plan owes as repayment for conditional or mistaken Medicare primary payments.

(b) Lack of acceptable documentation. If a GHP or LGHP fails to provide acceptable evidence or documentation that it has complied with the MSP prohibitions and requirements set forth in § 411.110, HCFA may make a determination of nonconformance for both the year in which the services were furnished and the year in which the request for information was made.

§ 411.114 Determination of nonconformance.

(a) Starting dates for determination of nonconformance. HCFA’s authority to determine nonconformance of GHPs begins on the following dates:

1. On January 1, 1987 for MSP provisions that affect the disabled.
2. On December 20, 1989 for MSP provisions that affect ESRD beneficiaries and the working aged.

(b) Special rule for failure to repay. A GHP that fails to comply with § 411.110 (a)(1), (a)(2), or (a)(3) in a particular year is nonconforming for that year. If, in a subsequent year, that plan fails to repay the resulting mistaken primary payments (in accordance with § 411.110(a)(4)), the plan is also nonconforming for the subsequent year. For example, if a plan paid secondary for the working aged in 1991, that plan was nonconforming for 1991. If in 1994 HCFA identifies mistaken primary payments attributable to the 1991 violation, and the plan refuses to repay, it is also nonconforming for 1994.

§ 411.115 Notice of determination of nonconformance.

(a) Notice to the GHP or LGHP. (1) If HCFA determines that a GHP or an LGHP is nonconforming with respect to a particular calendar year, HCFA mails to the plan written notice of the following:

(i) The determination.
(ii) The basis for the determination.
(iii) The right of the parties to request a hearing.
(iv) An explanation of the procedure for requesting a hearing.
(v) The tax that may be assessed by the IRS in accordance with section 5000 of the IRC.
(vi) The fact that if none of the parties requests a hearing within 65 days from the date of its notice, the determination is binding on all parties unless it is reopened in accordance with § 411.126.

(2) The notice also states that the plan must, within 30 days from the date of its notice, submit to HCFA the names and addresses of all employers and employee organizations that contributed to the plan during the calendar year for which HCFA has determined nonconformance.

(b) Notice to contributing employers and employee organizations. HCFA mails written notice of the determination, including all the information specified in paragraph (a)(1) of this section, to all contributing employers and employee organizations already known to HCFA or identified by the plan in accordance with paragraph (a)(2) of this section. Employers and employee organizations have 65 days from the date of their notice to request a hearing.

§ 411.121 Hearing procedures.
(a) Nature of hearing. (1) If any of the parties requests a hearing within 65 days from the date on the notice of the determination of nonconformance, the HCFA Administrator appoints a hearing officer.
(2) If no party files a request within the 65-day period, the initial determination of nonconformance is binding upon all parties unless it is reopened in accordance with § 411.126.
(3) If more than one party requests a hearing, the hearing officer conducts a single hearing in which all parties may participate.
(4) On the record review. Ordinarily, the hearing officer makes a decision based upon review of the data and documents on which HCFA based its determination of nonconformance and any other documentation submitted by any of the parties within 65 days from the date on the notice.
(5) Oral hearing. The hearing officer may provide for an oral hearing either on his or her own motion or in response to a party's request if the party demonstrates to the hearing officer's satisfaction that an oral hearing is necessary. Within 30 days of receipt of the request, the hearing officer gives all known parties written notice of the request and whether the request for oral hearing is granted.

(b) Notice of time and place of oral hearing. If the hearing officer provides an oral hearing, he or she gives all known parties written notice of the time and place of the hearing at least 30 days before the scheduled date.

(c) Prehearing discovery. (1) The hearing officer may permit prehearing discovery if it is requested by a party at least 30 days before the scheduled date of the hearing.
(2) If the hearing officer approves the request, he or she—
(i) Provides a reasonable time for inspection and reproduction of documents; and
(3) The hearing officer's orders on all discovery matters are final.
(d) Conduct of hearing. The hearing officer determines the conduct of the hearing, including the order in which
§ 411.122 Hearing officer's decision.

(a) Timing. (1) If the decision is based on a review of the record, the hearing officer mails the decision to all known parties within 120 days from the date of receipt of the request for hearing.

(2) If the decision is based on an oral hearing, the hearing officer mails the decision to all known parties within 120 days from the conclusion of the hearing.

(b) Basis, content, and distribution of hearing decision. (1) The written decision is based on substantial evidence and contains findings of fact, a statement of reasons, and conclusions of law.

(2) The hearing officer mails a copy of the decision to each of the parties, by certified mail, return receipt requested, and includes a notice that the administrator may review the hearing decision at the request of a party or on his or her own motion.

(c) Effect of hearing decision. The hearing officer's decision is the final Departmental decision and is binding upon all parties unless the Administrator chooses to review that decision in accordance with § 411.124 or it is reopened by the hearing officer in accordance with § 411.126.

§ 411.124 Administrator's review of hearing decision.

(a) Request for review. A party's request for review of a hearing officer's
decision must be in writing (not in facsimile or other electronic medium) and must be received by the Administrator within 25 days from the date on the decision.

(b) Office of the Attorney Advisor responsibility. The Office of the Attorney Advisor examines the hearing officer's decision, the requests made by any of the parties or HCFA, and any submission made in accordance with the provisions of this section in order to assist the Administrator in deciding whether to review the decision.

(c) Administrator's discretion. The Administrator may—

(1) Review or decline to review the hearing officer's decision;

(2) Exercise this discretion on his or her own motion or in response to a request from any of the parties; and

(3) Delegate review responsibility to the Deputy Administrator. (As used in this section, the term "Administrator" includes "Deputy Administrator" if review responsibility has been delegated.)

(d) Basis for decision to review. In deciding whether to review a hearing officer's decision, the Administrator considers—

(1) Whether the decision—

(i) Is based on a correct interpretation of law, regulation, or HCFA Ruling;

(ii) Is supported by substantial evidence;

(iii) Presents a significant policy issue having a basis in law and regulations;

(iv) Requires clarification, amplification, or an alternative legal basis for the decision; and

(v) Is within the authority provided by statute, regulation, or HCFA Ruling; and

(2) Whether review may lead to the issuance of a HCFA Ruling or other directive needed to clarify a statute or regulation.

(e) Notice of decision to review or not to review. (1) The Administrator gives all parties prompt written notice of his or her decision to review or not to review.

(2) The notice of a decision to review identifies the specific issues the Administrator will consider.

(f) Response to notice of decision to review. (1) Within 20 days from the date on a notice of the Administrator's decision to review a hearing officer's decision, any of the parties may file with the Administrator any of the following:

(i) Proposed findings and conclusions.

(ii) Supporting views or exceptions to the hearing officer's decision.

(iii) Supporting reasons for the proposed findings and exceptions.

(iv) A rebuttal to another party's request for review or to other submissions already filed with the Administrator.

(2) The submissions must be limited to the issues the Administrator has decided to review and confined to the record established by the hearing officer.

(3) All communications from the parties concerning a hearing officer's decision being reviewed by the Administrator must be in writing (not in facsimile or other electronic medium) and must include a certification that copies have been sent to all other parties.

(4) The Administrator does not consider any communication that does not meet the requirements of this paragraph.

(g) Administrator's review decision. (1) The Administrator bases his or her decision on the following:

(i) The entire record developed by the hearing officer.

(ii) Any materials submitted in connection with the hearing or under paragraph (f) of this section.

(iii) Generally known facts not subject to reasonable dispute.

(2) The Administrator mails copies of the review decision to all parties within 120 days from the date of the hearing officer's decision.

(3) The Administrator's review decision may affirm, reverse, or modify the hearing decision or may remand the case to the hearing officer.

(h) Basis and effect of remand. (1) Basis. The bases for remand do not include the following:

(i) Evidence that existed at the time of the hearing and that was known or could reasonably have been expected to be known.

(ii) A court case that was either not available at the time of the hearing or was decided after the hearing.
(iii) Change of the parties’ representation.
(iv) An alternative legal basis for an issue in dispute.

(2) Effect of remand. (i) The Administrator may instruct the hearing officer to take further action with respect to the development of additional facts or new issues or to consider the applicability of laws or regulations other than those considered during the hearing.
(ii) The hearing officer takes the action in accordance with the Administrator’s instructions in the remand notice and again issues a decision.
(iii) The Administrator may review or decline to review the hearing officer’s remand decision in accordance with the procedures set forth in this section.

(i) Finality of decision. The Administrator’s review decision, or the hearing officer’s decision following remand, is the final Departmental decision and is binding on all parties unless the Administrator chooses to review the decision in accordance with this section, or the decision is reopened in accordance with §411.126.

§ 411.126 Reopening of determinations and decisions.

(a) A determination that a GHP or LGHP is a nonconforming GHP or the decision or revised decision of a hearing officer or of the HCFA Administrator may be reopened within 12 months from the date on the notice of determination or decision or revised decision, for any reason by the entity that issued the determination or decision.
(b) The decision to reopen or not to reopen is not appealable.

§ 411.130 Referral to Internal Revenue Service (IRS).

(a) HCFA responsibility. After HCFA determines that a plan has been a nonconforming GHP in a particular year, it refers its determination to the IRS, but only after the parties have exhausted all HCFA appeal rights with respect to the determination.

(b) IRS responsibility. The IRS administers section 5000 of the IRC, which imposes a tax on employers (other than governmental entities) and employee organizations that contribute to a nonconforming GHP. The tax is equal to 25 percent of the employer’s or employee organization’s expenses, incurred during the calendar year in which the plan is a nonconforming GHP, for each GHP, both conforming and nonconforming, to which the employer or employee organization contributes.

Subpart F—Special Rules: Individuals Eligible or Entitled on the Basis of ESRD, Who Are Also Covered Under Group Health Plans

§ 411.160 Scope.

This subpart sets forth special rules that apply to individuals who are eligible for, or entitled to, Medicare on the basis of ESRD. (Section 406.13 of this chapter contains the rules for eligibility and entitlement based on ESRD.)

[60 FR 45367, Aug. 31, 1995]

§ 411.161 Prohibition against taking into account Medicare eligibility or entitlement or differentiating benefits.

(a) Taking into account. (1) Basic rule. A GHP may not take into account that an individual is eligible for or entitled to Medicare benefits on the basis of ESRD during the coordination period specified in §411.162(b) and (c). Examples of actions that constitute taking into account Medicare entitlement are listed in §411.108(a).

(2) Applicability. This prohibition applies for ESRD-based Medicare eligibility to the same extent as for ESRD-based Medicare entitlement. An individual who has ESRD but who has not filed an application for entitlement to Medicare on that basis is eligible for Medicare based on ESRD for purposes of paragraphs (b)(2) and (c)(2) through (c)(4) of §411.162 if the individual meets the other requirements of §406.13 of this chapter.

(3) Relation to COBRA continuation coverage. This rule does not prohibit the termination of GHP coverage under title X of COBRA when termination of that coverage is expressly permitted, upon entitlement to Medicare, under 26 U.S.C. 4980B(f)(2)(B)(iv); 29 U.S.C. 4980B(f)(2)(B)(iv); 29 U.S.C.
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COBRA requires that certain group health plans offer continuation of plan coverage for 18 to 36 months after the occurrence of certain "qualifying events," including loss of employment or reduction of employment hours. Those are events that otherwise would result in loss of group health plan coverage unless the individual is given the opportunity to elect, and does so elect, to continue plan coverage at his or her own expense. With one exception, the COBRA amendments expressly permit termination of continuation coverage upon entitlement to Medicare. The exception is that the plan may not terminate continuation coverage of an individual (and his or her qualified dependents) if the individual retires on or before the date the employer substantially eliminates regular plan coverage by filing for Chapter 11 bankruptcy (26 U.S.C. 4980B (g)(3)(D) and 29 U.S.C. 1167.13(C)).

Mr. Smith works for employer A, and he and his wife are covered through employer A's GHP (Plan A). Neither is eligible for Medicare nor has ESRD. Mrs. Smith works for employer B, and is also covered by employer B's plan (Plan B). Plan B is more comprehensive than Plan A and covers certain items and services which Plan B does not cover, such as prescription drugs. If Mrs. Smith obtains a medical service, Plan B pays primary and Plan A pays secondary. That is, Plan A covers Plan B copayment amounts and items and services that Plan A covers but that Plan B does not.

Mr. Jones also works for employer A, and he and his wife are covered through Plan A. Mrs. Jones develops ESRD and becomes entitled to Medicare on that basis. Plan A pays primary to Medicare during the first 18 months of Medicare entitlement based on ESRD. When Medicare becomes the primary payer, the plan converts Mrs. Jones' coverage to a Medicare supplement policy. That policy pays Medicare deductible and coinsurance amounts but does not pay for items and services not covered by Medicare, which plan

1 COBRA requires that certain group health plans offer continuation of plan coverage for 18 to 36 months after the occurrence of certain "qualifying events," including loss of employment or reduction of employment hours. Those are events that otherwise would result in loss of group health plan coverage unless the individual is given the opportunity to elect, and does so elect, to continue plan coverage at his or her own expense. With one exception, the COBRA amendments expressly permit termination of continuation coverage upon entitlement to Medicare. The exception is that the plan may not terminate continuation coverage of an individual (and his or her qualified dependents) if the individual retires on or before the date the employer substantially eliminates regular plan coverage by filing for Chapter 11 bankruptcy (26 U.S.C. 4980B (g)(3)(D) and 29 U.S.C. 1167.13(C)).

1Situation in which Medicare is secondary to COBRA continuation coverage are set forth in §411.162(a)(3).

(b) Nondifferentiation. (1) A GHP may not differentiate in the benefits it provides between individuals who have ESRD and others enrolled in the plan, on the basis of the existence of ESRD, or the need for renal dialysis, or in any other manner.

(2) GHP actions that constitute differentiation in plan benefits (and that may therefore constitute "taking into account" Medicare eligibility or entitlement) include, but are not limited to the following:

(i) Terminating coverage of individuals with ESRD, when there is no basis for such termination unrelated to ESRD (such as failure to pay plan premiums) that would result in termination for individuals who do not have ESRD.

(ii) Imposing on persons who have ESRD, but not on others enrolled in the plan, benefit limitations such as less comprehensive health plan coverage, reductions in benefits, exclusions of benefits, a higher deductible or coinsurance, a longer waiting period, a lower annual or lifetime benefit limit, or more restrictive preexisting illness limitations.

(iii) Charging individuals with ESRD higher premiums.

(iv) Paying providers and suppliers less for services furnished to individuals who have ESRD than for the same services furnished to those who do not have ESRD, such as paying 80 percent of the Medicare rate for renal dialysis on behalf of a plan enrollee who has ESRD and the usual, reasonable and customary charge for renal dialysis on behalf of an enrollee who does not have ESRD.

(v) Failure to cover routine maintenance dialysis or kidney transplants, when a plan covers other dialysis services or other organ transplants.

(c) Uniform Limitations on particular services permissible. A plan is not prohibited from limiting covered utilization of a particular service as long as the limitation applies uniformly to all plan enrollees. For instance, if a plan limits its coverage of renal dialysis sessions to 30 per year for all plan enrollees, the plan would not be differentiating in the benefits it provides between plan enrollees who have ESRD and those who do not.

(d) Benefits secondary to Medicare. (1) The prohibition against differentiation of benefits does not preclude a plan from paying benefits secondary to Medicare after the expiration of the coordination period described in §411.162(b) and (c), but a plan may not otherwise differentiate, as described in paragraph (b) of this section, in the benefits it provides.

(2) Example—

Mr. Jones also works for employer A, and he and his wife are covered through employer A's GHP (Plan A). Neither is eligible for Medicare nor has ESRD. Mrs. Jones works for employer B, and is also covered by employer B's plan (Plan B). Plan A is more comprehensive than Plan B and covers certain items and services which Plan B does not cover, such as prescription drugs. If Mrs. Jones obtains a medical service, Plan B pays primary and Plan A pays secondary. That is, Plan A covers Plan B copayment amounts and items and services that Plan A covers but that Plan B does not.

Mr. Jones also works for employer A, and he and his wife are covered through Plan A. Mrs. Jones does not have other GHP coverage. Mrs. Jones develops ESRD and becomes entitled to Medicare on that basis. Plan A pays primary to Medicare during the first 18 months of Medicare entitlement based on ESRD. When Medicare becomes the primary payer, the plan converts Mrs. Jones' coverage to a Medicare supplement policy. That policy pays Medicare deductible and coinsurance amounts but does not pay for items and services not covered by Medicare, which plan.
§ 411.162 Medicare benefits secondary to group health plan benefits.

(a) General provisions. (1) Basic rule. Except as provided in §411.163 (with respect to certain individuals who are also entitled on the basis of age or disability), Medicare is secondary to any GHP (including a retirement plan), with respect to benefits that are payable to an individual who is entitled to Medicare on the basis of ESRD, for services furnished during any coordination period determined in accordance with paragraphs (b) and (c) of this section. (No Medicare benefits are payable on behalf of an individual who is eligible but not yet entitled.)

(2) Medicare benefits secondary without regard to size of employer and beneficiary’s employment status. The size of employer and employment status requirements of the MSP provisions for the aged and disabled do not apply with respect to ESRD beneficiaries.

(3) COBRA continuation coverage. Medicare is secondary payer for benefits that a GHP--

(i) Is required to keep in effect under COBRA continuation requirements (as explained in the footnote to §411.161(a)(3)), even after the individual becomes entitled to Medicare; or

(ii) Voluntarily keeps in effect after the individual becomes entitled to Medicare on the basis of ESRD, even though not obligated to do so under the COBRA provisions.

(4) Medicare payments during the coordination period. During the coordination period, HCFA makes Medicare payments as follows:

(i) Primary payments only for Medicare covered services that are—

(A) Furnished to Medicare beneficiaries who have declined to enroll in the GHP;

(B) Not covered under the plan;

(C) Covered under the plan but not available to particular enrollees because they have exhausted their benefits; or

(D) Furnished to individuals whose COBRA continuation coverage has been terminated because of the individual’s Medicare entitlement.

(ii) Secondary payments, within the limits specified in §§411.32 and 411.33, to supplement the amount paid by the GHP if that plan pays only a portion of the charge for the services.

(b) Beginning of coordination period. (1) For individuals who start a course of maintenance dialysis or who receive a kidney transplant before December 1989, the coordination period begins with the earlier of—

(i) The month in which the individual initiated a regular course of renal dialysis; or

(ii) In the case of an individual who received a kidney transplant, the first month in which the individual became entitled to Medicare, or, if earlier, the first month for which the individual would have been entitled to Medicare benefits if he or she had filed an application for such benefits.

(2) For individuals other than those specified in paragraph (b)(1) of this section, the coordination period begins with the earlier of—

(i) The first month in which the individual becomes entitled to Medicare part A on the basis of ESRD; or

(ii) The first month the individual would have become entitled to Medicare part A on the basis of ESRD if he or she had filed an application for such benefits.

(c) End of coordination period. (1) For individuals who start a regular course of renal dialysis or who receive a kidney transplant before December 1989, the coordination period ends with the earlier of the end of the 12th month of dialysis or the end of the 12th month of a transplant. The 12th month of dialysis may be any time from the 9th month through the 12th month of Medicare entitlement, depending on the extent to which the individual was

1HCFA does not pay if noncoverage of services constitutes differentiation as prohibited by §411.161(b).
subject to a waiting period before becoming entitled to Medicare.

(2) The coordination period for the following individuals ends with the earlier of the 12th month of eligibility or the 12th month of entitlement to Medicare part A:

(i) Individuals, other than those specified in paragraph (c)(1) of this section, who became entitled to Medicare part A solely on the basis of ESRD during December 1989 and January 1990.

(ii) Individuals, other than those specified in paragraph (c)(1) of this section, who could have become entitled to Medicare Part A solely on the basis of ESRD during December 1989 and January 1990 if they had filed an application.

(iii) Individuals who become entitled to Medicare part A on the basis of ESRD after September 1997.

(iv) Individuals who can become entitled to Medicare part A on the basis of ESRD after September 1997.

(3) The coordination period for the following individuals ends with the earlier of the 18th month of eligibility or the 18th month of entitlement to Medicare part A:

(i) Individuals, other than those specified in paragraph (c)(1) of this section, who become entitled to Medicare part A on the basis of ESRD from February 1990 through April 1997.

(ii) Individuals, other than those specified in paragraph (c)(1) of this section, who could become entitled to Medicare part A on the basis of ESRD from February 1990 through April 1997 if they would file an application.

(4) The coordination periods for the following individuals ends September 30, 1998:

(i) Individuals who become entitled to Medicare part A on the basis of ESRD from May 1997 through September 1997.

(ii) Individuals who could become entitled to Medicare part A on the basis of ESRD from May 1997, through September 1997, if they would file an application.

(d) Examples. Based on the rules specified in paragraphs (b) and (c) of this section and the rules specified in §406.13 of this subchapter, the following examples illustrate how to determine, in different situations, the number of months during which Medicare is secondary payer.

(1) An individual began dialysis on November 4, 1989. He did not initiate a course in self-dialysis training nor did he receive a kidney transplant during the first 3 calendar months of dialysis. Thus, he became entitled to Medicare on February 1, 1990. Since this individual began dialysis before December 1989, the 12-month period began with the first month of dialysis, November 1989, and ended October 31, 1990. The coordination period in this case is 9 months, February 1990 through October 1990.

(2) An individual began dialysis on January 29, 1990. He did not initiate a course in self-dialysis training nor did he receive a kidney transplant during the first 3 calendar months of dialysis. Thus, he became entitled to Medicare on April 1, 1990. Since the individual began dialysis after November 1989, and became entitled to Medicare after January 1990, the coordination period began with the first month of entitlement, April 1990, and ended September 30, 1991, the end of the 18th month of entitlement.

(3) An individual began a regular course of maintenance dialysis on February 10, 1990. He did not initiate a course of self-dialysis training nor did he receive a kidney transplant during the first 3 calendar months of dialysis. Thus, he became entitled to Medicare on May 1, 1990. Medicare is secondary payer from May 1, 1990 through October 1991, a total of 18 months.

(4) The same facts exist as in the example under paragraph (d)(3), except that the individual began a course of self-dialysis training during the first 3 calendar months of dialysis. Thus, the effective date of his Medicare entitlement is February 1, 1990, and Medicare is secondary payer from February 1, 1990 through July 1991, a total of 18 months.

(5) An individual began dialysis on September 15, 1990. He did not initiate a course of self-dialysis training nor did he receive a kidney transplant during the first 3 calendar months of dialysis. Thus, he became entitled to Medicare effective December 1, 1990.
Medicare is secondary payer from December 1, 1990 through May 1992, a total of 18 months.


(7) An individual began a regular course of dialysis on December 10, 1990. He does not initiate a course of self-dialysis training nor does he receive a kidney transplant. He decides to delay his enrollment in Medicare because his employer group health plan pays charges in full and he does not wish to incur part B premiums at this time. However, in March 1992, he files for part A and part B Medicare entitlement, and stipulates that he wants his Medicare entitlement to be effective March 1, 1992 (one year later than he could have become entitled). Since this individual could have been entitled to Medicare as early as March 1, 1991, Medicare is secondary payer only from March 1, 1992 through August 1992, a period of 6 months.

(While Medicare is secondary payer for only the last 6 months of this period, the Medicare program is effectively secondary payer for the full coordination period, due to the fact that the individual delayed his Medicare enrollment on account of his employer plan pays charges in full and he does not wish to incur part B premiums at this time. However, in March 1992, he files for part A and part B Medicare entitlement, and stipulates that he wants his Medicare entitlement to be effective March 1, 1992 (one year later than he could have become entitled). Since this individual could have been entitled to Medicare as early as March 1, 1991, Medicare is secondary payer only from March 1, 1992 through August 1992, a period of 6 months.)

(8) The same facts exist as in the example under paragraph (d)(7) of this section, except that the individual defers Medicare entitlement beyond August 1992. (For purposes of this example, Medicare entitlement is not retroactive, but rather takes effect after August 1992.) There would be no period during which Medicare is secondary payer in this situation. This is because Medicare entitlement does not begin until after the 18-month period expires as specified in paragraph (c)(3)(ii) of this section. Medicare would become primary payer as of the effective date of Medicare entitlement. The employer plan is required to pay primary from December 1, 1990, through August 1992, a total of 21 months.

(9) An individual becomes entitled to Medicare on December 1, 1997. The employer plan is primary payer, and Medicare is secondary payer, from December 1, 1997, through November 30, 1998, a period of 12 months. Medicare becomes primary payer on December 1, 1998, because the extension of the coordination period from 12 to 18 months applies only to items and services furnished before October 1, 1998.


(e) [Reserved]

(f) Determinations for subsequent periods of ESRD eligibility. If an individual has more than one period of eligibility based on ESRD, a coordination period will be determined for each period of eligibility in accordance with this section.

first month of dual eligibility or entitlement, regardless of when dual eligibility or entitlement began.

(2) First month of ESRD-based eligibility or entitlement and first month of dual eligibility/entitlement after February 1992 and before August 10, 1993. Except as provided in paragraph (b)(4) of this section, if the first month of ESRD-based eligibility or entitlement and first month of dual eligibility/entitlement were after February 1992 and before August 10, 1993, Medicare—

(i) Is primary payer from the first month of dual eligibility/entitlement through August 9, 1993;

(ii) Is secondary payer from August 10, 1993, through the 18th month of ESRD-based eligibility or entitlement; and

(iii) Again becomes primary payer after the 18th month of ESRD-based eligibility or entitlement.

(3) First month of ESRD-based eligibility or entitlement after February 1992 and first month of dual eligibility/entitlement after August 9, 1993. Except as provided in paragraph (b)(4) of this section, if the first month of ESRD-based eligibility or entitlement is after February 1992, and the first month of dual eligibility/entitlement is after August 9, 1993, the rules of § 411.162(b) and (c) apply; that is, Medicare—

(i) Is secondary payer during the first 18 months of ESRD-based eligibility or entitlement; and

(ii) Becomes primary after the 18th month of ESRD-based eligibility or entitlement.

(4) Medicare continues to be primary after an aged or disabled beneficiary becomes eligible on the basis of ESRD. (i) Applicability of the rule. Medicare remains the primary payer when an individual becomes eligible for Medicare based on ESRD if all of the following conditions are met:

(A) The individual is already entitled on the basis of age or disability when he or she becomes eligible on the basis of ESRD.

(B) The MSP prohibition against "taking into account" age-based or disability-based entitlement does not apply because plan coverage was not "by virtue of current employment status" or the employer had fewer than 20 employees (in the case of the aged) or fewer than 100 employees (in the case of the disabled).

(C) The plan is paying secondary to Medicare because the plan had justifiably taken into account the age-based or disability-based entitlement.

(ii) Effect of the rule. The plan may continue to pay benefits secondary to Medicare under paragraph (b)(4)(i) of this section. However, the plan may not differentiate in the services covered and the payments made between persons who have ESRD and those who do not.

(c) Examples. (1) (Rule (b)(1).) Mr. A, who is covered by a GHP, became entitled to Medicare on the basis of ESRD in January 1992. On December 20, 1992, Mr. A attained age 65 and became entitled on the basis of age. Since prior law was still in effect (OBRA ’93 amendment was effective in August 1993), Medicare became primary payer as of December 1992, when dual entitlement began.

(2) (Rule (b)(2).) Miss B, who has GHP coverage, became entitled to Medicare on the basis of ESRD in July 1992, and also entitled on the basis of disability in June 1993. Medicare was primary payer from June 1993 through August 9, 1993, because the plan permissibly took into account the ESRD-based entitlement (ESRD was not the "sole" basis of Medicare entitlement); secondary payer from August 10, 1993, through December 1993, the 18th month of ESRD-based entitlement (the plan is no longer permitted to take into account ESRD-based entitlement that is not the "sole" basis of Medicare entitlement); and again became primary payer beginning January 1994.

(3) (Rule (b)(3).) Mr. C, who is 67 years old and entitled to Medicare on the basis of age, has GHP coverage by virtue of current employment status. Mr. C is diagnosed as having ESRD and begins a course of maintenance dialysis on June 27, 1993. Effective September 1, 1993, Mr. C is eligible for Medicare on the basis of ESRD. Medicare, which was secondary because Mr. C’s GHP coverage was by virtue of current employment, continues to be secondary payer through February 1995, the 18th month of ESRD-based eligibility, and becomes primary payer beginning March 1995.
(4) (Rule (b)(3).) Mr. D retired at age 62 and maintained GHP coverage as a retiree. In January 1994, at the age of 64, Mr. D became entitled to Medicare based on ESRD. Seven months into the 18-month coordination period (July 1994), Mr. D turned age 65. The coordination period continues without regard to age-based entitlement, with the retirement plan continuing to pay primary benefits through June 1995, the 18th month of ESRD-based entitlement. Thereafter, Medicare becomes the primary payer.

(5) (Rule (b)(3).) Mrs. E retired at age 62 and maintained GHP coverage as a retiree. In July 1994, she simultaneously became eligible for Medicare based on ESRD (maintenance dialysis began in April 1994) and entitled based on age. The retirement plan must pay benefits primary to Medicare from July 1994 through December 1995, the first 18 months of ESRD-based eligibility. Thereafter, Medicare becomes the primary payer.

(6) (Rule (b)(3).) Mr. F, who is 67 years of age, is working and has GHP coverage because of his employment status, subsequently develops ESRD, and begins a course of maintenance dialysis in October 1994. He becomes eligible for Medicare based on ESRD effective January 1, 1995. Under the working aged provision, the plan continues to pay primary to Medicare through December 1994. On January 1, 1995, the working aged provision ceases to apply and the ESRD MSP provision takes effect. In September 1995, Mr. F retires. The GHP must ignore Mr. F’s retirement status and continue to pay primary to Medicare through June 1996, the end of the 18-month coordination period.

(7) (Rule (b)(4).) Mrs. G, who is 67 years of age, is retired. She has GHP retirement coverage through her former employer. Her plan permissibly took into account her age-based Medicare entitlement when she retired and is paying benefits secondary to Medicare. Mrs. G subsequently develops ESRD and begins a course of maintenance dialysis in October 1995. She automatically becomes entitled for Medicare based on ESRD effective January 1, 1996. The plan continues to be secondary on the basis of Mrs. G’s age-based entitlement as long as the plan does not differentiate in the services it provides to Mrs. G and does not do anything else that would constitute “taking into account” her ESRD-based eligibility.

[60 FR 45369, Aug. 31, 1995; 60 FR 53876, Oct. 18, 1995]

§ 411.165 Basis for conditional Medicare payments.

(a) General rule. Except as specified in paragraph (b) of this section, the Medicare intermediary or carrier may make a conditional payment if—

(1) The beneficiary, the provider, or the supplier that has accepted assignment files a proper claim under the group health plan and the plan denies the claim in whole or in part; or

(2) The beneficiary, because of physical or mental incapacity, fails to file a proper claim.

(b) Exception. Medicare does not make conditional primary payments under either of the following circumstances:

(1) The claim is denied for one of the following reasons:

(i) It is alleged that the group health plan is secondary to Medicare.

(ii) The group health plan limits its payments when the individual is entitled to Medicare.

(iii) Failure to file a proper claim if that failure is for any reason other than the physical or mental incapacity of the beneficiary.

(2) The group health plan fails to furnish information requested by HCFA and necessary to determine whether the employer plan is primary to Medicare.


Subpart G—Special Rules: Aged Beneficiaries and Spouses Who Are Also Covered Under Group Health Plans

§ 411.170 General provisions.

(a) Basis. (1) This subpart is based on certain provisions of section 1862(b) of the Act, which impose specific requirements and limitations with respect to—
§ 411.172 Medicare benefits secondary to group health plan benefits.

(a) Conditions that the individual must meet. Medicare Part A and Part B benefits are secondary to benefits payable by a GHP for services furnished during any month in which the individual—

(1) Is aged;

(2) Is entitled to Medicare Part A benefits under § 406.10 of this chapter; and

(3) Meets one of the following conditions:

(i) Is covered under a GHP of an employer that has at least 20 employees (including a multi-employer plan in which at least one of the participating employers meets that condition), and coverage under the plan is by virtue of the individual’s current employment status.

(ii) Is the aged spouse (including a divorced or common-law spouse) of an individual (of any age) who is covered under a GHP described in paragraph (a)(3)(i) of this section by virtue of the individual’s current employment status.

(b) Special rule for multi-employer plans. The requirements and limitations of paragraph (a) of this section and of (a)(2)(iii) of § 411.170 do not apply with respect to individuals enrolled in a multi-employer plan if—

(1) The individuals are covered by virtue of current employment status with an employer that has fewer than 20 employees; and

(2) The plan requests an exception and identifies the individuals for whom it requests the exception as meeting the conditions specified in paragraph (b)(1) of this section.

(c) Refusal to accept group health plan coverage. An employee or spouse may refuse the health plan offered by the employer. If the employee or spouse refuses the plan—

(1) Medicare is primary payer for that individual; and

(2) The plan may not offer that individual coverage complementary to Medicare.

(d) Reemployed retiree or annuitant. A reemployed retiree or annuitant who is covered by a GHP and who performs...
§ 411.175 Basis for Medicare primary payments.

(a) General rule. HCFA makes Medicare primary payments for covered services that are—

(1) Furnished to Medicare beneficiaries who have declined to enroll in the GHP;

(2) Not covered by the plan for any individuals or spouses who are enrolled by virtue of the individual’s current employment status;

(3) Covered under the plan but not available to particular individuals or spouses enrolled by virtue of current employment status because they have exhausted their benefits under the plan;

(4) Furnished to individuals whose COBRA continuation coverage has been terminated because of the individual’s Medicare entitlement; or

(5) Covered under COBRA continuation coverage notwithstanding the individual’s Medicare entitlement.

(b) Conditional Medicare payments: Basic rule. Except as provided in paragraph (c) of this section, Medicare may make a conditional primary payment if—

(1) The beneficiary, the provider, or the supplier that has accepted assignment has filed a proper claim under the group health plan and the plan has denied the claim in whole or in part; or

(2) The beneficiary, because of physical or mental incapacity, failed to file proper claim.

(c) Conditional primary payments: Exception. Medicare does not make conditional primary payments under either of the following circumstances:

(1) The claim is denied for one of the following reasons:

(i) It is alleged that the group health plan is secondary to Medicare.

(ii) The plan limits its payments when the individual is entitled to Medicare.

(iii) The plan covers the services for individuals or spouses who are enrolled in the plan by virtue of current employment status and are under age 65 but not for individuals and spouses who are enrolled on the same basis but are age 65 or older.

(iv) Failure to file a proper claim if that failure is for any reason other than physical or mental incapacity of the beneficiary.

(2) The group health plan fails to furnish information requested by HCFA and necessary to determine whether

(3) Covered under the plan but not available to particular individuals or spouses enrolled by virtue of current employment status because they have exhausted their benefits under the plan;
§ 411.200 Basis.

(a) This subpart is based on certain provisions of section 1862(b) of the Act, which impose specific requirements and limitations with respect to—

(1) Individuals who are entitled to Medicare on the basis of disability; and

(2) Large group health plans (LGHPs) that cover those individuals.

(b) Under these provisions, the LGHP may not take into account the Medicare entitlement of a disabled individual who is covered (or seeks to be covered) under the plan by virtue of his or her own current employment status or that of a member of his or her family. (§ 411.108 gives examples of actions that constitute taking into account.)

§ 411.201 Definitions.

As used in this subpart—

Entitled to Medicare on the basis of disability means entitled or deemed entitled on the basis of entitlement to social security disability benefits or railroad retirement disability benefits. (§ 406.12 of this chapter explains the requirements an individual must meet in order to be entitled or deemed to be entitled to Medicare on the basis of disability.)

Family member means a person who is enrolled in an LGHP based on another person’s enrollment; for example, the enrollment of the named insured individual. Family members may include a spouse (including a divorced or common-law spouse), a natural, adopted, foster, or stepchild, a parent, or a sibling.

§ 411.204 Medicare benefits secondary to LGHP benefits.

(a) Medicare benefits are secondary to benefits payable by an LGHP for services furnished during any month in which the individual—

(1) Is entitled to Medicare Part A benefits under § 406.12 of this chapter;

(2) Is covered under an LGHP; and

(3) Has LGHP coverage by virtue of his or her own or a family member’s current employment status.

(b) Individuals entitled to Medicare on the basis of disability who are also eligible for, or entitled to, Medicare on the basis of ESRD. If a disabled individual is, or could upon filing an application become, entitled to Medicare on the basis of ESRD, the coordination of benefits rules of subpart F of this part apply.

§ 411.206 Basis for Medicare primary payments and limits on secondary payments.

(a) General rule. HCFA makes Medicare primary payments for services furnished to disabled beneficiaries covered under the LGHP by virtue of their own or a family member’s current employment status if the services are—

(1) Furnished to Medicare beneficiaries who have declined to enroll in the GHP;

(2) Not covered under the plan for the disabled individual or similarly situated individuals;

(3) Covered under the plan but not available to particular disabled individuals because they have exhausted their benefits under the plan;

(4) Furnished to individuals whose COBRA continuation coverage has been terminated because of the individual’s Medicare entitlement; or

(5) Covered under COBRA continuation coverage notwithstanding the individual’s Medicare entitlement.

(b) Conditional primary payments: Basic rule. Except as provided in paragraph (c) of this section, HCFA may make a conditional Medicare primary payment for any of the following reasons:

(1) The beneficiary, the provider, or the supplier that has accepted assignment has filed a proper claim with the LGHP and the LGHP has denied the claim in whole or in part.

(2) The beneficiary, because of physical or mental incapacity, failed to file a proper claim.
(c) Conditional primary payments: Exceptions. HCFA does not make conditional Medicare primary payments if—

(1) The LGHP denies the claim in whole or in part for one of the following reasons:
   (i) It is alleged that the LGHP is secondary to Medicare.
   (ii) The LGHP limits its payments when the individual is entitled to Medicare.
   (iii) The LGHP does not provide the benefits to individuals who are entitled to Medicare on the basis of disability and covered under the plan by virtue of current employment status but does provide the benefits to other similarly situated individuals enrolled in the plan.
   (iv) The LGHP takes into account entitlement to Medicare in any other way.
   (v) There was failure to file a proper claim for any reason other than physical or mental incapacity of the beneficiary.

(2) The LGHP, an employer or employee organization, or the beneficiary fails to furnish information that is requested by HCFA and that is necessary to determine whether the LGHP is primary to Medicare.

(d) Limit on secondary payments. The provisions of §411.172(e) also apply to services furnished to the disabled under this subpart.

Subpart I—[Reserved]

Subpart J—Physician Ownership of, and Referral of Patients or Laboratory Specimens to, Entities Furnishing Clinical Laboratory or Other Health Services

SOURCE: 60 FR 41978, Aug. 14, 1995, unless otherwise noted.

§ 411.350 Scope of subpart.

(a) This subpart implements section 1877 of the Act, which generally prohibits a physician from making a referral under Medicare for clinical laboratory services to an entity with which the physician or a member of the physician’s immediate family has a financial relationship.

(b) This subpart does not provide for exceptions or immunity from civil or criminal prosecution or other sanctions applicable under any State laws or under Federal law other than section 1877 of the Act. For example, although a particular arrangement involving a physician’s financial relationship with an entity may not prohibit the physician from making referrals to the entity under this subpart, the arrangement may nevertheless violate another provision of the Act or other laws administered by HHS, the Federal Trade Commission, the Securities and Exchange Commission, the Internal Revenue Service, or any other Federal or State agency.

(c) This subpart requires, with some exceptions, that certain entities furnishing covered items or services under Part A or Part B report information concerning their ownership, investment, or compensation arrangements in the form, manner, and at the times specified by HCFA.

§ 411.351 Definitions.

As used in this subpart, unless the context indicates otherwise:

Clinical laboratory services means the biological, microbiological, serological, chemical, immunohematological, hematological, biophysical, cytological, pathological, or other examination of materials derived from the human body for the purpose of providing information for the diagnosis, prevention, or treatment of any disease or impairment of, or the assessment of the health of, human beings. These examinations also include procedures to determine, measure, or otherwise describe the presence or absence of various substances or organisms in the body.

Compensation arrangement means any arrangement involving any remuneration, direct or indirect, between a physician (or a member of a physician’s immediate family) and an entity.

Direct supervision means supervision by a physician who is present in the office suite and immediately available to provide assistance and direction throughout the time services are being performed.

Employee means any individual who, under the usual common law rules that
Health Care Financing Administration, HHS § 411.351

apply in determining the employer-employee relationship (as applied for purposes of section 3121(d)(2) of the Internal Revenue Code of 1986), is considered to be employed by, or an employee of, an entity. (Application of these common law rules is discussed at 20 CFR 404.1007 and 26 CFR 31.3121(d)-1(c).)

Entity means a sole proprietorship, trust, corporation, partnership, foundation, not-for-profit corporation, or unincorporated association.

Fair market value means the value in arm's-length transactions, consistent with the general market value. With respect to rentals or leases, fair market value means the value of rental property for general commercial purposes (not taking into account its intended use). In the case of a lease of space, this value may not be adjusted to reflect the additional value the prospective lessee or lessor would attribute to the proximity or convenience to the lessor when the lessor is a potential source of patient referrals to the lessee.

Financial relationship refers to a direct or indirect relationship between a physician (or a member of a physician's immediate family) and an entity in which the physician or family member has—

(1) An ownership or investment interest that exists in the entity through equity, debt, or other means and includes an interest in an entity that holds an ownership or investment interest in any entity providing laboratory services; or

(2) A compensation arrangement with the entity.

Group practice means a group of two or more physicians, legally organized as a partnership, professional corporation, foundation, not-for-profit corporation, faculty practice plan, or similar association, that meets the following conditions:

(1) Each physician who is a member of the group, as defined in this section, furnishes substantially the full range of patient care services that the physician routinely furnishes including medical care, consultation, diagnosis, and treatment through the joint use of shared office space, facilities, equipment, and personnel.

(2) Except as provided in paragraphs (2)(i) and (2)(ii) of this definition, substantially all of the patient care services of the physicians who are members of the group (that is, at least 75 percent of the total patient care services of the group practice members) are furnished through the group and billed in the name of the group and the amounts received are treated as receipts of the group. "Patient care services" are measured by the total patient care time each member spends on these services. For example, if a physician practices 40 hours a week and spends 30 hours on patient care services for a group practice, the physician has spent 75 percent of his or her time providing countable patient care services.

(i) The "substantially all" test does not apply to any group practice that is located solely in an HPSA, as defined in this section, and

(ii) For group practices located outside of an HPSA (as defined in this section) any time spent by group practice members providing services in an HPSA should not be used to calculate whether the group practice located outside the HPSA has met the "substantially all" test, regardless of whether the members' time in the HPSA is spent in a group practice, clinic, or office setting.

(3) The practice expenses and income are distributed in accordance with methods previously determined. In the case of faculty practice plans associated with a hospital, institution of higher education, or medical school that has an approved medical residency training program in which faculty practice plan physicians perform specialty and professional services, both within and outside the faculty practice, as well as perform other tasks such as research, this definition applies only to those services that are furnished within the faculty practice plan. Hospital means any separate legally organized operating entity plus any subsidiary, related, or other entities that perform services for the hospital's patients and for which the hospital bills. A "hospital" does not include entities that perform services for hospital patients "under arrangements" with the hospital.
HPSA means, for purposes of this regulation, an area designated as a health professional shortage area under section 332(a)(1)(A) of the Public Health Service Act for primary medical care professionals (in accordance with the criteria specified in 42 CFR part 5, appendix A, part I—Geographic Areas). In addition, with respect to dental, mental health, vision care, podiatric, and pharmacy services, an HPSA means an area designated as a health professional shortage area under section 332(a)(1)(A) of the Public Health Service Act for dental professionals, mental health professionals, vision care professionals, podiatric professionals, and pharmacy professionals, respectively.

Immediate family member or member of a physician's immediate family means husband or wife; natural or adoptive parent, child, or sibling; stepparent, stepchild, stepbrother, or stepsister; father-in-law, mother-in-law, son-in-law, daughter-in-law, brother-in-law, or sister-in-law; grandparent or grandchild; and spouse of a grandparent or grandchild.

Laboratory means an entity furnishing biological, microbiological, serological, chemical, immunohematological, hematological, biophysical, cytological, pathological, or other examination of materials derived from the human body for the purpose of providing information for the diagnosis, prevention, or treatment of any disease or impairment of, or the assessment of the health of, human beings. These examinations also include procedures to determine, measure, or otherwise describe the presence or absence of various substances or organisms in the body. Entities only collecting or preparing specimens (or both) or only serving as a mailing service and not performing testing are not considered laboratories.

Members of the group means physician partners and full-time and part-time physician contractors and employees during the time they furnish services to patients of the group practice that are furnished through the group and are billed in the name of the group.

Patient care services means any tasks performed by a group practice member that address the medical needs of specific patients, regardless of whether they involve direct patient encounters. They can include, for example, the services of physicians who do not directly treat patients, time spent by a physician consulting with other physicians, or time spent reviewing laboratory tests.

Physician incentive plan means any compensation arrangement between an entity and a physician or physician group that may directly or indirectly have the effect of reducing or limiting services furnished with respect to individuals enrolled with the entity.

Plan of care means the establishment by a physician of a course of diagnosis or treatment (or both) for a particular patient, including the ordering of items or services.

Referral—
(1) Means either of the following:
(i) Except as provided in paragraph (2) of this definition, the request by a physician for, or ordering of, any item or service for which payment may be made under Medicare Part B, including a request for a consultation with another physician and any test or procedure ordered by or to be performed by (or under the supervision of) that other physician.
(ii) Except as provided in paragraph (2) of this definition, a request by a physician that includes the provision of laboratory services or the establishment of a plan of care by a physician that includes the provision of laboratory services.

(2) Does not include a request by a pathologist for clinical diagnostic laboratory tests and pathological examination services if—
(i) The request is part of a consultation initiated by another physician; and
(ii) The tests or services are furnished by or under the supervision of the pathologist.

Referring physician means a physician (or group practice) who makes a referral as defined in this section.

Remuneration means any payment, discount, forgiveness of debt, or other benefit made directly or indirectly, overtly or covertly, in cash or in kind, except that the following are not considered remuneration:
§ 411.355 General exceptions to referral prohibitions related to both ownership/investment and compensation.

The prohibition on referrals set forth in § 411.353 does not apply to the following types of services:

(a) Physicians’ services, as defined in § 410.20(a), that are furnished personally by (or under the personal supervision of) another physician in the same group practice as the referring physician.

(b) In-office ancillary services. Services that meet the following conditions:

(1) They are furnished personally by one of the following individuals:

(i) The referring physician.

(ii) A physician who is a member of the same group practice as the referring physician.

(iii) Individuals who are directly supervised by the referring physician or, in the case of group practices, by another physician in the same group practice as the referring physician.

(2) They are furnished in one of the following locations:

(i) A building in which the referring physician (or another physician who is a member of the same group practice) furnishes physicians’ services unrelated to the furnishing of clinical laboratory services.

(ii) A building that is used by the group practice for the provision of some or all of the group’s clinical laboratory services.

(3) They are billed by one of the following:

(1) The forgiveness of amounts owed for inaccurate tests or procedures, mistakenly performed tests or procedures, or the correction of minor billing errors.

(2) The furnishing of items, devices, or supplies that are used solely to collect, transport, process, or store specimens for the entity furnishing the items, devices, or supplies or are used solely to order or communicate the results of tests or procedures for the entity.

(3) A payment made by an insurer or a self-insured plan to a physician to satisfy a claim, submitted on a fee-for-service basis, for the furnishing of health services by that physician to an individual who is covered by a policy with the insurer or by the self-insured plan, if—

(i) The health services are not furnished, and the payment is not made, under a contract or other arrangement between the insurer or the plan and the physician;

(ii) The payment is made to the physician on behalf of the covered individual and would otherwise be made directly to the individual; and

(iii) The amount of the payment is set in advance, does not exceed fair market value, and is not determined in a manner that takes into account directly or indirectly the volume or value of any referrals.

Transaction means an instance or process of two or more persons doing business. An isolated transaction is one involving a single payment between two or more persons. A transaction that involves long-term or installment payments is not considered an isolated transaction.

§ 411.353 Prohibition on certain referrals by physicians and limitations on billing.

(a) Prohibition on referrals. Except as provided in this subpart, a physician who has a financial relationship with an entity, or who has an immediate family member who has a financial relationship with the entity, may not make a referral to that entity for the furnishing of clinical laboratory services for which payment otherwise may be made under Medicare.

(b) Limitations on billing. An entity that furnishes clinical laboratory services under a referral that is prohibited by paragraph (a) of this section may not present or cause to be presented a claim or bill to the Medicare program or to any individual, third party payer, or other entity for the clinical laboratory services performed under that referral.

(c) Denial of payment. No Medicare payment may be made for a clinical laboratory service that is furnished under a prohibited referral.

(d) Refunds. An entity that collects payment for a laboratory service that was performed under a prohibited referral must refund all collected amounts on a timely basis.
§ 411.356 Exceptions to referral prohibitions related to ownership or investment interests.

For purposes of § 411.353, the following ownership or investment interests do not constitute a financial relationship:

(a) Publicly traded securities. Ownership of investment securities (including shares of bonds, debentures, notes, or other debt instruments) that may be purchased on terms generally available to the public and that meet the requirements of paragraphs (a)(1) and (a)(2) of this section.

(i) Listed for trading on the New York Stock Exchange, the American Stock Exchange, or any regional exchange in which quotations are published on a daily basis, or foreign securities listed on a recognized foreign, national, or regional exchange in which quotations are published on a daily basis; or

(ii) Traded under an automated interdealer quotation system operated by the National Association of Securities Dealers.

(b) Mutual funds. Ownership of shares in a regulated investment company as defined in section 851(a) of the Internal Revenue Code of 1986, if the company had, at the end of its most recent fiscal year, or on average during the previous 3 fiscal years, total assets exceeding $75 million.

(c) Specific providers. Ownership or investment interest in the following entities:

(i) A laboratory that is located in a rural area (that is, a laboratory that is not located in an urban area as defined in § 412.62(f)(1) of this chapter) and that meets the following criteria:

(A) The testing that is referred by a physician who has (or whose immediate family member has) an ownership or investment interest in the rural laboratory is either—

(B) If not performed by the rural laboratory or

(C) Performed on the premises of the rural laboratory; or

(ii) Substantially all of the testing billed by the Medicare program directly for the testing.

§ 411.356 Exceptions to referral prohibitions related to ownership or investment interests.

For purposes of § 411.353, the following ownership or investment interests do not constitute a financial relationship:

(a) Publicly traded securities. Ownership of investment securities (including shares of bonds, debentures, notes, or other debt instruments) that may be purchased on terms generally available to the public and that meet the requirements of paragraphs (a)(1) and (a)(2) of this section.

(i) Listed for trading on the New York Stock Exchange, the American Stock Exchange, or any regional exchange in which quotations are published on a daily basis, or foreign securities listed on a recognized foreign, national, or regional exchange in which quotations are published on a daily basis; or

(ii) Traded under an automated interdealer quotation system operated by the National Association of Securities Dealers.

(b) Mutual funds. Ownership of shares in a regulated investment company as defined in section 851(a) of the Internal Revenue Code of 1986, if the company had, at the end of its most recent fiscal year, or on average during the previous 3 fiscal years, total assets exceeding $75 million.

(c) Specific providers. Ownership or investment interest in the following entities:

(i) A laboratory that is located in a rural area (that is, a laboratory that is not located in an urban area as defined in § 412.62(f)(1) of this chapter) and that meets the following criteria:

(A) The testing that is referred by a physician who has (or whose immediate family member has) an ownership or investment interest in the rural laboratory is either—

(B) Performed on the premises of the rural laboratory; or

(C) Performed on the premises of the rural laboratory; or

(D) If not performed by the rural laboratory or

(E) Substantially all of the testing billed by the Medicare program directly for the testing.

(ii) Substantially all of the testing billed by the Medicare program directly for the testing.

(iii) Substantially all of the testing billed by the Medicare program directly for the testing.

(iv) Substantially all of the testing billed by the Medicare program directly for the testing.

§ 411.356 Exceptions to referral prohibitions related to ownership or investment interests.

For purposes of § 411.353, the following ownership or investment interests do not constitute a financial relationship:

(a) Publicly traded securities. Ownership of investment securities (including shares of bonds, debentures, notes, or other debt instruments) that may be purchased on terms generally available to the public and that meet the requirements of paragraphs (a)(1) and (a)(2) of this section.

(i) Listed for trading on the New York Stock Exchange, the American Stock Exchange, or any regional exchange in which quotations are published on a daily basis, or foreign securities listed on a recognized foreign, national, or regional exchange in which quotations are published on a daily basis; or

(ii) Traded under an automated interdealer quotation system operated by the National Association of Securities Dealers.

(b) Mutual funds. Ownership of shares in a regulated investment company as defined in section 851(a) of the Internal Revenue Code of 1986, if the company had, at the end of its most recent fiscal year, or on average during the previous 3 fiscal years, total assets exceeding $75 million.

(c) Specific providers. Ownership or investment interest in the following entities:

(i) A laboratory that is located in a rural area (that is, a laboratory that is not located in an urban area as defined in § 412.62(f)(1) of this chapter) and that meets the following criteria:

(A) The testing that is referred by a physician who has (or whose immediate family member has) an ownership or investment interest in the rural laboratory is either—

(B) Performed on the premises of the rural laboratory; or

(C) If not performed by the rural laboratory or

(D) Substantially all of the testing billed by the Medicare program directly for the testing.

(ii) Substantially all of the testing billed by the Medicare program directly for the testing.

(iii) Substantially all of the testing billed by the Medicare program directly for the testing.

(iv) Substantially all of the testing billed by the Medicare program directly for the testing.

§ 411.356 Exceptions to referral prohibitions related to ownership or investment interests.

For purposes of § 411.353, the following ownership or investment interests do not constitute a financial relationship:

(a) Publicly traded securities. Ownership of investment securities (including shares of bonds, debentures, notes, or other debt instruments) that may be purchased on terms generally available to the public and that meet the requirements of paragraphs (a)(1) and (a)(2) of this section.

(i) Listed for trading on the New York Stock Exchange, the American Stock Exchange, or any regional exchange in which quotations are published on a daily basis, or foreign securities listed on a recognized foreign, national, or regional exchange in which quotations are published on a daily basis; or

(ii) Traded under an automated interdealer quotation system operated by the National Association of Securities Dealers.

(b) Mutual funds. Ownership of shares in a regulated investment company as defined in section 851(a) of the Internal Revenue Code of 1986, if the company had, at the end of its most recent fiscal year, or on average during the previous 3 fiscal years, total assets exceeding $75 million.

(c) Specific providers. Ownership or investment interest in the following entities:

(i) A laboratory that is located in a rural area (that is, a laboratory that is not located in an urban area as defined in § 412.62(f)(1) of this chapter) and that meets the following criteria:

(A) The testing that is referred by a physician who has (or whose immediate family member has) an ownership or investment interest in the rural laboratory is either—

(B) Performed on the premises of the rural laboratory; or

(C) If not performed by the rural laboratory or

(D) Substantially all of the testing billed by the Medicare program directly for the testing.

(ii) Substantially all of the testing billed by the Medicare program directly for the testing.

(iii) Substantially all of the testing billed by the Medicare program directly for the testing.

(iv) Substantially all of the testing billed by the Medicare program directly for the testing.
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(2) A hospital that is located in Puerto Rico.

(3) A hospital that is located outside of Puerto Rico if one of the following conditions is met:
   (i) The referring physician is authorized to perform services at the hospital, and the physician's ownership or investment interest is in the entire hospital and not merely in a distinct part or department of the hospital.
   (ii) Until January 1, 1995, the referring physician's ownership or investment interest does not relate (directly or indirectly) to the furnishing of clinical laboratory services.

§411.357 Exceptions to referral prohibitions related to compensation arrangements.

For purposes of §411.353, the following compensation arrangements do not constitute a financial relationship:

(a) Rental of office space. Payments for the use of office space made by a lessee to a lessor if there is a rental or lease agreement that meets the following requirements:
   (1) The agreement is set out in writing and is signed by the parties and specifies the premises covered by the lease.
   (2) The term of the agreement is at least 1 year.
   (3) The space rented or leased does not exceed that which is reasonable and necessary for the legitimate business purposes of the lease or rental and is used exclusively by the lessee when being used by the lessee, except that the lessee may make payments for the use of space consisting of common areas if the payments do not exceed the lessee's pro rata share of expenses for the space based upon the ratio of the space used exclusively by the lessee to the total amount of space (other than common areas) occupied by all persons using the common areas.
   (4) The rental charges over the term of the lease are set in advance and are consistent with fair market value.
   (5) The charges are not determined in a manner that takes into account the volume or value of any referrals or other business generated between the parties.
   (6) The agreement would be commercially reasonable even if no referrals were made between the lessee and the lessor.

(b) Rental of equipment. Payments made by a lessee to a lessor for the use of equipment under the following conditions:
   (1) A rental or lease agreement is set out in writing and signed by the parties and specifies the equipment covered by the lease.
   (2) The equipment rented or leased does not exceed that which is reasonable and necessary for the legitimate business purposes of the lease or rental and is used exclusively by the lessee when being used by the lessee.
   (3) The lease provides for a term of rental or lease of at least 1 year.
   (4) The rental charges over the term of the lease are set in advance, are consistent with fair market value, and are not determined in a manner that takes into account the volume or value of any referrals or other business generated between the parties.
   (5) The lease would be commercially reasonable even if no referrals were made between the parties.

(c) Bona fide employment relationships. Any amount paid by an employer to a physician (or immediate family member) who has a bona fide employment relationship with the employer for the provision of services if the following conditions are met:
   (1) The employment is for identifiable services.
   (2) The amount of the remuneration under the employment is—
      (i) Consistent with the fair market value of the services; and
      (ii) Except as provided in paragraph (c)(4) of this section, is not determined in a manner that takes into account (directly or indirectly) the volume or value of any referrals by the referring physician.
   (3) The remuneration is provided under an agreement that would be commercially reasonable even if no referrals were made to the employer.
   (4) Paragraph (c)(2)(ii) of this section does not prohibit payment of remuneration in the form of a productivity bonus based on services performed personally by the physician (or immediate family member of the physician).

(d) Personal service arrangements.

   (1) General. Remuneration from an entity
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under an arrangement to a physician or immediate family member of the physician, including remuneration for specific physicians’ services furnished to a nonprofit blood center, if the following conditions are met:

(i) The arrangement is set out in writing, is signed by the parties, and specifies the services covered by the arrangement.

(ii) The arrangement covers all of the services to be furnished by the physician (or an immediate family member of the physician) to the entity.

(iii) The aggregate services contracted for do not exceed those that are reasonable and necessary for the legitimate business purposes of the arrangement.

(iv) The term of the arrangement is for at least 1 year.

(v) The compensation to be paid over the term of the arrangement is set in advance, does not exceed fair market value, and, except in the case of a physician incentive plan, is not determined in a manner that takes into account the volume or value of any referrals or other business generated between the parties.

(vi) The services to be furnished under the arrangement do not involve the counseling or promotion of a business arrangement or other activity that violates any State or Federal law.

(2) Physician incentive plan exception. In the case of a physician incentive plan between a physician and an entity, the compensation may be determined in a manner (through a withhold, capitation, bonus, or otherwise) that takes into account directly or indirectly the volume or value of any referrals the physician generates for the hospital, if the plan meets the following requirements:

(i) No specific payment is made directly or indirectly under the plan to a physician or a physician group as an inducement to reduce or limit medically necessary services furnished with respect to a specific individual enrolled in the entity.

(ii) In the case of a plan that places a physician or a physician group at substantial financial risk as determined by the Secretary under section 1877(e)(1)(A)(ii) of the Act, the plan complies with any requirements the Secretary has imposed under that section.

(iii) Upon request by the Secretary, the entity provides the Secretary with access to descriptive information regarding the plan, in order to permit the Secretary to determine whether the plan is in compliance with the requirements of paragraph (d)(2) of this section.

(3) Until January 1, 1995, the provisions in paragraphs (d)(1) and (2) of this section do not apply to any arrangements that meet the requirements of section 1877(e)(2) or section 1877(e)(3) of the Act as they read before they were amended by the Omnibus Budget Reconciliation Act of 1993 (Public Law 103-66).

(e) Physician recruitment. Remuneration provided by a hospital to recruit a physician that is intended to induce the physician to relocate to the geographic area served by the hospital in order to become a member of the hospital’s medical staff, if all of the following conditions are met:

(1) The arrangement and its terms are in writing and signed by both parties.

(2) The arrangement is not conditioned on the physician’s referral of patients to the hospital.

(3) The hospital does not determine (directly or indirectly) the amount or value of the remuneration to the physician based on the volume or value of any referrals the physician generates for the hospital.

(4) The physician is not precluded from establishing staff privileges at another hospital or referring business to another entity.

(f) Isolated transactions. Isolated financial transactions, such as a, one-time sale of property or a practice, if all of the conditions set forth in paragraphs (c)(2) and (c)(3) of this section are met with respect to an entity in the same manner as they apply to an employer. There can be no additional transactions between the parties for 6 months after the isolated transaction, except for transactions which are specifically excepted under the other provisions in §§ 411.355 through 411.357.

(g) Arrangements with hospitals. (1) Until January 1, 1996, any compensation arrangement between a hospital
§ 411.360 Group practice attestation.

(a) Except as provided in paragraph (b) of this section, a group practice (as defined in section 1877(h)(4) of the Act and §411.351) must submit a written statement to its carrier annually to attest that, during the most recent 12-month period (calendar year, fiscal year, or immediately preceding 12-month period) 75 percent of the total patient care services of group practice members was furnished through the group, was billed under a billing number assigned to the group, and the amounts so received were treated as receipts of the group.

(b) A newly-formed group practice (one in which physicians have recently begun to practice together) or any group practice that has been unable in the past to meet the requirements of section 1877(h)(4) of the Act must—

(1) Submit a written statement to attest that, during the next 12-month period (calendar year, fiscal year, or next 12 months), it expects to meet the 75-percent standard and will take measures to ensure the standard is met; and

(2) At the end of the 12-month period, submit a written statement to attest that it met the 75-percent standard during that period, billed for those services under a billing number assigned to the group, and treated amounts received for those services as receipts of the group. If the group did not meet the standard, any Medicare payments made for clinical laboratory services furnished by the group during the 12-month period that were conditioned upon the standard being met are overpayments.

(c) Once any group has chosen whether to use its fiscal year, the calendar year, or some other 12-month period, the group practice must adhere to this choice.

(d) The attestation must contain a statement that the information furnished in the attestation is true and accurate and must be signed by a group representative.

(e) A group that intends to meet the definition of a group practice in order to qualify for an exception described in §§411.355 through 411.357, must submit the attestation required by paragraph (a) or paragraph (b)(1) of this section, as applicable, to its carrier no later
than 60 days after receipt of the attestation instructions from its carrier.

§ 411.361 Reporting requirements.

(a) Basic rule. Except as provided in paragraph (b) of this section, all entities furnishing items or services for which payment may be made under Medicare must submit information to HCFA concerning their financial relationships (as defined in paragraph (d) of this section), in such form, manner, and at such times as HCFA specifies.

(b) Exception. The requirements of paragraph (a) of this section do not apply to entities that provide 20 or fewer Part A and Part B items and services during a calendar year, or to designated health services provided outside the United States.

(c) Required information. The information submitted to HCFA under paragraph (a) of this section must include at least the following:

(1) The name and unique physician identification number (UPIN) of each physician who has a financial relationship with the entity;

(2) The name and UPIN of each physician who has an immediate relative (as defined in § 411.351) who has a financial relationship with the entity;

(3) The covered items and services provided by the entity; and

(4) With respect to each physician identified under paragraphs (c)(1) and (c)(2) of this section, the nature of the financial relationship (including the extent and/or value of the ownership or investment interest or the compensation arrangement, if requested by HCFA).

(d) Reportable financial relationships. For purposes of this section, a financial relationship is any ownership or investment interest or any compensation arrangement, as described in section 1877 of the Act.

(e) Form and timing of reports. Entities that are subject to the requirements of this section must submit the required information on a HCFA-prescribed form within the time period specified by the servicing carrier or intermediary. Entities are given at least 30 days from the date of the carrier's or intermediary's request to provide the initial information. Thereafter, an entity must provide updated information within 60 days from the date of any change in the submitted information. Entities must retain documentation sufficient to verify the information provided on the forms and, upon request, must make that documentation available to HCFA or the OIG.

(f) Consequences of failure to report. Any person who is required, but fails, to submit information concerning his or her financial relationships in accordance with this section is subject to a civil money penalty of up to $10,000 for each day of the period beginning on the day following the applicable deadline established under paragraph (e) of this section until the information is submitted. Assessment of these penalties will comply with the applicable provisions of part 1003 of this title.

(g) Public disclosure. Information furnished to HCFA under this section is subject to public disclosure in accordance with the provisions of part 401 of this chapter.

§ 411.370 Advisory opinions relating to physician referrals.

(a) Period during which HCFA will accept requests. The provisions of §§ 411.370 through 411.389 apply to requests for advisory opinions that are submitted to HCFA after November 3, 1997, and before August 21, 2000, and to any requests submitted during any other time period during which HCFA is required by law to issue the advisory opinions described in this subpart.

(b) Matters that qualify for advisory opinions and who may request one. Any individual or entity may request a written advisory opinion from HCFA concerning whether a physician's referral relating to designated health services (other than clinical laboratory services) is prohibited under section 1877 of the Act. In the advisory opinion, HCFA determines whether a business arrangement described by the parties to that arrangement appears to constitute a “financial relationship” (as defined in section 1877(a)(2) of the Act) that could potentially restrict a physician's referrals, and whether the arrangement or the designated health services at issue appear to qualify for any of the exceptions to the referral arrangement.
§ 411.372 Procedure for submitting a request.

(a) Format for a request. A party or parties must submit a request for an advisory opinion to HCFA in writing, including an original request and 2 copies. The request must be addressed to: Health Care Financing Administration, Department of Health and Human Services, Attention: Advisory Opinions, P.O. Box 26505, Baltimore, MD 21207.

(b) Information HCFA requires with all submissions. The request must include the following:
(1) The name, address, telephone number, and Taxpayer Identification Number of the requestor.
(2) The names and addresses, to the extent known, of all other actual and potential parties to the arrangement that is the subject of the request.
(3) The name, title, address, and daytime telephone number of a contact person who will be available to discuss the request with HCFA on behalf of the requestor.
(4) A complete and specific description of all relevant information bearing on the arrangement, including—
(i) A complete description of the arrangement that the requestor is undertaking, or plans to undertake, including: the purpose of the arrangement; the nature of each party's (including each entity's) contribution to the arrangement; the direct or indirect relationships between the parties, with an emphasis on the relationships between physicians involved in the arrangement (or their immediate family members who are involved) and any entities that provide designated health services; the types of services for which a physician wishes to refer, and whether the referrals will involve Medicare or Medicaid patients;
(ii) Complete copies of all relevant documents or relevant portions of documents that affect or could affect the arrangement, such as personal services...
or employment contracts, leases, deeds, pension or insurance plans, financial statements, or stock certificates (or, if these relevant documents do not yet exist, a complete description, to the best of the requestor’s knowledge, of what these documents are likely to contain); (iii) Detailed statements of all collateral or oral understandings, if any; and (iv) Descriptions of any other arrangements or relationships that could affect HCFA’s analysis.

(5) Complete information on the identity of all entities involved either directly or indirectly in the arrangement, including their names, addresses, legal form, ownership structure, nature of the business (products and services) and, if relevant, their Medicare and Medicaid provider numbers. The requestor must also include a brief description of any other entities that could affect the outcome of the opinion, including those with which the requestor, the other parties, or the immediate family members of involved physicians, have any financial relationships (either direct or indirect, and as defined in section 1877(a)(2) of the Act and §411.351), or in which any of the parties holds an ownership or control interest as defined in section 1124(a)(3) of the Act.

(6) A discussion of the specific issues or questions the requestor would like HCFA to address including, if possible, a description of why the requestor believes the referral prohibition in section 1877 of the Act might or might not be triggered by the arrangement and which, if any, exceptions to the prohibition the requestor believes might apply. The requestor should attempt to designate which facts are relevant to each issue or question raised in the request and should cite the provisions of law under which each issue or question arises.

(7) An indication of whether the parties involved in the request have also asked for or are planning to ask for an advisory opinion on the arrangement in question from the OIG under section 1128D(b) of the Act (42 U.S.C. 1320a-7d(b)) and whether the arrangement is or is not, to the best of the requestor’s knowledge, the subject of an investigation.

(8) The certification(s) described in §411.373. The certification(s) must be signed by— (i) The requestor, if the requestor is an individual; (ii) The chief executive officer, or comparable officer, of the requestor, if the requestor is a corporation; (iii) The managing partner of the requestor, if the requestor is a partnership; or (iv) A managing member, if the requestor is a limited liability company.

(9) A check or money order payable to HCFA in the amount described in §411.375(a).

(c) Additional information HCFA might require. If the request does not contain all of the information required by paragraph (b) of this section, or, if either before or after accepting the request, HCFA believes it needs more information in order to render an advisory opinion, it may request whatever additional information or documents it deems necessary. Additional information must be provided in writing, signed by the same person who signed the initial request (or by an individual in a comparable position), and be certified as described in §411.373.
§ 411.375 Fees for the cost of advisory opinions.

(a) Initial payment. Parties must include with each request for an advisory opinion submitted through December 31, 1998, a check or money order payable to HCFA for $250. For requests submitted after this date, parties must include a check or money order in this amount, unless HCFA has revised the amount of the initial fee in a program issuance, in which case, the requestor must include the revised amount. This initial payment is nonrefundable.

(b) How costs are calculated. Before issuing the advisory opinion, HCFA calculates the costs the Department has incurred in responding to the request. The calculation includes the costs of salaries, benefits, and overhead for analysts, attorneys, and others who have worked on the request, as well as administrative and supervisory support for these individuals.

(c) Agreement to pay all costs. (1) By submitting the request for an advisory opinion, the requestor agrees, except as indicated in paragraph (c)(3) of this section, to pay all costs the Department incurs in responding to the request for an advisory opinion.

(2) In its request for an advisory opinion, the requestor may designate a triggering dollar amount. If HCFA estimates that the costs of processing the advisory opinion request have reached or are likely to exceed the designated triggering dollar amount, HCFA notifies the requestor.

(3) If HCFA notifies the requestor that the actual or estimated cost of processing the request has reached or is likely to exceed the triggering dollar amount, HCFA stops processing the request until the requestor makes a written request for HCFA to continue. If HCFA is delayed in processing the request for an advisory opinion because of this procedure, the time within which HCFA must issue an advisory opinion is suspended until the requestor asks HCFA to continue working on the request.

(4) If the requestor chooses not to pay for HCFA to complete an advisory opinion, or withdraws the request, the requestor is still obligated to pay for all costs HCFA has identified as costs it incurred in processing the request for an advisory opinion, up to that point.

(5) If the costs HCFA has incurred in responding to the request are greater than the amount the requestor has paid, HCFA, before issuing the advisory opinion, notifies the requestor of any additional amount that is due. HCFA does not issue an advisory opinion until the requestor has paid the full amount that is owed. Once the requestor has paid HCFA the total amount due for the costs of processing the request, HCFA issues the advisory opinion. The time period HCFA has for issuing advisory opinions is suspended from the time HCFA notifies the requestor of the amount owed until the time HCFA receives full payment.

(d) Fees for outside experts. (1) In addition to the fees identified in this section, the requestor also must pay any required fees for expert opinions, if any, from outside sources, as described in §411.377.

(2) The time period for issuing an advisory opinion is suspended from the time that HCFA notifies the requestor that it needs an outside expert opinion until the time HCFA receives that opinion.

§ 411.377 Expert opinions from outside sources.

(a) HCFA may request expert advice from qualified sources if HCFA believes that the advice is necessary to respond to a request for an advisory opinion. For example, HCFA may require the use of accountants or business experts to assess the structure of a complex business arrangement or to ascertain a physician’s or immediate family member’s financial relationship with entities that provide designated health services.

(b) If HCFA determines that it needs to obtain expert advice in order to issue a requested advisory opinion, HCFA notifies the requestor of that
§411.378 Withdrawing a request.

The party requesting an advisory opinion may withdraw the request before HCFA issues a formal advisory opinion. This party must submit the withdrawal in writing to the same address as the request, as indicated in §411.372(a). Even if the party withdraws the request, the party must pay the costs the Department has expended in processing the request, as discussed in §411.375(c).

[63 FR 1657, Jan. 9, 1998]

§411.379 When HCFA accepts a request.

(a) Upon receiving a request for an advisory opinion, HCFA promptly makes an initial determination of whether the request includes all of the information it will need to process the request.

(b) Within 15 working days of receiving the request, HCFA—

(1) Formally accepts the request for an advisory opinion;

(2) Notifies the requestor about the additional information it needs, or

(3)Declines to formally accept the request.

(c) If the requestor provides the additional information HCFA has requested, or otherwise resubmits the request, HCFA processes the resubmission in accordance with paragraphs (a) and (b) of this section as if it were an initial request for an advisory opinion.

(d) Upon accepting the request, HCFA notifies the requestor by regular U.S. mail of the date that HCFA formally accepted the request.

(e) The 90-day period that HCFA has to issue an advisory opinion set forth in §411.380(c) does not begin until HCFA has formally accepted the request for an advisory opinion.

[63 FR 1657, Jan. 9, 1998]

§411.380 When HCFA issues a formal advisory opinion.

(a) HCFA considers an advisory opinion to be issued once it has received payment and once the opinion has been dated, numbered, and signed by an authorized HCFA official.

(b) An advisory opinion contains a description of the material facts known to HCFA that relate to the arrangement that is the subject of the advisory opinion, and states HCFA’s opinion about the subject matter of the request based on those facts. If necessary, HCFA includes in the advisory opinion material facts that could be considered confidential information or trade secrets within the meaning of 18 U.S.C. 1065.

(c)(1) HCFA issues an advisory opinion, in accordance with the provisions of this part, within 90 days after it has formally accepted the request for an advisory opinion, or, for requests that HCFA determines, in its discretion, involve complex legal issues or highly complicated fact patterns, within a reasonable time period.

(2) If the 90th day falls on a Saturday, Sunday, or Federal holiday, the time period ends at the close of the first business day following the weekend or holiday;

(3) The 90-day period is suspended from the time HCFA—

(i) Notifies the requestor that the costs have reached or are likely to exceed the triggering amount as described in §411.375(c)(2) until HCFA receives written notice from the requestor to continue processing the request;

(ii) Requests additional information from the requestor until HCFA receives the additional information;
§ 411.382 HCFA's right to rescind advisory opinions.

Any advice HCFA gives in an opinion does not prejudice its right to reconsider the questions involved in the opinion and, if it determines that it is in the public interest, to rescind or revoke the opinion. HCFA provides notice to the requestor of its decision to rescind or revoke the opinion so that the requestor and the parties involved in the requestor's arrangement may discontinue any course of action they have taken in accordance with the advisory opinion. HCFA does not proceed against the requestor with respect to any action the requestor and the involved parties have taken in good faith reliance upon HCFA's advice under this part, provided—

(a) The requestor presented to HCFA a full, complete and accurate description of all the relevant facts; and
(b) The parties promptly discontinue the action upon receiving notice that HCFA had rescinded or revoked its approval, or discontinue the action within a reasonable "wind down" period, as determined by HCFA.

[63 FR 1657, Jan. 9, 1998]

§ 411.384 Disclosing advisory opinions and supporting information.

(a) Advisory opinions that HCFA issues and releases in accordance with the procedures set forth in this subpart are available to the public.

(b) Promptly after HCFA issues an advisory opinion and releases it to the requestor, HCFA makes available a copy of the advisory opinion for public inspection during its normal hours of operation and on the DHHS/HCFA web site.

(c) Any predecisional document, or part of such predecisional document, that is prepared by HCFA, the Department of Justice, or any other Department or agency of the United States in connection with an advisory opinion request under the procedures set forth in this part is exempt from disclosure under 5 U.S.C. 552, and will not be made publicly available.

(d) Documents submitted by the requestor to HCFA in connection with a request for an advisory opinion are available to the public to the extent they are required to be made available by 5 U.S.C. 552, through procedures set forth in 45 CFR part 5.

(e) Nothing in this section limits HCFA's obligation, under applicable laws, to publicly disclose the identity of the requesting party or parties, and the nature of the action HCFA has taken in response to the request.

[63 FR 1657, Jan. 9, 1998]

§ 411.386 HCFA's advisory opinions as exclusive.

The procedures described in this subpart constitute the only method by which any individuals or entities can obtain a binding advisory opinion on the subject of a physician's referrals, as described in § 411.370. HCFA has not and does not issue a binding advisory opinion on the subject matter in § 411.370, in either oral or written form, except through written opinions it issues in accordance with this subpart.

[63 FR 1658, Jan. 9, 1998]

§ 411.387 Parties affected by advisory opinions.

An advisory opinion issued by HCFA does not apply in any way to any individual or entity that does not join in the request for the opinion. Individuals or entities other than the requestor(s) may not rely on an advisory opinion.

[63 FR 1658, Jan. 9, 1998]

§ 411.388 When advisory opinions are not admissible evidence.

The failure of a party to seek or to receive an advisory opinion may not be introduced into evidence to prove that
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the party either intended or did not intend to violate the provisions of sections 1128, 1128A or 1128B of the Act. [63 FR 1658, Jan. 9, 1998]

§ 411.389 Range of the advisory opinion.

(a) An advisory opinion states only HCFA’s opinion regarding the subject matter of the request. If the subject of an advisory opinion is an arrangement that must be approved by or is regulated by any other agency, HCFA’s advisory opinion cannot be read to indicate HCFA’s views on the legal or factual issues that may be raised before that agency.

(b) An advisory opinion that HCFA issues under this part does not bind or obligate any agency other than the Department. It does not affect the requester’s, or anyone else’s, obligations to any other agency, or under any statutory or regulatory provision other than that which is the specific subject matter of the advisory opinion. [63 FR 1658, Jan. 9, 1998]

Subpart K—Payment for Certain Excluded Services

§ 411.400 Payment for custodial care and services not reasonable and necessary.

(a) Conditions for payment. Notwithstanding the exclusions set forth in §411.15 (g) and (k), Medicare pays for “custodial care” and “services not reasonable and necessary” if the following conditions are met:

(1) The services were furnished by a provider or by a practitioner or supplier that had accepted assignment of benefits for those services.

(2) Neither the beneficiary nor the provider, practitioner, or supplier knew, or could reasonably have been expected to know, that the services were excluded from coverage under §411.15 (g) or (k).

(b) Time limits on payment. (1) Basic rule. Except as provided in paragraph (b)(2) of this section, payment may not be made for inpatient hospital care, posthospital SNF care, or home health services furnished after the earlier of the following:

(i) The day on which the beneficiary has been determined, under §411.404, to have knowledge, actual or imputed, that the services were excluded from coverage by reason of §411.15(g) or §411.15(k).

(ii) The day on which the provider has been determined, under §411.406 to have knowledge, actual or imputed, that the services are excluded from coverage by reason of §411.15(g) or §411.15(k).

(2) Exception. Payment may be made for services furnished during the first day after the limit established in paragraph (b)(1) of this section, if the PRO or the intermediary determines that the additional period of one day is necessary for planning post-discharge care. If the PRO or the intermediary determines that yet another day is necessary for planning post-discharge care, payment may be made for services furnished during the second day after the limit established in paragraph (b)(1) of this section.

§ 411.402 Indemnification of beneficiary.

(a) Conditions for indemnification. If Medicare payment is precluded because the conditions of §411.400(a)(2) are not met, Medicare indemnifies the beneficiary (and recovers from the provider, practitioner, or supplier), if the following conditions are met:

(1) The beneficiary paid the provider, practitioner, or supplier some or all of the charges for the excluded services.

(2) The beneficiary did not know and could not reasonably have been expected to know that the services were not covered.

(3) The provider, practitioner, or supplier knew, or could reasonably have been expected to know that the services were not covered.

(4) The beneficiary files a proper request for indemnification before the end of the sixth month after whichever of the following is later:

(i) The month in which the beneficiary paid the provider, practitioner, or supplier.

(ii) The month in which the intermediary or carrier notified the beneficiary (or someone on his or her behalf) that the beneficiary would not be liable for the services.
§ 411.406 Criteria for determining that a provider, practitioner, or supplier knew that services were excluded from coverage as custodial care or as not reasonable and necessary.

(a) Basic rule. A provider, practitioner, or supplier that furnished services which constitute custodial care under §411.15(g) or that are not reasonable and necessary under §411.15(k), is considered to have known that the services were not covered if any one of the conditions specified in paragraphs (b) through (e) of this section is met.

(b) Notice from the PRO, intermediary or carrier. The PRO, intermediary, or carrier had informed the provider, practitioner, or supplier that the services furnished were not covered, or that similar or reasonably comparable services were not covered.

(c) Notice from the utilization review committee or the beneficiary’s attending physician. The utilization review group or committee for the provider or the beneficiary’s attending physician had informed the provider that these services were not covered.

(d) Notice from the provider, practitioner, or supplier to the beneficiary. Before the services were furnished, the provider, practitioner or supplier informed the beneficiary that—

(1) The services were not covered; or
(2) The beneficiary no longer needed covered services.

(e) Knowledge based on experience, actual notice, or constructive notice. It is clear that the provider, practitioner, or supplier could have been expected to have known that the services were excluded from coverage on the basis of the following:

(1) Its receipt of HCFA notices, including manual issuances, bulletins, or other written guides or directives from intermediaries, carriers, or PROs, including notification of PRO screening criteria specific to the condition of the beneficiary for whom the furnished services are at issue and of medical procedures subject to preadmission review by a PRO.
§411.408 Refunds of amounts collected for physician services not reasonable and necessary, payment not accepted on an assignment-related basis.

(a) Basic rule. Except as provided in paragraph (d) of this section, a physician who furnishes a beneficiary services for which the physician does not undertake to claim payment on an assignment-related basis must refund any amounts collected from the beneficiary for services otherwise covered if Medicare payment is denied because the services are found to be not reasonable and necessary under §411.15(k).

(b) Time limits for making refunds. A timely refund of any incorrectly collected amounts of money must be made to the beneficiary to whom the services were furnished. A refund is timely if—

(1) A physician who does not request a review within 30 days after receipt of the denial notice makes the refund within that time period; or

(2) A physician who files a request for review within 30 days after receipt of the denial notice makes the refund within 15 days after receiving notice of an initial adverse review determination, whether or not the physician further appeals the initial adverse review determination.

(c) Notices and appeals. If payment is denied for nonassignment-related claims because the services are found to be not reasonable and necessary, a notice of denial will be sent to both the physician and the beneficiary. The physician who does not accept assignment will have the same rights as a physician who submits claims on an assignment-related basis, as detailed in subpart H of part 405 and subpart B of part 473, to appeal the determination, and will be subject to the same time limitations.

(d) When a refund is not required. A refund of any amounts collected for services not reasonable and necessary is not required if—

(1) The physician did not know, and could not reasonably have been expected to know, that Medicare would not pay for the service; or

(2) Before the service was provided—

(i) The physician informed the beneficiary, or someone acting on the beneficiary’s behalf, in writing that the physician believed Medicare was likely to deny payment for the specific service; and

(ii) The beneficiary (or someone eligible to sign for the beneficiary under §424.36(b) of this chapter) signed a statement agreeing to pay for that service.

(e) Criteria for determining that a physician knew that services were excluded as not reasonable and necessary. A physician will be determined to have known that furnished services were excluded from coverage as not reasonable and necessary if one or more of the conditions in §411.406 of this subpart are met.

(f) Acceptable evidence of prior notice to a beneficiary that Medicare was likely to deny payment for a particular service. To qualify for waiver of the refund requirement under paragraph (d)(2) of this section, the physician must inform the beneficiary (or person acting on his or her behalf) that the physician believes Medicare is likely to deny payment.

(1) The notice must—

(i) Be in writing, using approved notice language;

(ii) Cite the particular service or services for which payment is likely to be denied; and

(iii) Cite the physician’s reasons for believing Medicare payment will be denied.

(2) The notice is not acceptable evidence if—

(i) The physician routinely gives this notice to all beneficiaries for whom he or she furnishes services; or

(ii) The notice is no more than a statement to the effect that there is a possibility that Medicare may not pay for the service.

(g) Applicability of sanctions to physicians who fail to make refunds under this
A physician who knowingly and willfully fails to make refunds as required by this section may be subject to sanctions as provided for in chapter V, parts 1001, 1002, and 1003 of this title.

[55 FR 24568, June 18, 1990; 55 FR 35142, 35143, Aug. 28, 1990]

PART 412—PROSPECTIVE PAYMENT SYSTEMS FOR INPATIENT HOSPITAL SERVICES

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AUTHORITY: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh).

SOURCE: 50 FR 12741, Mar. 29, 1985, unless otherwise noted.

Subpart A—General Provisions

§ 412.2 Basis of payment.

(a) Payment on a per discharge basis. Under both the inpatient operating and inpatient capital-related prospective payment systems, hospitals are paid a predetermined amount per discharge for inpatient hospital services furnished to Medicare beneficiaries. The
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prospective payment rate for each discharge (as defined in § 412.4) is determined according to the methodology described in subpart D, E, or G of this part, as appropriate, for operating costs, and according to the methodology described in subpart M of this part for capital-related costs. An additional payment is made for both inpatient operating and inpatient capital-related costs, in accordance with subpart F of this part, for cases that are extraordinarily costly to treat.

(b) Payment in full. (1) The prospective payment amount paid for inpatient hospital services is the total Medicare payment for the inpatient operating costs (as described in paragraph (c) of this section) and the inpatient capital-related costs (as described in paragraph (d) of this section) incurred in furnishing services covered by the Medicare program.

(2) The full prospective payment amount, as determined under subpart D, E, or G and under subpart M of this part, is made for each stay during which there is at least one Medicare payable day of care. Payable days of care, for purposes of this paragraph include the following:

(i) Limitation of liability days payable under the payment procedures for custodial care and services that are not reasonable and necessary as specified in § 411.400 of this chapter.

(ii) Guarantee of payment days, as authorized under § 409.68 of this chapter, for inpatient hospital services furnished to an individual whom the hospital has reason to believe is entitled to Medicare benefits at the time of admission.

(c) Inpatient operating costs. The prospective payment system provides a payment amount for inpatient operating costs, including—

(1) Operating costs for routine services (as described in § 413.53(b) of this chapter), such as the costs of room, board, and routine nursing services;

(2) Operating costs for ancillary services, such as radiology and laboratory services furnished to hospital inpatients;

(3) Special care unit operating costs (intensive care type unit services, as described in § 413.53(b) of this chapter);

(4) Malpractice insurance costs related to services furnished to inpatients; and

(5) Preadmission services otherwise payable under Medicare Part B furnished to a beneficiary during the 3 calendar days immediately preceding the date of the beneficiary’s admission to the hospital that meet the following conditions:

(i) The services are furnished by the hospital or by an entity wholly owned or operated by the hospital. An entity is wholly owned by the hospital if the hospital is the sole owner of the entity. An entity is wholly operated by a hospital if the hospital has exclusive responsibility for conducting and overseeing the entity’s routine operations, regardless of whether the hospital also has policymaking authority over the entity.

(ii) For services furnished after January 1, 1991, the services are diagnostic (including clinical diagnostic laboratory tests).

(iii) For services furnished on or after October 1, 1991, the services are furnished in connection with the principal diagnosis that requires the beneficiary to be admitted as an inpatient and are not the following:

(A) Ambulance services.

(B) Maintenance renal dialysis.

(d) Inpatient capital-related costs. For cost reporting periods beginning on or after October 1, 1991, the capital prospective payment system provides a payment amount for inpatient hospital capital-related costs as described in part 413, subpart G of this chapter.

(e) Excluded costs. The following inpatient hospital costs are excluded from the prospective payment amounts and are paid for on a reasonable cost basis:

(1) Capital-related costs for cost reporting periods beginning before October 1, 1991, and an allowance for return on equity, as described in §§ 413.130 and 413.157, respectively, of this chapter.

(2) Direct medical education costs for approved nursing and allied health education programs as described in § 413.85 of this chapter.

(3) Costs for direct medical and surgical services of physicians in teaching hospitals exercising the election in § 405.521 of this chapter.
(4) Heart, kidney, liver, lung, and pancreas acquisition costs incurred by approved transplantation centers.

(5) The costs of qualified nonphysician anesthetists’ services, as described in §412.113(c).

(f) Additional payments to hospitals. In addition to payments based on the prospective payment rates for inpatient operating costs and inpatient capital-related costs, hospitals receive payments for the following:

(1) Outlier cases, as described in subpart F of this part.

(2) The indirect costs of graduate medical education, as specified in subparts F and G of this part and in §412.105 for inpatient operating costs and in §412.322 for inpatient capital-related costs.

(3) Costs excluded from the prospective payment rates under paragraph (e) of this section, as provided in §412.115.

(4) Bad debts of Medicare beneficiaries, as provided in §412.115(a).

(5) ESRD beneficiary discharges if such discharges are ten percent or more of the hospital’s total Medicare discharges, as provided in §412.104.

(6) Serving a disproportionate share of low-income patients, as provided in §412.106 for inpatient operating costs and §412.320 for inpatient capital-related costs.

(7) The direct graduate medical education costs for approved residency programs in medicine, osteopathy, dentistry, and podiatry as described in §413.86 of this chapter.

(8) For discharges on or after June 19, 1990, and before October 1, 1994, and for discharges on or after October 1, 1997, a payment amount per unit for blood clotting factor provided to Medicare inpatients who have hemophilia.

§ 412.4 Discharges and transfers.

(a) Discharges. Subject to the provisions of paragraphs (b) and (c) of this section, a hospital inpatient is considered discharged from a hospital paid under the prospective payment system when—

(1) The patient is formally released from the hospital; or

(2) The patient dies in the hospital.

(b) Transfer—Basic rule. A discharge of a hospital inpatient is considered to be a transfer for purposes of payment under this part if the discharge is made under any of the following circumstances:

(1) From a hospital to the care of another hospital that is—

(i) Paid under the prospective payment system; or

(ii) Excluded from being paid under the prospective payment system because of participation in an approved Statewide cost control program as described in subpart C of part 403 of this chapter.

(2) From one inpatient area or unit of a hospital to another inpatient area or unit of the hospital that is paid under the prospective payment system.

(c) Transfers—Special 10 DRG rule. For discharges occurring on or after October 1, 1998, a discharge of a hospital inpatient is considered to be a transfer for purposes of this part when the patient’s discharge is assigned, as described in §412.60(c), to one of the qualifying diagnosis-related groups (DRGs) listed in paragraph (d) of this section and the discharge is made under any of the following circumstances—

(1) To a hospital or distinct part hospital unit excluded from the prospective payment system under subpart B of this part.

(2) To a skilled nursing facility.

(3) To home under a written plan of care for the provision of home health services from a home health agency and those services begin within 3 days after the date of discharge.

(d) Qualifying DRGs. The qualifying DRGs for purposes of paragraph (c) of this section are DRGs 14, 113, 210, 211, 236, 263, 264, 429, and 483.

(e) Payment for discharges. The hospital discharging an inpatient (under paragraph (a) of this section) is paid in full, in accordance with §412.2(b).

(f) Payment for transfers. (1) General rule. Except as provided in paragraph (f)(2) or (f)(3) of this section, a hospital

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that transfers an inpatient under the circumstances described in paragraph (b) or (c) of this section, is paid a graduated per diem rate for each day of the patient’s stay in that hospital, not to exceed the amount that would have been paid under subparts D and M of this part if the patient had been discharged to another setting. The per diem rate is determined by dividing the appropriate prospective payment rate (as determined under subparts D and M of this part) by the geometric mean length of stay for the specific DRG to which the case is assigned. Payment is graduated by paying twice the per diem amount for the first day of the stay, and the per diem amount for each subsequent day, up to the full DRG payment.

(2) Special rule for DRGs 209, 210, and 211. A hospital that transfers an inpatient under the circumstances described in paragraph (c) of this section and the transfer is assigned to DRGs 209, 210 or 211 is paid as follows:

(i) 50 percent of the appropriate prospective payment rate (as determined under subparts D and M of this part) for the first day of the stay; and

(ii) 50 percent of the amount calculated under paragraph (f)(1) of this section for each day of the stay, up to the full DRG payment.

(3) Transfer assigned to DRG 385. If a transfer is classified into DRG 385 (Neonates, died or transferred) the transferring hospital may qualify for an additional payment for extraordinary high-cost cases that meet the criteria for cost outliers as described in subpart F of this part.

§ 412.8 Publication of schedules for determining prospective payment rates.

(a) Initial prospective payment rates—

(1) For inpatient operating costs. Initial prospective payment rates for inpatient operating costs (for the period October 1, 1983 through September 30, 1984) were determined in accordance with documents published in the Federal Register on September 1, 1983 (48 FR 39839) and January 3, 1984 (49 FR 324).

(2) For inpatient capital-related costs. Initial prospective payment rates for inpatient capital-related costs (for the
§ 412.20 Hospital services subject to the prospective payment systems.

(a) Except for services described in paragraph (b) of this section, all covered inpatient hospital services furnished to beneficiaries during subject cost reporting periods are paid for under the prospective payment systems.

(b) Annual publication of schedule for determining prospective payment rates. (1) HCFA proposes changes in the methods, amounts, and factors used to determine inpatient prospective payment rates in a Federal Register document published for public comment not later than the April 1 before the beginning of the Federal fiscal year in which the proposed changes would apply.

(2) HCFA publishes a Federal Register document setting forth final methods, amounts, and factors for determining inpatient prospective payment rates not later than the August 1 before the Federal fiscal year in which the rates would apply.

§ 412.10 Changes in the DRG classification system.

(a) General rule. HCFA issues changes in the DRG classification system in a Federal Register notice at least annually. Except as specified in paragraphs (c) and (d) of this section, the DRG changes are effective prospectively with discharges occurring on or after the same date the payment rates are effective.

(b) Basis for changes in the DRG classification system. All changes in the DRG classification system are made using the principles established for the DRG system. This means that cases are classified so each DRG is—

(1) Clinically coherent; and

(2) Embraces an acceptable range of resource consumption.

(c) Interim coverage changes—(1) Criteria. HCFA makes interim changes to the DRG classification system during the Federal fiscal year to incorporate items and services newly covered under Medicare.

(2) Implementation and effective date. HCFA issues interim coverage changes through its administrative issuance system and makes the change effective as soon as administratively feasible.

(3) Publication for comment. HCFA publishes any change made under paragraph (c)(1) of this section in the next annual notice of changes to the DRG classification system published in accordance with paragraph (a) of this section.

(d) Interim changes to correct omissions and inequities—(1) Criteria. HCFA makes interim changes to the DRG classification system to correct a serious omission or inequity in the system only if failure to make the changes would have—

(i) A potentially substantial adverse impact on the health and safety of beneficiaries; or

(ii) A significant and unwarranted fiscal impact on hospitals or the Medicare program.

(2) Publication and effective date. HCFA publishes these changes in the Federal Register in a final notice with comment period with a prospective effective date. The change is also published for public information in the next annual notice of changes to the DRG classification system published in accordance with paragraph (a) of this section.

(e) Review by ProPAC. Changes published annually in accordance with paragraph (a) of this section are subject to review and comment by ProPAC upon publication. Interim changes to the DRG classification system that are made in accordance with paragraphs (c) and (d) of this section are subject to review by ProPAC before implementation.

§ 412.22 Excluded hospitals and hospital units: General rules.

(a) Criteria. Subject to the criteria set forth in paragraph (e) of this section, a hospital is excluded from the prospective payment systems if it meets the criteria for one or more of the excluded classifications described in § 412.23.

(b) Cost reimbursement. Except for those hospitals specified in paragraph (c) of this section, all excluded hospitals (and excluded hospital units, as described in §§ 412.23 through 412.29) are reimbursed under the cost reimbursement rules set forth in part 413 of this chapter, and are subject to the ceiling on the rate of hospital cost increases described in § 413.40 of this chapter.

(c) Special payment provisions. The following classifications of hospitals are paid under special provisions and therefore are not generally subject to the cost reimbursement or prospective payment rules of this chapter:

(1) Veterans Administration hospitals.

(2) Hospitals reimbursed under State cost control systems approved under part 403 of this chapter.

(3) Hospitals reimbursed in accordance with demonstration projects authorized under section 402(a) of Public Law 90-248 (42 U.S.C. 1395b-1) or section 222(a) of Public Law 92-603 (42 U.S.C. 1395b-1(note)).

(4) Nonparticipating hospitals furnishing emergency services to Medicare beneficiaries.

(d) Changes in hospitals’ status. For purposes of exclusion from the prospective payment systems under this subpart, the status of each currently participating hospital (excluded or not excluded) is determined at the beginning of each cost reporting period and is effective for the entire cost reporting period. Any changes in the status of the hospital are made only at the start of a cost reporting period.

(e) Hospitals within hospitals. Except as provided in paragraph (f) of this section, for cost reporting periods beginning on or after October 1, 1997, a hospital that occupies space in a building also used by another hospital, or in one or more entire buildings located on the same campus as buildings used by another hospital, must meet the following criteria in order to be excluded from the prospective payment system:

(1) Separate governing body. The hospital has a governing body that is separate from the governing body of the hospital occupying space in the same building or on the same campus. The hospital’s governing body is not under the control of the hospital occupying space in the same building or on the same campus, or of any third entity that controls both hospitals.

(2) Separate chief medical officer. The hospital has a single chief medical officer who reports directly to the governing body and who is responsible for all medical staff activities of the hospital. The chief medical officer of the hospital is not employed by or under contract with either the hospital occupying space in the same building or on the same campus or any third entity that controls both hospitals.

(3) Separate medical staff. The hospital has a medical staff that is separate from the medical staff of the hospital occupying space in the same building or on the same campus. The hospital’s medical staff is directly accountable to the governing body for the quality of medical care provided in the hospital.
and adopts and enforces bylaws governing medical staff activities, including criteria and procedures for recommending to the governing body the privileges to be granted to individual practitioners.

(4) Chief executive officer. The hospital has a single chief executive officer through whom all administrative authority flows, and who exercises control and surveillance over all administrative activities of the hospital. The chief executive officer is not employed by, or under contract with, either the hospital occupying space in the same building or on the same campus or any third entity that controls both hospitals.

(5) Performance of basic hospital functions. The hospital meets one of the following criteria:

(i) The hospital performs the basic functions specified in §§ 482.21 through 482.27, 482.30, and 482.42 of this chapter through the use of employees or under contracts or other agreements with entities other than the hospital occupying space in the same building or on the same campus, or a third entity that controls both hospitals. Food and dietetic services and housekeeping, maintenance, and other services necessary to maintain a clean and safe physical environment could be obtained under contracts or other agreements with the hospital occupying space in the same building or on the same campus, or with a third entity that controls both hospitals.

(ii) For the same period of at least 6 months used to determine compliance with the criterion regarding the age of inpatients in §412.23(d)(2) or the length-of-stay criterion in §412.23(e)(2), or for hospitals other than children’s or long-term care hospitals, for the period of at least 6 months immediately preceding the first cost reporting period for which exclusion is sought, the hospital has an inpatient population of whom at least 75 percent were referred to the hospital from a source other than another hospital occupying space in the same building or on the same campus.

(f) Application for certain hospitals. If a hospital was excluded from the prospective payment systems under the provisions of this section on or before September 30, 1995, and at that time occupied space in a building also used by another hospital, or in one or more entire buildings located on the same campus as buildings used by another hospital, the criteria in paragraph (e) of this section do not apply to the hospital.

(g) Definition of control. For purposes of this section, control exists if an individual or an organization has the power, directly or indirectly, significantly to influence or direct the actions or policies of an organization or institution.

(h) Satellite facilities. (1) For purposes of paragraphs (h)(2) through (h)(4) of this section, a satellite facility is a part of a hospital that provides inpatient services in a building also used by another hospital, or in one or more entire buildings located on the same campus as buildings used by another hospital.

(2) Except as provided in paragraph (h)(3) of this section, effective for cost reporting periods beginning on or after October 1, 1999, a hospital that has a satellite facility is a part of a hospital that provides inpatient services in a building also used by another hospital, or in one or more entire buildings located on the same campus as buildings used by another hospital.

(i) In the case of a hospital (other than a children's hospital) that was excluded from the prospective payment systems for the most recent cost reporting period beginning before October 1, 1997, the hospital’s number of...
§412.23 State-licensed and Medicare-certified beds, including those at the satellite facilities, does not exceed the hospital’s number of State-licensed and Medicare-certified beds on the last day of the hospital’s last cost reporting period beginning before October 1, 1997.

(ii) The satellite facility independently complies with—
(A) For psychiatric hospitals, the requirements under §412.23(a);
(B) For rehabilitation hospitals, the requirements under §412.23(b)(2);
(C) For children’s hospitals, the requirements under §412.23(d)(2); or
(D) For long-term care hospitals, the requirements under §§412.23(e)(1) through (e)(3)(i).

(iii) The satellite facility meets all of the following requirements:
(A) It maintains admission and discharge records that are separately identified from those of the hospital in which it is located and are readily available.
(B) It has beds that are physically separate from (that is, not commingled with) the beds of the hospital in which it is located.
(C) It is serviced by the same fiscal intermediary as the hospital of which it is a part.
(D) It is treated as a separate cost center of the hospital of which it is a part.
(E) For cost reporting and apportionment purposes, it uses an accounting system that properly allocates costs and maintains adequate statistical data to support the basis of allocation.
(F) It reports its costs on the cost report of the hospital of which it is a part, covering the same fiscal period and using the same method of apportionment as the hospital of which it is a part.

(3) Except as provided in paragraph (h)(4) of this section, the provisions of paragraph (h)(2) of this section do not apply to—

(i) Any hospital structured as a satellite facility on September 30, 1999, and excluded from the prospective payment systems on that date, to the extent the hospital continues operating under the same terms and conditions, including the number of beds and square footage considered, for purposes of Medicare participation and payment, to be part of the hospital, in effect on September 30, 1999; or
(ii) Any hospital excluded from the prospective payment systems under §412.23(e)(2).

(4) In applying the provisions of paragraph (h)(3) of this section, any hospital structured as a satellite facility on September 30, 1999, may increase or decrease the square footage of the satellite facility or may decrease the number of beds in the satellite facility if these changes are made necessary by relocation of a facility—

(i) To permit construction or renovation necessary for compliance with changes in Federal, State, or local law; or
(ii) Because of catastrophic events such as fires, floods, earthquakes, or tornadoes.


§412.23 Excluded hospitals: Classifications.

Hospitals that meet the requirements for the classifications set forth in this section may not be reimbursed under the prospective payment systems.

(a) Psychiatric hospitals. A psychiatric hospital must—

(1) Be primarily engaged in providing, by or under the supervision of a psychiatrist, psychiatric services for the diagnosis and treatment of mentally ill persons; and
(2) Meet the conditions of participation for hospitals and special conditions of participation for psychiatric hospitals set forth in part 482 of this chapter.

(b) Rehabilitation hospitals. A rehabilitation hospital must meet the following requirements:

(1) Have a provider agreement under part 489 of this chapter to participate as a hospital.
(2) Except in the case of a newly participating hospital seeking exclusion for its first 12-month cost reporting period, as described in paragraph (b)(8) of this section, show that during its most recent 12-month cost reporting period, it served an inpatient population of whom at least 75 percent required intensive rehabilitative services for the
treatment of one or more of the following conditions:
(i) Stroke.
(ii) Spinal cord injury.
(iii) Congenital deformity.
(iv) Amputation.
(v) Major multiple trauma.
(vi) Fracture of femur (hip fracture).
(vii) Brain injury.
(viii) Polyarthritis, including rheumatoid arthritis.
(ix) Neurological disorders, including multiple sclerosis, motor neuron diseases, polyneuropathy, muscular dystrophy, and Parkinson's disease.
(x) Burns.

(3) Have in effect a preadmission screening procedure under which each prospective patient's condition and medical history are reviewed to determine whether the patient is likely to benefit significantly from an intensive inpatient hospital program or assessment.

(4) Ensure that the patients receive close medical supervision and furnish, through the use of qualified personnel, rehabilitation nursing, physical therapy, and occupational therapy, plus, as needed, speech therapy, social or psychological services, and orthotic and prosthetic services.

(5) Have a director of rehabilitation who—
(i) Provides services to the hospital and its inpatients on a full-time basis;
(ii) Is a doctor of medicine or osteopathy;
(iii) Is licensed under State law to practice medicine or surgery; and
(iv) Has had, after completing a one-year hospital internship, at least two years of training or experience in the medical management of inpatients requiring rehabilitation services.

(6) Have a plan of treatment for each inpatient that is established, reviewed, and revised as needed by a physician in consultation with other professional personnel who provide services to the patient.

(7) Use a coordinated multidisciplinary team approach in the rehabilitation of each inpatient, as documented by periodic clinical entries made in the patient's medical record to note the patient's status in relationship to goal attainment, and that team conferences are held at least every two weeks to determine the appropriateness of treatment.

(8) A hospital that seeks exclusion as a rehabilitation hospital for the first full 12-month cost reporting period that occurs after it becomes a Medicare participating hospital may provide a written certification that the inpatient population it intends to serve meets the requirements of paragraph (b)(2) of this section, instead of showing that it has treated such a population during its most recent 12-month cost reporting period. The written certification is also effective for any cost reporting period of not less than one month and not more than 11 months occurring between the date the hospital began participating in Medicare and the start of the hospital's regular 12-month cost reporting period.

(9) For cost reporting periods beginning on or after October 1, 1991, if a hospital is excluded from the prospective payment systems for a cost reporting period under paragraph (b)(8) of this section, but the inpatient population it actually treated during that period does not meet the requirements of paragraph (b)(2) of this section, HCFA adjusts payments to the hospital retroactively in accordance with the provisions in §412.130 of this part.

c) [Reserved]

d) Children's hospitals. A children's hospital must—
(1) Have a provider agreement under part 489 of this chapter to participate as a hospital; and
(2) Be engaged in furnishing services to inpatients who are predominantly individuals under the age of 18.

e) Long-term care hospitals. A long-term care hospital must meet the requirements of paragraphs (e)(1) or (e)(2) of this section, and, where applicable, the additional requirements §412.22(e).

(1) The hospital must have a provider agreement under part 489 of this chapter to participate as a hospital and an average inpatient length of stay greater than 25 days as calculated under paragraph (e)(3) of this section.

(2) For cost reporting periods beginning on or after August 5, 1997, a hospital that was first excluded from the prospective payment system under this section in 1986 must have an average inpatient length of stay of greater than
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20 days, as calculated under paragraph (e)(3) of this section, and must demonstrate that at least 80 percent of its annual Medicare inpatient discharges in the 12-month cost reporting period ending in fiscal year 1997 have a principal diagnosis that reflects a finding of neoplastic disease as defined in paragraph (f)(1)(iv) of this section.

(3) The average inpatient length of stay is calculated—

(i) By dividing the number of total inpatient days (less leave or pass days) by the number of total discharges for the hospital’s most recent complete cost reporting period;

(ii) If a change in the hospital’s average length-of-stay is indicated, by the same method for the immediately preceding 6-month period; or

(iii) If a hospital has undergone a change of ownership (as described in § 489.18 of this chapter) at the start of a cost reporting period or at any time within the preceding 6 months, the hospital may be excluded from the prospective payment system as a long-term care hospital for a cost reporting period if, for the 6 months immediately preceding the start of the period (including time before the change of ownership), the hospital has the required average length of stay, continuously operated as a hospital, and continuously participated as a hospital in Medicare.

(f) Cancer hospitals—(1) General rule.

Except as provided in paragraph (f)(2) of this section, if a hospital meets the following criteria, it is classified as a cancer hospital and is excluded from the prospective payment systems beginning with its first cost reporting period.

(i) It was recognized as a comprehensive cancer center or clinical cancer research center by the National Cancer Institute of the National Institutes of Health as of April 20, 1983.

(ii) It is classified on or before December 31, 1989, was not operating a demonstration project under section 1814(b) of the Act; and

(iii) It was recognized as a comprehensive cancer center or clinical cancer research center by the National Cancer Institute of the National Institutes of Health as of April 20, 1983.

(iii) It demonstrates that the entire facility is organized primarily for treatment of and research on cancer (that is, the facility is not a subunit of an acute general hospital or university-based medical center).

(iv) It shows that at least 50 percent of its total discharges have a principal diagnosis that reflects a finding of neoplastic disease. (The principal diagnosis for this purpose is defined as the condition established after study to be chiefly responsible for occasioning the admission of the patient to the hospital. For the purposes of meeting this definition, only discharges with ICD-9-CM principal diagnosis codes of 140 through 239, V58.0, V58.1, V66.1, V66.2, or 990 will be considered to reflect neoplastic disease.)

(2) Alternative. A hospital that applied for and was denied, on or before December 31, 1990, classification as a cancer hospital under the criteria set forth in paragraph (f)(1) of this section is classified as a cancer hospital and is excluded from the prospective payment systems beginning with its first cost reporting period.

(i) Licensed for fewer than 50 acute care beds as of August 5, 1997;

(ii) Is located in a State that as of December 19, 1989, was not operating a demonstration project under section 1814(b) of the Act; and

(iii) Demonstrates that, for the 4-year period ending on December 31, 1996, at least 50 percent of its total discharges have a principal diagnosis that reflects a finding of neoplastic disease as defined in paragraph (f)(1)(iv) of this section.

(g) Hospitals outside the 50 States, the District of Columbia, or Puerto Rico.

A hospital is excluded from the prospective payment systems if it is not located in one of the fifty States, the District of Columbia, or Puerto Rico.

(h) Hospitals reimbursed under special arrangements.

A hospital must be excluded from prospective payment for
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inpatient hospital services if it is reimbursed under special arrangement as provided in §412.22(c).

§ 412.25 Excluded hospital units: Common requirements.

(a) Basis for exclusion. In order to be excluded from the prospective payment system, a psychiatric or rehabilitation unit must meet the following requirements.

(1) Be part of an institution that—
(i) Has in effect an agreement under part 489 of this chapter to participate as a hospital;
(ii) Is not excluded in its entirety from the prospective payment systems; and
(iii) Has enough beds that are not excluded from the prospective payment systems to permit the provision of adequate cost information, as required by §413.24(c) of this chapter.

(2) Have written admission criteria that are applied uniformly to both Medicare and non-Medicare patients.

(3) Have admission and discharge records that are separately identified from those of the hospital in which it is located and are readily available.

(4) Have policies specifying that necessary clinical information is transferred to the unit when a patient of the hospital is transferred to the unit.

(5) Meet applicable State licensure laws.

(6) Have utilization review standards applicable for the type of care offered in the unit.

(7) Have beds physically separate from (that is, not commingled with) the hospital’s other beds.

(8) Be serviced by the same fiscal intermediary as the hospital.

(9) Be treated as a separate cost center for cost finding and apportionment purposes.

(10) Use an accounting system that properly allocates costs.

(11) Maintain adequate statistical data to support the basis of allocation.

(12) Report its costs in the hospital’s cost report covering the same fiscal period and using the same method of apportionment as the hospital.

(13) As of the first day of the first cost reporting period for which all other exclusion requirements are met, the unit is fully equipped and staffed and is capable of providing hospital inpatient psychiatric or rehabilitation care regardless of whether there are any inpatients in the unit on that date.

(b) Changes in the size of excluded units. For purposes of exclusions from the prospective payment systems under this section, changes in the number of beds and square footage considered to be part of each excluded unit are allowed as specified in paragraphs (b)(1) through (b)(3) of this section.

(1) Increase in size. Except as described in paragraph (b)(3) of this section, the number of beds and square footage of an excluded unit may be increased only at the start of a cost reporting period.

(2) Decrease in size. Except as described in paragraph (b)(3) of this section, the number of beds and square footage of an excluded unit may be decreased at any time during a cost reporting period if the hospital notifies its fiscal intermediary and the HCFA Regional Office in writing of the planned decrease at least 30 days before the date of the decrease, and maintains the information needed to accurately determine costs that are attributable to the excluded unit. Any decrease in the number of beds or square footage considered to be part of an excluded unit made during a cost reporting period must remain in effect for the rest of that cost reporting period.

(3) Exception to changes in square footage and bed size. The number of beds in an excluded unit may be decreased, and the square footage considered to be part of the unit may be either increased or decreased, at any time, if these changes are made necessary by relocation of a unit—
(i) To permit construction or renovation necessary for compliance with changes in Federal, State, or local law affecting the physical facility; or
(ii) Because of catastrophic events such as fires, floods, earthquakes, or tornadoes.

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(c) Changes in the status of hospital units. For purposes of exclusions from the prospective payment systems under this section, the status of each hospital unit (excluded or not excluded) is determined as specified in paragraphs (c)(1) and (c)(2) of this section.

(1) The status of a hospital unit may be changed from not excluded to excluded only at the start of the cost reporting period. If a unit is added to a hospital after the start of a cost reporting period, it cannot be excluded from the prospective payment systems before the start of a hospital’s next cost reporting period.

(2) The status of a hospital unit may be changed from excluded to not excluded at any time during a cost reporting period, but only if the hospital notifies the fiscal intermediary and the HCFA Regional Office in writing of the change at least 30 days before the date of the change, and maintains the information needed to accurately determine costs that are or are not attributable to the excluded unit. A change in the status of a unit from excluded to not excluded that is made during a cost reporting period must remain in effect for the rest of that cost reporting period.

(d) Number of excluded units. Each hospital may have only one unit of each type (psychiatric or rehabilitation) excluded from the prospective payment systems.

(e) Satellite facilities. (1) For purposes of paragraphs (e)(2) through (e)(4) of this section, a satellite facility is a part of a hospital unit that provides inpatient services in a building also used by another hospital, or in one or more entire buildings located on the same campus as buildings used by another hospital.

(2) Except as provided in paragraph (e)(3) of this section, effective for cost reporting periods beginning on or after October 1, 1999, a hospital unit that establishes a satellite facility must meet the following requirements in order to be excluded from the prospective payment systems for any period:

(i) In the case of a unit excluded from the prospective payment systems for the most recent cost reporting period beginning before October 1, 1997, the unit’s number of State-licensed and Medicare-certified beds, including those at the satellite facility, does not exceed the unit’s number of State-licensed and Medicare-certified beds on the last day of the unit’s last cost reporting period beginning before October 1, 1997.

(ii) The satellite facility independently complies with—

(A) For a rehabilitation unit, the requirements under §412.23(b)(2); or

(B) For a psychiatric unit, the requirements under §412.27(a).

(iii) The satellite facility meets all of the following requirements:

(A) It maintains admission and discharge records that are separately identified from those of the hospital in which it is located and are readily available.

(B) It has beds that are physically separate from (that is, not commingled with) the beds of the hospital in which it is located.

(C) It is serviced by the same fiscal intermediary as the hospital unit of which it is a part.

(D) It is treated as a separate cost center of the hospital unit of which it is a part.

(E) For cost reporting and apportionment purposes, it uses an accounting system that properly allocates costs and maintains adequate statistical data to support the basis of allocation.

(F) It reports its costs on the cost report of the hospital of which it is a part, covering the same fiscal period and using the same method of apportionment as the hospital of which it is a part.

(3) Except as specified in paragraph (e)(4) of this section, the provisions of paragraph (e)(2) of this section do not apply to any unit structured as a satellite facility on September 30, 1999, and excluded from the prospective payment systems on that date, to the extent the unit continues operating under the same terms and conditions, including the number of beds and square footage considered to be part of the unit, in effect on September 30, 1999.

(4) In applying the provisions of paragraph (h)(3) of this section, any unit structured as a satellite facility as of
§ 412.27 Excluded psychiatric units: Additional requirements.

In order to be excluded from the prospective payment systems, a psychiatric unit must meet the following requirements:

(a) Admit only patients whose admission to the unit is required for active treatment, of an intensity that can be provided appropriately only in an inpatient hospital setting, of a psychiatric principal diagnosis that is listed in the Third Edition of the American Psychiatric Association’s Diagnostic and Statistical Manual, or in Chapter Five ("Mental Disorders") of the International Classification of Diseases, Ninth Revision, Clinical Modification.

(b) Furnish, through the use of qualified personnel, psychological services, social work services, psychiatric nursing, occupational therapy, and recreational therapy.

(c) Maintain medical records that permit determination of the degree and intensity of the treatment provided to individuals who are furnished services in the unit, and that meet the following requirements:

(1) Development of assessment/diagnostic data. Medical records must stress the psychiatric components of the record, including history of findings and treatment provided for the psychiatric condition for which the inpatient is treated in the unit.

(ii) The identification data must include the inpatient’s legal status.

(2) Psychiatric evaluation. Each inpatient must receive a psychiatric evaluation that must—

(i) Be completed within 60 hours of admission;

(ii) Include a medical history;

(iii) Contain a record of mental status;

(iv) Note the onset of illness and the circumstances leading to admission;

(v) Describe attitudes and behavior;

(vi) Estimate intellectual functioning, memory functioning, and orientation; and

(vii) Include an inventory of the inpatient’s assets in descriptive, not interpretative fashion.

(3) Treatment plan.

(i) Each inpatient must have an individual comprehensive treatment plan that must be based on an inventory of the inpatient’s strengths and disabilities. The written plan must include a substantiated diagnosis; short-term and long-term goals; the specific treatment modalities utilized; the responsibilities of each member of the treatment team; and adequate documentation to justify the diagnosis and the treatment and rehabilitation activities carried out; and

(ii) The treatment received by the inpatient must be documented in such a way as to assure that all active therapeutic efforts are included.

(4) Recording progress. Progress notes must be recorded by the doctor of medicine or osteopathy responsible for the care of the inpatient, a nurse, social worker and, when appropriate, others at the time of admission, and must include the diagnoses of intercurrent diseases as well as the psychiatric diagnoses.

(iii) The reasons for admission must be clearly documented as stated by the inpatient or others significantly involved, or both.

(iv) The social service records, including reports of interviews with inpatients, family members, and others must provide an assessment of home plans and family attitudes, and community resource contacts as well as a social history.

(v) When indicated, a complete neurological examination must be recorded at the time of the admission physical examination.

(2) Psychiatric evaluation. Each inpatient must receive a psychiatric evaluation that must—

(i) Be completed within 60 hours of admission;

(ii) Include a medical history;

(iii) Contain a record of mental status;

(iv) Note the onset of illness and the circumstances leading to admission;

(v) Describe attitudes and behavior;

(vi) Estimate intellectual functioning, memory functioning, and orientation; and

(vii) Include an inventory of the inpatient’s assets in descriptive, not interpretative fashion.

(3) Treatment plan.

(i) Each inpatient must have an individual comprehensive treatment plan that must be based on an inventory of the inpatient’s strengths and disabilities. The written plan must include a substantiated diagnosis; short-term and long-term goals; the specific treatment modalities utilized; the responsibilities of each member of the treatment team; and adequate documentation to justify the diagnosis and the treatment and rehabilitation activities carried out; and

(ii) The treatment received by the inpatient must be documented in such a way as to assure that all active therapeutic efforts are included.

(4) Recording progress. Progress notes must be recorded by the doctor of medicine or osteopathy responsible for the care of the inpatient, a nurse, social worker and, when appropriate, others
significantly involved in active treatment modalities. The frequency of progress notes is determined by the condition of the inpatient but must be recorded at least weekly for the first two months and at least once a month thereafter and must contain recommendations for revisions in the treatment plan as indicated as well as precise assessment of the inpatient’s progress in accordance with the original or revised treatment plan.

(5) Discharge planning and discharge summary. The record of each patient who has been discharged must have a discharge summary that includes a recapitulation of the inpatient’s hospitalization in the unit and recommendations from appropriate services concerning follow-up or aftercare as well as a brief summary of the patient’s condition on discharge.

(d) Meet special staff requirements in that the unit must have adequate numbers of qualified professional and supportive staff to evaluate inpatients, formulate written, individualized, comprehensive treatment plans, provide active treatment measures and engage in discharge planning, as follows:

(1) Personnel. The unit must employ or undertake to provide adequate numbers of qualified professional, technical, and consultative personnel to—

(i) Evaluate inpatients;

(ii) Formulate written, individualized, comprehensive treatment plans;

(iii) Provide active treatment measures; and

(iv) Engage in discharge planning.

(2) Director of inpatient psychiatric services: Medical staff. Inpatient psychiatric services must be under the supervision of a clinical director, service chief, or equivalent who is qualified to provide the leadership required for an intensive treatment program. The number and qualifications of doctors of medicine and osteopathy must be adequate to provide essential psychiatric services.

(i) The clinical director, service chief, or equivalent must meet the training and experience requirements for examination by the American Board of Psychiatry and Neurology or the American Osteopathic Board of Neurology and Psychiatry.

(ii) The director must monitor and evaluate the quality and appropriateness of services and treatment provided by the medical staff.

(3) Nursing services. The unit must have a qualified director of psychiatric nursing services. In addition to the director of nursing, there must be adequate numbers of registered nurses, licensed practical nurses, and mental health workers to provide nursing care necessary under each inpatient’s active treatment program and to maintain progress notes on each inpatient.

(i) The director of psychiatric nursing services must be a registered nurse who has a master’s degree in psychiatric or mental health nursing, or its equivalent, from a school of nursing accredited by the National League for Nursing, or be qualified by education and experience in the care of the mentally ill. The director must demonstrate competence to participate in interdisciplinary formulation of individual treatment plans; to give skilled nursing care and therapy; and to direct, monitor, and evaluate the nursing care furnished.

(ii) The staffing pattern must ensure the availability of a registered nurse 24 hours each day. There must be adequate numbers of registered nurses, licensed practical nurses, and mental health workers to provide the nursing care necessary under each inpatient’s active treatment program.

(4) Psychological services. The unit must provide or have available psychological services to meet the needs of the inpatients. The services must be furnished in accordance with acceptable standards of practice, service objectives, and established policies and procedures.

(5) Social services. There must be a director of social services who monitors and evaluates the quality and appropriateness of social services furnished. The services must be furnished in accordance with accepted standards of practice and established policies and procedures. Social service staff responsibilities must include, but are not limited to, participating in discharge planning, arranging for follow-up care, and developing mechanisms for exchange of appropriate information with sources outside the hospital.
§ 412.30 Exclusion of new rehabilitation units and expansion of units already excluded.

(a) Bed capacity in units. A decrease in bed capacity must remain in effect for at least a full 12-month cost reporting period before an equal or lesser number of beds can be added to the hospital's licensure and certification and considered "new" under paragraph (b) of this section. Thus, when a hospital seeks to establish a new unit under the criteria under paragraph (b) of this section, or to enlarge an existing unit under the criteria under paragraph (d) of this section, the regional office will review its records on the facility to determine whether any beds have been delicensed and decertified during the 12-month cost reporting period before the period for which the hospital seeks to add the beds. To the extent bed capacity was removed from the hospital's licensure and certification during that period, that amount of bed capacity may not be considered "new" under paragraph (b) of this section.

(b) New units. 

(1) A hospital unit is considered a new unit if the hospital—

(i) Has not previously sought exclusion for any rehabilitation unit; and

(ii) Has obtained approval, under State licensure and Medicare certification, for an increase in its hospital bed capacity that is greater than 50 percent of the number of beds in the unit.

(2) A hospital that seeks exclusion of a new rehabilitation unit may provide

§ 412.29 Excluded rehabilitation units: Additional requirements.

In order to be excluded from the prospective payment systems, a rehabilitation unit must meet the following requirements:

(a) Have met either the requirements for—

(1) New units under §412.30(a); or

(2) Converted units under §412.30(b).

(b) Have in effect a preadmission screening procedure under which each prospective patient's condition and medical history are reviewed to determine whether the patient is likely to benefit significantly from an intensive inpatient program or assessment.

(c) Ensure that the patients receive close medical supervision and furnish, through the use of qualified personnel, rehabilitation nursing, physical therapy, and occupational therapy, plus, as needed, speech therapy, social services or psychological services, and orthotic and prosthetic services.

(d) Have a plan of treatment for each inpatient that is established, reviewed, and revised as needed by a physician in consultation with other professional personnel who provide services to the patient.

(e) Use a coordinated multidisciplinary team approach in the rehabilitation of each inpatient, as documented by periodic clinical entries made in the patient's medical record to note the patient's status in relationship to goal attainment, and that team conferences are held at least every two weeks to determine the appropriateness of treatment.

(f) Have a director of rehabilitation who—

(1) Provides services to the unit and to its inpatients for at least 20 hours per week;

(2) Is a doctor of medicine or osteopathy;

(3) Is licensed under State law to practice medicine or surgery; and

(4) Has had, after completing a one-year hospital internship, at least two years of training or experience in the medical management of inpatients requiring rehabilitation services.
§ 412.30  a written certification that the inpatient population the hospital intends the unit to serve meets the requirements of §412.23(b)(2) instead of showing that the unit has treated such a population during the hospital’s most recent cost reporting period.

(3) The written certification described in paragraph (a)(2) of this section is effective for the first full cost reporting period during which the unit is used to provide hospital inpatient care. If the hospital has not previously participated in the Medicare program as a hospital, the written certification also is effective for any cost reporting period of not less than 1 month and not more than 11 months occurring between the date the hospital began participating in Medicare and the start of the hospital’s regular 12-month cost reporting period.

(4) If a hospital that has not previously participated in the Medicare program seeks exclusion of a rehabilitation unit, it may designate certain beds as a new rehabilitation unit for the first full 12-month cost reporting period that occurs after it becomes a Medicare-participating hospital. The written certification described in paragraph (b)(2) of this section also is effective for any cost reporting period of not less than 1 month and not more than 11 months occurring between the date the hospital began participating in Medicare and the start of the hospital’s regular 12-month cost reporting period.

(5) A hospital that has undergone a change of ownership or leasing as defined in §489.18 of this chapter is not considered to have participated previously in the Medicare program.

(1) New bed capacity. The beds that a hospital seeks to add to its excluded rehabilitation unit are considered new beds only if—

(i) The hospital’s State-licensed and Medicare-certified bed capacity increases at the start of the cost reporting period for which the hospital seeks to increase the size of its excluded rehabilitation unit, or at any time after the start of the preceding cost reporting period; and

(ii) The hospital has obtained approval, under State licensure and Medicare certification, for an increase in its hospital bed capacity that is greater than 50 percent of the number of beds it seeks to add to the unit.

(2) Conversion of existing bed capacity. A hospital may increase the size of its excluded rehabilitation unit through conversion of existing bed capacity only if it shows that, for all of the hospital’s most recent cost reporting period of at least 12 months, the beds have been used to treat an inpatient population meeting the requirements of §412.23(b)(2).

(e) Retroactive adjustments for certain units. For cost reporting periods beginning on or after October 1, 1991, if a hospital has a new rehabilitation unit excluded from the prospective payment systems for a cost reporting period under paragraph (a) of this section or expands an existing rehabilitation unit under paragraph (c) of this section, but the inpatient population actually treated in the new unit or the beds added to the existing unit during that cost reporting period does not meet the requirements in §412.23(b)(2), HCFA adjusts payments to the hospital retroactively in accordance with the provisions in §412.130 of this part.

Subpart C—Conditions for Payment Under the Prospective Payment Systems for Inpatient Operating Costs and Inpatient Capital-Related Costs

§ 412.40 General requirements.

(a) A hospital must meet the conditions of this subpart to receive payment under the prospective payment systems for inpatient hospital services furnished to Medicare beneficiaries.

(b) If a hospital fails to comply fully with these conditions with respect to inpatient hospital services furnished to one or more Medicare beneficiaries, HCFA may, as appropriate—

(1) Withhold Medicare payment (in full or in part) to the hospital until the hospital provides adequate assurances of compliance; or

(2) Terminate the hospital’s provider agreement.

[50 FR 12741, Mar. 29, 1985, as amended at 57 FR 39821, Sept. 1, 1992]

§ 412.42 Limitations on charges to beneficiaries.

(a) Prohibited charges. A hospital may not charge a beneficiary for any services for which payment is made by Medicare, even if the hospital’s costs of furnishing services to that beneficiary are greater than the amount the hospital is paid under the prospective payment systems.

(b) Permitted charges—Stay covered. A hospital receiving payment under the prospective payment systems for a covered hospital stay (that is, a stay that includes at least one covered day) may charge the Medicare beneficiary or other person only for the following:

(1) The applicable deductible and coinsurance amounts under §§ 409.82, 409.83, and 409.87 of this chapter.

(2) Noncovered items and services, furnished at any time during a covered stay, unless they are excluded from coverage only on the basis of the following:

(i) The exclusion of custodial care under § 405.310(g) of this chapter (see paragraph (c) of this section for when charges may be made for custodial care).

(ii) The exclusion of medically unnecessary items and services under § 405.310(k) of this chapter (see paragraphs (c) and (d) of this section for when charges may be made for medically unnecessary items and services).

(iii) The exclusion under § 405.310(m) of this chapter of nonphysician services furnished to hospital inpatients by other than the hospital or a provider or supplier under arrangements made by the hospital.

(iv) The exclusion of items and services furnished when the patient is not entitled to Medicare Part A benefits under subpart A of part 406 of this chapter (see paragraph (e) of this section for when charges may be made for items and services furnished when the patient is not entitled to benefits).

(v) The exclusion of items and services furnished after Medicare Part A benefits are exhausted under § 409.61 of this chapter (see paragraph (e) of this section for when charges may be made for items and services furnished after benefits are exhausted).

(c) Custodial care and medically unnecessary inpatient hospital care. A hospital may charge a beneficiary for services excluded from coverage on the basis of § 411.15(g) of this chapter (custodial care) or § 411.15(k) of this chapter (medically unnecessary services) and furnished by the hospital after all of the following conditions have been met:

(1) The hospital (acting directly or through its utilization review committee) determines that the beneficiary no longer requires inpatient hospital care. (The phrase “inpatient hospital care” includes cases where a beneficiary needs a SNF level of care, but, under Medicare criteria, a SNF-level bed is not available. This also means that a hospital may find that a patient awaiting SNF placement no longer requires inpatient hospital care because either a SNF-level bed has become available or the patient no longer requires SNF-level care.)

(2) The attending physician agrees with the hospital’s determination in writing (for example, by issuing a written discharge order). If the hospital believes that the beneficiary does not require inpatient hospital care but is unable to obtain the agreement of the physician, it may request an immediate review of the case by the PRO.
Concurrence by the PRO in the hospital’s determination will serve in lieu of the physician’s agreement.

(3) The hospital (acting directly or through its utilization review committee) notifies the beneficiary (or person acting on his or her behalf) in writing that—
   (i) In the hospital’s opinion, and with the attending physician’s concurrence or that of the PRO, the beneficiary no longer requires inpatient hospital care;
   (ii) Customary charges will be made for continued hospital care beyond the second day following the date of the notice;
   (iii) The PRO will make a formal determination on the validity of the hospital’s finding if the beneficiary remains in the hospital after he or she is liable for charges;
   (iv) The determination of the PRO made after the beneficiary received the purportedly noncovered services will be appealable by the hospital, the attending physician, or the beneficiary under the appeals procedures that apply to PRO determinations affecting Medicare Part A payment; and
   (v) The charges for continued care will be invalid and refunded if collected by the hospital, to the extent that a finding is made that the beneficiary required continued care beyond the point indicated by the hospital.

(4) If the beneficiary remains in the hospital after the appropriate notification, and the hospital, the physician who concurred in the hospital determination on which the notice was based, or PRO subsequently finds that the beneficiary requires an acute level of inpatient hospital care, the hospital may not charge the beneficiary for continued care until the hospital once again determines that the beneficiary no longer requires inpatient care, secures concurrence, and notifies the beneficiary, as required in paragraphs (c)(1), (c)(2), and (c)(3) of this section.

(d) Medically unnecessary diagnostic and therapeutic services. A hospital may charge a beneficiary for diagnostic procedures and studies, and therapeutic procedures and courses of treatment (for example, experimental procedures) that are excluded from coverage under §405.310(k) of this chapter (medically unnecessary items and services), even though the beneficiary requires continued inpatient hospital care, if those services are furnished after the beneficiary (or the person acting on his or her behalf) has acknowledged in writing that the hospital (acting directly or through its utilization review committee and with the concurrence of the intermediary) has informed him or her as follows:

   (1) In the hospital’s opinion, which has been agreed to by the intermediary, the services to be furnished are not considered reasonable and necessary under Medicare.
   (2) Customary charges will be made if he or she receives the services.
   (3) If the beneficiary receives the services, a formal determination on the validity of the hospital’s finding is made by the intermediary and, to the extent that the decision requires the exercise of medical judgment, the PRO.
   (4) The determination is appealable by the hospital, the attending physician, or the beneficiary under the appeals procedure that applies to determinations affecting Medicare Part A payment.
   (5) The charges for the services will be invalid and, to the extent collected, will be refunded by the hospital if the services are found to be covered by Medicare.

(e) Services furnished on days when the individual is not entitled to Medicare Part A benefits or has exhausted the available benefits. The hospital may charge the beneficiary its customary charges for noncovered items and services furnished on outlier days (as described in Subpart F of this part) for which payment is denied because the beneficiary is not entitled to Medicare Part A or his or her Medicare Part A benefits are exhausted. (1) If payment is considered for outlier days, the entire stay is reviewed and days up to the number of days in excess of the outlier threshold may be denied on the basis of non-entitlement to Part A or exhaustion of benefits. (2) In applying this rule, the latest days will be denied first.

(f) Differential for private room or other luxury services. The hospital may charge the beneficiary the customary charge differential for a private room or other luxury service that is more expensive than is medically required and
§ 412.48 Denial of payment as a result of admissions and quality review.

(a) If HCFA determines, on the basis of information supplied by a PRO that a hospital has misrepresented admissions, discharges, or billing information, or has taken an action that results in the unnecessary admission of an individual entitled to benefits under Part A, unnecessary multiple admissions of an individual, or other inappropriate medical or other practices with respect to beneficiaries or billing for services furnished to beneficiaries, HCFA may as appropriate—

(1) Deny payment (in whole or in part) under Part A with respect to inpatient hospital services provided with respect to such an unnecessary admission or subsequent readmission of an individual; or

(2) Require the hospital to take other corrective action necessary to prevent or correct the inappropriate practice.

(b) When payment with respect to admission of an individual patient is denied by a PRO under paragraph (a)(1) of this section, and liability is not waived in accordance with §§405.330 through

§ 412.46 Medical review requirements: Physician acknowledgement.

(a) Basis. Because payment under the prospective payment system is based in part on each patient’s principal and secondary diagnoses and major procedures performed, as evidenced by the physician’s entries in the patient’s medical record, physicians must complete an acknowledgement statement to this effect.

(b) Content of physician acknowledgement statement. When a claim is submitted, the hospital must have on file a signed and dated acknowledgement from the attending physician that the physician has received the following notice:

Notice to Physicians: Medicare payment to hospitals is based in part on each patient’s principal and secondary diagnoses and the major procedures performed on the patient, as attested to by the patient’s attending physician by virtue of his or her signature in the medical record. Anyone who misrepresents, falsifies, or conceals essential information required for payment of Federal funds, may be subject to fine, imprisonment, or civil penalty under applicable Federal laws.

(c) Completion of acknowledgement. The acknowledgement must be completed by the physician at the time that the physician is granted admitting privileges at the hospital, or before or at the time the physician admits his or her first patient. Existing acknowledgements signed by physicians already on staff remain in effect as long as the physician has admitting privileges at the hospital.

[60 FR 45847, Sept. 1, 1995]

§ 412.44 Medical review requirements: Admissions and quality review.

Beginning on November 15, 1984, a hospital must have an agreement with a PRO to have the PRO review, on an ongoing basis, the following:

(a) The medical necessity, reasonableness and appropriateness of hospital admissions and discharges.

(b) The medical necessity, reasonableness and appropriateness of inpatient hospital care for which additional payment is sought under the outlier provisions of §§412.82 and 412.84 of this chapter.

(c) The validity of the hospital’s diagnostic and procedural information.

(d) The completeness, adequacy, and quality of the services furnished in the hospital.

(e) Other medical or other practices with respect to beneficiaries or billing for services furnished to beneficiaries.


§ 412.48 Denial of payment as a result of admissions and quality review.

(a) If HCFA determines, on the basis of information supplied by a PRO that a hospital has misrepresented admissions, discharges, or billing information, or has taken an action that results in the unnecessary admission of an individual entitled to benefits under Part A, unnecessary multiple admissions of an individual, or other inappropriate medical or other practices with respect to beneficiaries or billing for services furnished to beneficiaries, HCFA may as appropriate—

(1) Deny payment (in whole or in part) under Part A with respect to inpatient hospital services provided with respect to such an unnecessary admission or subsequent readmission of an individual; or

(2) Require the hospital to take other corrective action necessary to prevent or correct the inappropriate practice.

(b) When payment with respect to admission of an individual patient is denied by a PRO under paragraph (a)(1) of this section, and liability is not waived in accordance with §§405.330 through

§ 412.46 Medical review requirements: Physician acknowledgement.

(a) Basis. Because payment under the prospective payment system is based in part on each patient’s principal and secondary diagnoses and major procedures performed, as evidenced by the physician’s entries in the patient’s medical record, physicians must complete an acknowledgement statement to this effect.

(b) Content of physician acknowledgement statement. When a claim is submitted, the hospital must have on file a signed and dated acknowledgement from the attending physician that the physician has received the following notice:

Notice to Physicians: Medicare payment to hospitals is based in part on each patient’s principal and secondary diagnoses and the major procedures performed on the patient, as attested to by the patient’s attending physician by virtue of his or her signature in the medical record. Anyone who misrepresents, falsifies, or conceals essential information required for payment of Federal funds, may be subject to fine, imprisonment, or civil penalty under applicable Federal laws.

(c) Completion of acknowledgement. The acknowledgement must be completed by the physician at the time that the physician is granted admitting privileges at the hospital, or before or at the time the physician admits his or her first patient. Existing acknowledgements signed by physicians already on staff remain in effect as long as the physician has admitting privileges at the hospital.

[60 FR 45847, Sept. 1, 1995]
§ 412.50 Furnishing of inpatient hospital services directly or under arrangements.

(a) The applicable payments made under the prospective payment systems, as described in subparts H and M of this part, are payment in full for all inpatient hospital services, as defined in § 409.10 of this chapter. Inpatient hospital services do not include the following types of services:

(1) Physician services that meet the requirements of § 415.102(a) of this chapter for payment on a fee schedule basis.

(2) Physician assistant services, as defined in section 1861(s)(2)(K)(i) of the Act.

(3) Nurse practitioner and clinical nurse specialist services, as defined in section 1861(s)(2)(K)(ii) of the Act.

(4) Certified nurse mid-wife services, as defined in section 1861(gg) of the Act.

(5) Qualified psychologist services, as defined in section 1861(ii) of the Act.

(6) Services of an anesthetist, as defined in § 410.69 of this chapter.

(b) HCFA does not pay any provider or supplier other than the hospital for services furnished to a beneficiary who is an inpatient, except for the services described in paragraphs (a)(1) through (a)(6) of this section.

(c) The hospital must furnish all necessary covered services to the beneficiary either directly or under arrangements (as defined in § 409.3 of this chapter).


§ 412.52 Reporting and recordkeeping requirements.

All hospitals participating in the prospective payment systems must meet the recordkeeping and cost reporting requirements of §§ 413.20 and 413.24 of this chapter.


Subpart D—Basic Methodology for Determining Prospective Payment Federal Rates for Inpatient Operating Costs

§ 412.60 DRG classification and weighting factors.

(a) Diagnosis-related groups. HCFA establishes a classification of inpatient hospital discharges by Diagnosis-Related Groups (DRGs).

(b) DRG weighting factors. HCFA assigns, for each DRG, an appropriate weighting factor that reflects the estimated relative cost of hospital resources used with respect to discharges classified within that group compared to discharges classified within other groups.

(c) Assignment of discharges to DRGs. HCFA establishes a methodology for classifying specific hospital discharges within DRGs which ensures that each hospital discharge is appropriately assigned to a single DRG based on essential data abstracted from the inpatient bill for that discharge.

(1) The classification of a particular discharge is based, as appropriate, on the patient’s age, sex, principal diagnosis (that is, the diagnosis established after study to be chiefly responsible for causing the patient’s admission to the hospital), secondary diagnoses, procedures performed, and discharge status.

(2) Each discharge is assigned to only one DRG (related, except as provided in paragraph (c)(3) of this section, to the

§ 412.62 Federal rates for inpatient operating costs for fiscal year 1984.

(a) General rule. HCFA determines national adjusted DRG prospective payment rates for operating costs, for each inpatient hospital discharge in fiscal year 1984 involving inpatient hospital services of a hospital in the United States subject to the prospective payment system under subpart B of this part, and determines regional adjusted DRG prospective payment rates for inpatient operating costs for such discharges in each region, for which payment may be made under Medicare Part A. Such rates are determined for hospitals located in urban or rural areas within the United States and within each such region, respectively, as described in paragraphs (b) through (k) of this section.

(b) Determining allowable individual hospital inpatient operating costs. HCFA determines the Medicare allowable operating costs per discharge of inpatient hospital services for each hospital in the data base for the most recent cost reporting period for which data are available.

(c) Updating for fiscal year 1984. HCFA updates each amount determined under paragraph (b) of this section for fiscal year 1984 by—

(1) Updating for fiscal year 1983 by the estimated average rate of change of hospital costs industry-wide between the cost reporting period used under paragraph (b) of this section and fiscal year 1983; and

(2) Projecting for fiscal year 1984 by the applicable percentage increase in the hospital market basket for fiscal year 1984.

(d) Standardizing amounts. HCFA standardizes the amount updated under paragraph (c) of this section for each hospital by—

(1) Adjusting for area variations in case mix among hospitals;

(2) Excluding an estimate of indirect medical education costs;

(3) Adjusting for area variations in hospital wage levels; and

(4) Adjusting for the effects of a higher cost of living for hospitals located in Alaska and Hawaii.

(e) Computing urban and rural averages. HCFA computes an average of the standardized amounts determined under paragraph (d) of this section for urban and rural hospitals in the United States and for urban and rural hospitals in each region.

(f) Geographic classifications. (1) For purposes of paragraph (e) of this section, the following definitions apply:

(i) The term region means one of the nine census divisions, comprising the...
fifty States and the District of Columbia, established by the Bureau of the Census for statistical and reporting purposes.

(ii) The term urban area means—

(A) A Metropolitan Statistical Area (MSA) or New England County Metropolitan Area (NECMA), as defined by the Executive Office of Management and Budget; or

(B) The following New England counties, which are deemed to be parts of urban areas under section 601(g) of the Social Security Amendments of 1983 (Pub. L. 98-21, 42 U.S.C. 1395ww (note)): Litchfield County, Connecticut; York County, Maine; Sagadahoc County, Maine; Merrimack County, New Hampshire; and Newport County, Rhode Island.

(iii) The term rural area means any area outside an urban area.

(iv) The phrase hospital reclassified as rural means a hospital located in a county that was part of an MSA or NECMA, as defined by the Executive Office of Management and Budget, but is not part of an MSA or NECMA as a result of an Executive Office of Management and Budget redesignation occurring after April 20, 1983.

(2) For hospitals within an MSA or NECMA that crosses census division boundaries, the following provisions apply:

(i) The MSA or NECMA is deemed to belong to the census division in which most of the hospitals within the MSA or NECMA are located.

(ii) If a hospital would receive a lower Federal rate because most of the hospitals are located in a census division with a lower Federal rate than the rate applicable to the census division in which the hospital is located, the payment rate will not be reduced for the hospital's cost reporting period beginning before October 1, 1984.

(iii) If an equal number of hospitals within the MSA or NECMA are located in each census division, such hospitals are deemed to be in the census division with the higher Federal rate.

(g) Adjusting the average standardized amounts. HCFA adjusts each of the average standardized amounts determined under paragraphs (c), (d), and (e) of this section by factors representing HCFA’s estimates of the following:

(1) The amount of payment that would have been made under Medicare Part B for nonphysician services to hospital inpatients during the first cost reporting period subject to prospective payment were it not for the fact that such services must be furnished either directly by hospitals or under arrangements in order for any Medicare payment to be made after September 30, 1983 (the effective date of §405.310(m) of this chapter).

(2) The amount of FICA taxes that would be incurred during the first cost reporting period subject to the prospective payment system, by hospitals that had not incurred such taxes for any or all of their employees during the base period described in paragraph (c) of this section.

(h) Reducing for value of outlier payments. HCFA reduces each of the adjusted average standardized amounts determined under paragraphs (c) through (g) of this section by a proportion equal to the proportion (estimated by HCFA) of the total amount of payments based on DRG prospective payment rates that are additional payments for outlier cases under subpart F of this part.

(i) Maintaining budget neutrality. (1) HCFA adjusts each of the reduced standardized amounts determined under paragraphs (c) through (h) of this section as required for fiscal year 1984 so that the estimated amount of aggregate payments made, excluding the hospital-specific portion (that is, the total of the Federal portion of transition payments, plus any adjustments and special treatment of certain classes of hospitals for Federal fiscal year 1984) is not greater or less than 25 percent of the payment amounts that would have been payable for the inpatient operating costs for those same hospitals for fiscal year 1984 under the Social Security Act as in effect on April 19, 1983.

(2) The aggregate payments considered under this paragraph exclude payments for per case review by a utilization and quality control peer review organization, as allowed under section 1866(a)(1)(F) of the Act.

(j) Computing Federal rates for inpatient operating costs for urban and rural hospitals in the United States and in each
region. For each discharge classified within a DRG, HCFA establishes a national prospective payment rate for inpatient operating costs and a regional prospective payment rate for inpatient operating costs for each region, as follows:

(1) For hospitals located in an urban area in the United States or in that region respectively, the rate equals the product of—

(i) The adjusted average standardized amount (computed under paragraphs (c) through (i) of this section) for hospitals located in an urban area in the United States or in that region; and

(ii) The weighting factor determined under §412.60(b) for that DRG.

(2) For hospitals located in a rural area in the United States or in that region respectively, the rate equals the product of—

(i) The adjusted average standardized amount (computed under paragraphs (c) through (i) of this section) for hospitals located in a rural area in the United States or in that region; and

(ii) The weighting factor determined under §412.60(b) for that DRG.

(k) Adjusting for different area wage levels. HCFA adjusts the proportion (as estimated by HCFA from time to time) of Federal rates computed under paragraph (j) of this section that are attributable to wages and labor-related costs, for area differences in hospital wage levels by a factor (established by HCFA) reflecting the relative hospital wage level in the geographic area (that is, urban or rural area as determined under the provisions of paragraph (f) of this section) of the hospital compared to the national average hospital wage level.

§412.63 Federal rates for inpatient operating costs for fiscal years after Federal fiscal year 1984.

(a) General rule. (1) HCFA determines a national adjusted prospective payment rate for inpatient operating costs for each inpatient hospital discharge in a Federal fiscal year after fiscal year 1984 involving inpatient hospital services of a hospital in the United States subject to the prospective payment system, and determines a regional adjusted prospective payment rate for operating costs for such discharges in each region, for which payment may be made under Medicare Part A.

(2) Each such rate is determined for hospitals located in urban or rural areas within the United States and within each such region respectively, as described in paragraphs (b) through (g) of this section.

(b) Geographic classifications. (1) For purposes of this section, the definitions set forth in §412.62(f) apply, except that, effective January 1, 2000, a hospital reclassified as rural may mean a reclassification that results from a geographic redesignation as set forth in §412.62(f)(1)(iv) or a reclassification that results from an urban hospital applying for reclassification as rural as set forth in §412.103.

(2) For hospitals within an MSA or NECMA that crosses census division boundaries, the following provisions apply:

(i) The MSA or NECMA is deemed to belong to the census division in which most of the hospitals within the MSA or NECMA are located.

(ii) A hospital that met the conditions specified in §412.62(f)(2)(ii) and therefore did not receive a lower Federal rate that would have applied for cost reporting periods beginning before October 1, 1984, receives the lower Federal rate applicable to all hospitals in the MSA or NECMA in which it is located effective with the hospital's cost reporting period that begins on or after October 1, 1984.

(iii) The higher Federal rate is payable to all hospitals in the MSA or NECMA if an equal number of hospitals within the MSA or NECMA are located in each census division.

(3) For discharges occurring on or after October 1, 1988, a hospital located in a rural county adjacent to one or more urban areas is deemed to be located in an urban area and receives the Federal payment amount for the urban area to which the greater number of workers in the county commute if the rural county would otherwise be considered part of an urban area, under the standards for designating MSAs or NECMAs if the commuting rates used
in determining outlying counties were determined on the basis of the aggregate number of resident workers who commute to (and, if applicable under the standards, from) the central county or central counties of all adjacent MSAs or NECMAs. These EOMB standards are set forth in the notice of final standards for classification of MSAs published in the Federal Register on January 3, 1980 (45 FR 956), and available from HCFA, East High Rise Building, room 132, 6325 Security Boulevard, Baltimore, Maryland 21207.

(4) For purposes of this section, any change in an MSA or NECMA designation is recognized on the October 1 following the effective date of the change.

(5) For discharges occurring on or after October 1, 1988, for hospitals that consist of two or more separately located inpatient hospital facilities the national adjusted prospective payment rate is based on the geographic location of the hospital facility at which the discharge occurs.

(c) Updating previous standardized amounts.

(1) HCFA computes an average standardized amount for hospitals in urban areas and rural areas within the United States, and urban areas and rural areas within each region.

(2) Each of those amounts is equal to the respective adjusted average standardized amount computed for fiscal year 1984 under §412.62(g)—

(i) Increased for fiscal year 1985 by the applicable percentage increase in the hospital market basket;

(ii) Adjusted by the estimated amount of Medicare payment for non-physician services furnished to hospital inpatients that would have been paid under Part B were it not for the fact that such services must be furnished either directly by hospitals or under arrangements;

(iii) Reduced by a proportion equal to the proportion (estimated by HCFA) of the total amount of prospective payments that are additional payment amounts attributable to outlier cases under subpart F of this part, and for discharges occurring on or after October 1, 1986, reduced by a proportion (estimated by HCFA) of the amount of payments that, based on the total amount of prospective payments for urban hospitals and the total amount of prospective payments for rural hospitals, are additional payments attributable to outlier cases in such hospitals under subpart F of this part.

(3) For fiscal year 1986 and thereafter, HCFA computes average standardized amounts for hospitals located in large urban areas, other urban areas, and rural areas. The term large urban area means an MSA with a population of more than 1,000,000 or an NECMA, with a population of more than 970,000 based on the most recent available population data published by the Bureau of the Census.
(d) Applicable percentage change for fiscal year 1986. (1) The applicable percentage change for fiscal year 1986 is—
(i) For discharges occurring on or after October 1, 1985 and before May 1, 1986, zero percent; and
(ii) For discharges occurring on or after May 1, 1986, one-half of one percent.
(2) For purposes of determining the standardized amounts for discharges occurring on or after October 1, 1986, the applicable percentage increase for fiscal year 1986 is deemed to have been one-half of one percent.

(e) Applicable percentage change for fiscal year 1987. The applicable percentage change for fiscal year 1987 is 1.15 percent.

(f) Applicable percentage change for fiscal year 1988. (1) The applicable percentage change for fiscal year 1988 is—
(i) For discharges occurring on or after October 1, 1987 and before November 21, 1987, zero percent;
(ii) For discharges occurring on or after November 21, 1987 and before April 1, 1988, 2.7 percent; and
(iii) For discharges occurring on or after April 1, 1988 and before October 1, 1988—
(A) 3.0 percent for hospitals located in rural areas;
(B) 1.5 percent for hospitals located in large urban areas; and
(C) 1.0 percent for hospitals located in other urban areas.
(2) For purposes of determining the standardized amounts for discharges occurring on or after October 1, 1988 (for Federal fiscal year 1989), the applicable percentage change for fiscal year 1988 is deemed to have been—
(i) 3.0 percent for hospitals located in rural areas;
(ii) 1.5 percent for hospitals located in large urban areas; and
(iii) 1.0 percent for hospitals located in other urban areas.

(g) Applicable percentage change for fiscal year 1989. The applicable percentage change for fiscal year 1989 is the percentage increase in the market basket index (as defined in §413.40(a)(3) of this chapter)—
(1) Minus 1.5 percentage points for hospitals located in rural areas;
(2) Minus 2.0 percentage points for hospitals in large urban areas; and
(3) Minus 2.5 percentage points for hospitals in other urban areas.

(h) Applicable percentage change for fiscal year 1990. (1) The applicable percentage change for fiscal year 1990 is—
(i) For discharges occurring on or after October 1, 1989 and before January 1, 1990, 5.5 percent; and
(ii) For discharges occurring on or after January 1, 1990 and before October 1, 1990—
(A) 9.72 percent for hospitals located in rural areas;
(B) 5.62 percent for hospitals located in large urban areas; and
(C) 4.97 percent for hospitals located in other urban areas.
(2) For purposes of determining the standardized amounts for discharges occurring on or after October 1, 1990, the applicable percentage change for fiscal year 1990 is deemed to have been the percentage change provided for in paragraph (h)(1)(ii) of this section.

(i) Applicable percentage change for fiscal year 1991. (1) The applicable percentage change for fiscal year 1991 is—
(i) For discharges occurring on or after October 1, 1990 and before October 21, 1990, 5.2 percent; and
(ii) For discharges occurring on or after October 21, 1990 and before January 1, 1991, 0.0 percent; and
(iii) For discharges occurring on or after January 1, 1991 and before October 1, 1991—
(A) 4.5 percent for hospitals located in rural areas; and
(B) 3.2 percent for hospitals located in large urban areas and other urban areas.
(2) For purposes of determining the standardized amounts for discharges occurring on or after October 1, 1991, the applicable percentage change for fiscal year 1991 is deemed to have been the percentage change provided for in paragraph (i)(1)(iii) of this section.

(j) Applicable percentage change for fiscal year 1992. The applicable percentage change for fiscal year 1992 is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a)(3) of this chapter)—
(1) Minus 0.6 percentage points for hospitals located in rural areas.
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(2) Minus 1.6 percentage points for hospitals located in large urban areas and other urban areas.

(k) Applicable percentage change for fiscal year 1993. The applicable percentage change for fiscal year 1993 is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a)(3) of this chapter)—

(1) Minus 0.55 percentage points for hospitals located in rural areas.

(2) Minus 1.55 percentage points for hospitals located in large urban areas and other urban areas.

(l) Applicable percentage change for fiscal year 1994. The applicable percentage change for fiscal year 1994 is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this chapter)—

(1) Minus 1.0 percentage point for hospitals located in rural areas.

(2) Minus 2.5 percentage points for hospitals located in large urban areas and other urban areas.

(m) Applicable percentage change for fiscal year 1995. The applicable percentage change for fiscal year 1995 is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this chapter)—

(1) Plus, for hospitals located in rural areas, the percentage increase necessary so that the average standardized amounts computed under paragraph (c) through (i) of this section are equal to the average standardized amounts for hospitals located in an urban area other than a large urban area.

(2) Minus 2.5 percentage points for hospitals located in large urban areas and other urban areas.

(n) Applicable percentage change for fiscal year 1996. The applicable percentage change for fiscal year 1996 is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this chapter) minus 2.0 percentage points for all areas.

(o) Applicable percentage change for fiscal year 1997. The applicable percentage change for fiscal year 1997 is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this chapter) minus 0.5 percentage point for all areas.

(p) Applicable percentage change for fiscal year 1998. The applicable percentage change for fiscal year 1998 is 0 percent for hospitals in all areas.

(q) Applicable percentage change for fiscal year 1999. The applicable percentage change for fiscal year 1999 is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this subchapter) minus 1.9 percentage points for hospitals in all areas.

(r) Applicable percentage change for fiscal year 2000. The applicable percentage change for fiscal year 2000 is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this chapter) minus 1.8 percentage points for hospitals in all areas.

(s) Applicable percentage change for fiscal year 2001. The applicable percentage change for fiscal year 2001 is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this subchapter) for sole community hospitals and the increase in the market basket index minus 1.1 percentage points for other hospitals in all areas.

(t) Applicable percentage change for fiscal year 2002. The applicable percentage change for fiscal year 2002 is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this subchapter) minus 1.1 percentage points for hospitals in all areas.

(u) Applicable percentage change for fiscal year 2003 and for subsequent years. The applicable percentage change for fiscal year 2003 and for subsequent years is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a)) for hospitals in all areas.

(v) Maintaining budget neutrality for fiscal year 1985. (1) For fiscal year 1985, HCFA will adjust each of the reduced standardized amounts determined under paragraph (c) of this section as required for fiscal year 1985 to ensure that the estimated amount of aggregate payments made, excluding the hospital-specific portion (that is, the total of the Federal portion of transition payments, plus any adjustments...
and special treatment of certain classes of hospitals for fiscal year 1985 is not greater or less than 50 percent of the payment amounts that would have been payable for the inpatient operating costs for those same hospitals for fiscal year 1985 under the law as in effect on April 19, 1983.

(2) The aggregate payments considered under this paragraph exclude payments for per case review by a utilization and quality control peer review organization, as allowed under section 1866(a)(3)(F) of the Act.

(w) Computing Federal rates for inpatient operating costs for hospitals located in large urban and other areas. For each discharge classified within a DRG, HCFA establishes for the fiscal year a national prospective payment rate and a regional prospective payment rate for inpatient operating costs, for each region, as follows:

(1) For hospitals located in a large urban area in the United States or that region respectively, the rate equals the product of—

(i) The adjusted average standardized amount (computed under paragraph (c) of this section) for the fiscal year for hospitals located in a large urban area in the United States or in that region; and

(ii) The weighting factor determined under §412.60(b) for that DRG.

(2) For hospitals located in an other area in the United States or that region respectively, the rate equals the product of—

(i) The adjusted average standardized amount (computed under paragraph (c) of this section) for the fiscal year for hospitals located in an other area in the United States or in that region; and

(ii) The weighting factor (determined under §412.60(b)) for that DRG.

(x) Adjusting for different area wage levels. (1) HCFA adjusts the proportion (as estimated by HCFA from time to time) of Federal rates for inpatient operating costs computed under paragraph (i) of this section that are attributable to wages and labor-related costs for area differences in hospital wage levels by a factor (established by HCFA based on survey data) reflecting the relative level of hospital wages and wage-related costs in the geographic area (that is, urban or rural area as determined under the provisions of paragraph (b) of this section) of the hospital compared to the national average level of hospital wages and wage-related costs. The wage index is updated annually.

(2)(i) HCFA makes a midyear correction to the wage index for an area only if a hospital can show that—

(A) The intermediary or HCFA made an error in tabulating the hospital’s data; and

(B) The hospital could not have known about the error, or did not have the opportunity to correct the error, before the beginning of the Federal fiscal year.

(ii) A midyear correction to the wage index is effective prospectively from the date the change is made to the wage index.

(3) Revisions to the wage index resulting from midyear corrections to the wage index values are incorporated in the wage index values for other areas at the beginning of the next Federal fiscal year.

(4) The effect on program payments of midyear corrections to the wage index values is taken into account in establishing the standardized amounts for the following Federal fiscal year.

(5) If a judicial decision reverses a HCFA denial of a hospital’s wage data revision request, HCFA pays the hospital by applying a revised wage index that reflects the revised wage data as if HCFA’s decision had been favorable rather than unfavorable.

[50 FR 12741, Mar. 29, 1985]

EDITORIAL NOTE: For Federal Register citations affecting §412.63, see the List of Sections Affected in the finding Aids section of this volume.

Subpart E—Determination of Transition Period Payment Rates for the Prospective Payment System for Inpatient Operating Costs

412.70 General description.

For discharges occurring on or after April 1, 1988, and before October 1, 1996, payments to a hospital are based on the greater of the national average standardized amount or the sum of 85
§ 412.71 Determination of base-year inpatient operating costs.

(a) Base-year costs. (1) For each hospital, the intermediary will estimate the hospital’s Medicare Part A allowable inpatient operating costs, as described in §412.2(c), for the 12-month or longer cost reporting period ending on or after September 30, 1982 and before September 30, 1983.

(2) If the hospital’s last cost reporting period ending before September 30, 1983 is for less than 12 months, the base period will be the hospital’s most recent 12-month or longer cost reporting period ending before such short reporting period, with an appropriate adjustment for inflation. (The rules applicable to new hospitals are set forth in §412.74.)

(b) Modifications to base-year costs. Prior to determining the hospital-specific rate, the intermediary will adjust the hospital’s estimated base-year inpatient operating costs, as necessary, to include malpractice insurance costs in accordance with §413.53(a)(1)(i) of this chapter, and exclude the following:

(1) Medical education costs as described in §413.85 of this chapter.

(2) Capital-related costs as described in §413.130 of this chapter.

(3) Kidney acquisition costs incurred by hospitals approved as renal transplantation centers as described in §412.100. Kidney acquisition costs in the base year will be determined by multiplying the hospital’s average kidney acquisition cost per kidney times the number of kidney transplants covered by Medicare Part A during the base period.

(4) Higher costs that were incurred for purposes of increasing base-year costs.

(5) One-time nonrecurring higher costs or revenue offsets that have the effect of distorting base-year costs as an appropriate basis for computing the hospital-specific rate.

(6) Higher costs that result from changes in hospital accounting principles initiated in the base year.

(7) The costs of qualified nonphysician anesthetists’ services, as described in §412.113(c).

(c) Hospital’s request for adjustment of base-year inpatient operating costs. (1) Before the date it becomes subject to the prospective payment system for inpatient operating costs, a hospital may request the intermediary to further adjust its estimated base-period costs to take into account the following:

(i) Services paid for under Medicare Part B during the hospital’s base year that will be paid for under prospective payments. The base-year costs may be increased to include estimated payments for certain services previously billed as physicians’ services before the effective date of §415.102(a) of this chapter, and estimated payments for nonphysicians’ services that were not furnished either directly or under arrangements before October 1, 1983 (the effective date of §405.310(m) of this chapter), but may not include the costs of anesthetists’ services for which a physician employer continues to bill under §405.553(b)(4) of this chapter.

(ii) The payment of FICA taxes during cost reporting periods subject to the prospective payment system, if the hospital had not paid such taxes for all its employees during its base period and will be required to participate effective January 1, 1984.

(2) If a hospital requests that its base-period costs be adjusted under paragraph (c)(1) of this section, it must timely provide the intermediary with sufficient documentation to justify the adjustment, and adequate data to compute the adjusted costs. The intermediary decides whether to use part or all of the data on the basis of audit, survey and other information available.

(d) Intermediary’s determination. The intermediary uses the best data available at the time in estimating each hospital’s base-year costs and the modifications to those costs authorized by paragraphs (b) and (c) of this section. The intermediary’s estimate of base-year costs and modifications thereto is final and may not be changed after the first day of the first
§ 412.72 Modification of base-year costs.

(a) Bases for modification of base-year costs. Base-year costs as determined under §412.71(d) may be modified under the following circumstances:

(1) Inadvertent omissions. (i) A hospital that becomes subject to the prospective payment system beginning on or after October 1, 1983 and before November 16, 1983 has until November 15, 1983 to request its intermediary to reestimate its base-period costs to take into account inadvertent omissions in its previous submissions to the intermediary related to changes made by the prospective payment legislation for purposes of estimating the base-period costs.

(ii) The intermediary may also initiate changes to the estimation—

(A) For any reason before the date the hospital becomes subject to prospective payment; and

(B) Before November 16, 1983, for corrections to take into account inadvertent omissions in the hospital’s previous submissions related to changes made by the prospective payment legislation for purposes of estimating the base-period costs.

(iii) Such omissions pertain to adjustments to exclude capital-related costs and the direct medical education costs of approved educational activities and to adjustments specified in §412.71(c).

(iv) The intermediary must notify the provider of any change to the hospital-specific amount as a result of the provider’s request within 30 days of receipt of the additional data.

(v) Any change to base-period costs made under this paragraph (a)(1) will be made effective retroactively, beginning with the first day of the affected hospital’s fiscal year.

(2) Correction of mathematical errors of calculations. (i) The hospital must report mathematical errors of calculations to the intermediary within 90 days of the intermediary’s notification to the hospital of the hospital’s payments rates.

(ii) The intermediary may also identify such errors and initiate their correction during this period.

(iii) The intermediary will either make an appropriate adjustment or notify the hospital that no adjustment is warranted within 30 days of receipt of the hospital’s report of an error.

(iv) Corrections of errors of calculation will be effective with the first day of the hospital’s first cost reporting period subject to the prospective payment system.

(3) Recognition of additional costs. (i) The intermediary may adjust base-period costs to take into account additional costs recognized as allowable costs for the hospital’s base year as the result of any of the following:

(A) A reopening and revision of the hospital’s base-year notice of amount of program reimbursement under §§405.1885 through 405.1889 of this chapter.

(B) A prehearing order or finding issued during the provider payment appeals process by the appropriate reviewing authority under §405.1821 or §405.1853 of this chapter that resolved a matter at issue in the hospital’s base-year notice of amount of program reimbursement.

(C) An affirmation, modification, or reversal of a Provider Reimbursement Review Board decision by the Administrator of HCFA under §405.1875 of this chapter that resolved a matter at issue in the hospital’s base-year notice of amount of program reimbursement.

(D) An administrative or judicial review decision under §§405.1831, 405.1871, or 405.1877 of this chapter that is final and no longer subject to review under applicable law or regulations by a higher reviewing authority, and that resolved a matter at issue in the hospital’s base-year notice of amount of program reimbursement.

(B) A prehearing order or finding issued during the provider payment appeals process by the appropriate reviewing authority under §405.1821 or §405.1853 of this chapter that resolved a matter at issue in the hospital’s base-year notice of amount of program reimbursement.

(C) An affirmation, modification, or reversal of a Provider Reimbursement Review Board decision by the Administrator of HCFA under §405.1875 of this chapter that resolved a matter at issue in the hospital’s base-year notice of amount of program reimbursement.

(D) An administrative or judicial review decision under §§405.1831, 405.1871, or 405.1877 of this chapter that is final and no longer subject to review under applicable law or regulations by a higher reviewing authority, and that resolved a matter at issue in the hospital’s base-year notice of amount of program reimbursement.

(E) The intermediary will either make an appropriate adjustment or notify the hospital that no adjustment is warranted within 30 days of receipt of the hospital’s report of an error.
§412.73  Determination of the hospital-specific rate based on a Federal fiscal year 1982 base period.

(a) Costs on a per discharge basis. The intermediary will determine the hospital's estimated adjusted base-year operating cost per discharge by dividing the total adjusted operating costs by the number of discharges in the base period.

(b) Case-mix adjustment. The intermediary will divide the adjusted base-year costs by the hospital's 1981 case-mix index. If the hospital's case-mix index is statistically unreliable (as determined by HCFA), the hospital's base-year costs will be divided by the lower of the following:

(1) The hospital's estimated case-mix index.

(2) The average case-mix index for the appropriate classifications of all hospitals subject to cost limits established under §413.30 of this chapter for cost reporting periods beginning on or after October 1, 1982 and before October 1, 1983.

(c) Updating base-year costs—(1) For Federal fiscal year 1984. The case-mix adjusted base-year cost per discharge will be updated by the applicable updating factor, that is, the rate-of-increase percentage determined under §413.40(c)(3) of this chapter, as adjusted for budget neutrality.

(2) For Federal fiscal year 1985. The amount determined under paragraph (c)(1) of this section will be updated by
the applicable updating factor, as adjusted for budget neutrality.

(3) For Federal fiscal year 1986. (i) The amount determined under paragraph (c)(2) of this section is updated by—
   (A) Zero percent for the first seven months of the hospital’s cost reporting period; and
   (B) One-half of one percent for the remaining five months of the hospital’s cost reporting period.

(ii) For purposes of determining the updated base-year costs for cost reporting periods beginning in Federal fiscal year 1987 (that is, on or after October 1, 1986 and before October 1, 1987), the update factor for the previous cost reporting period is deemed to have been one-half of one percent.

(4) For Federal fiscal year 1987. The amount determined under paragraph (c)(3)(ii) of this section is updated by 1.15 percent.

(5) For Federal fiscal year 1988. (i) For purposes of determining the prospective payment rates for sole community hospitals under §412.92(d) for cost reporting periods beginning in Federal fiscal year 1988 (that is, on or after October 1, 1987 and before October 1, 1988), the base-year cost per discharge is updated as follows:
   (A) For the first 51 days of the hospital’s cost reporting period, by zero percent.
   (B) For the next 132 days of the hospital’s cost reporting period, by 2.7 percent.
   (C) For the remainder of the hospital’s cost reporting period, by—
      (1) 3.0 percent for hospitals located in rural areas;
      (2) 1.5 percent for hospitals located in large urban areas; and
      (3) 1.0 percent for hospitals located in other urban areas.

(ii) For discharges occurring on or after October 21, 1990 and before January 1, 1991, the base-period cost per discharge, updated as set forth in paragraph (c)(7)(i) of this section, is reduced by 5.5 percent.

(iii) For purposes of determining the updated base-period costs for cost reporting periods beginning in Federal fiscal year 1991 (that is, beginning on or after October 1, 1990 and before October 1, 1991), the update factor for the cost reporting period beginning during Federal fiscal year 1990 is deemed to have been the percentage change provided for in paragraph (c)(7)(i)(B) of this section.

(6) For Federal fiscal year 1989. For cost reporting periods beginning in Federal fiscal year 1989, the update factor is determined using the methodology set forth in §412.63(g).

(7) For Federal fiscal year 1990. (i) Except as described in paragraph (c)(7)(ii) of this section, for cost reporting periods beginning in Federal fiscal year 1990, the base-period cost per discharge is updated as follows:
   (A) For cost reporting periods beginning on or after October 1, 1989 and before January 1, 1990, by 5.5 percent for discharges occurring before January 1, 1990 and by the factors set forth in paragraph (c)(7)(i)(B) of this section for discharges occurring on or after January 1, 1990.
   (B) For cost reporting periods beginning on or after January 1, 1990 and before October 1, 1990, by—
      (1) 9.72 percent for hospitals located in rural areas;
      (2) 5.62 percent for hospitals located in large urban areas; and
      (3) 4.97 percent for hospitals located in other urban areas.

(ii) For discharges occurring on or after October 21, 1990 and before January 1, 1991, the base-period cost per discharge is updated by 0.0 percent.

(8) For Federal fiscal year 1991. (i) Except as described in paragraph (c)(8)(ii) of this section, for cost reporting periods beginning in Federal fiscal year 1991, the base-period cost per discharge is updated by 5.2 percent.

(ii) For discharges occurring on or after October 21, 1990 and before January 1, 1991, the base-period cost per discharge is updated by 0.0 percent.
(iii) For purposes of determining the updated base period costs for cost reporting periods beginning in Federal fiscal year 1992, the update factor for the cost reporting period beginning during Federal fiscal year 1991 is deemed to have been the percentage change provided for in paragraph (c)(8)(i) of this section.

(9) For Federal fiscal years 1992 and 1993, the update factor is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this chapter).

(10) For Federal fiscal year 1994. For Federal fiscal year 1994, the update factor is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of the chapter) minus 2.3 percentage points. For purposes of determining the hospital-specific rate for Federal fiscal year 1994 and subsequent years, this update factor is adjusted to take into account the portion of the 12-month cost reporting period beginning during Federal fiscal year 1993 that occurs in Federal fiscal year 1994.

(11) For Federal fiscal year 1995. For Federal fiscal year 1995, the update factor is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this chapter) minus 2.2 percentage points.

(12) For Federal fiscal years 1996 through 2000. For Federal fiscal years 1996 through 2000, the update factor is the applicable percentage change for other prospective payment hospitals in each respective year as set forth in §§412.63(n) through (r).

(13) For Federal fiscal year 2001. For Federal fiscal year 2001, the update factor is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this chapter).

(14) For Federal fiscal year 2002. For Federal fiscal year 2002, the update factor is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this chapter) minus 1.1 percentage points.

(15) For Federal fiscal year 2003 and for subsequent years. For Federal fiscal year 2003 and subsequent years, the update factor is the percentage increase in the market basket index for prospective payment hospitals (as defined in §413.40(a) of this chapter).

(d) Budget neutrality—(1) For Federal fiscal year 1984. For cost reporting periods beginning on or after October 1, 1983 and before October 1, 1984, HCFA adjusts the target rate percentage used under paragraph (c)(1) of this section. This adjustment is based on a factor actuarially estimated to ensure that the estimated amount of aggregate Medicare payments based on the hospital-specific portion of the transition payment rates is neither greater nor less than 75 percent of the amounts that would have been payable for the inpatient operating costs for those same hospitals for fiscal year 1984 under the law in effect before April 20, 1983.

(2) For Federal fiscal year 1985. For cost reporting periods beginning on or after October 1, 1984 and before October 1, 1985, HCFA adjusts the target rate percentage used under paragraph (c)(2) of this section. This adjustment is based on a factor actuarially estimated to ensure that the estimated amount of aggregate Medicare payment based on the hospital-specific portion of the transition payment rates is neither greater nor less than 50 percent of the amounts that would have been payable for the inpatient operating costs for those same hospitals for fiscal year 1985 under the Social Security Act as in effect on April 19, 1983.

(e) DRG adjustment. The applicable hospital-specific cost per discharge is multiplied by the appropriate DRG weighting factor to determine the hospital-specific base payment amount (target amount) for a particular covered discharge.

§ 412.75 Determination of the hospital-specific rate for inpatient operating costs based on a Federal fiscal year 1987 base period.

(a) Base-period costs—(1) General rule. Except as provided in paragraph (a)(2) of this section, for each hospital, the intermediary determines the hospital’s Medicare part A allowable inpatient operating costs, as described in §412.2(c), for the 12-month or longer cost reporting period ending on or after September 30, 1987 and before September 30, 1988.

(2) Exceptions. (i) If the hospital’s last cost reporting period ending before September 30, 1988 is for less than 12 months, the base period is the hospital’s most recent 12-month or longer cost reporting period ending before the short period report.

(ii) If the hospital does not have a cost reporting period ending on or after September 30, 1987 and before September 30, 1988 and does have a cost reporting period beginning on or after October 1, 1986 and before October 1, 1987, that cost reporting period is the base period unless the cost reporting period is for less than 12 months. In that case, the base period is the hospital’s most recent 12-month or longer cost reporting period ending before the short cost reporting period.

(b) Costs on a per discharge basis. The intermediary determines the hospital’s average base-period operating cost per discharge by dividing the total operating costs by the number of discharges in the base period. For purposes of this section, a transfer as defined in §412.4(b) is considered to be a discharge.

(c) Case-mix adjustment. The intermediary divides the average base-period cost per discharge by the hospital’s case-mix index for the base period.

(d) Updating base-period costs. For purposes of determining the updated base-period costs for cost reporting periods beginning in Federal fiscal year 1988, the update factor is determined using the methodology set forth in §412.73(c)(15).

(e) DRG adjustment. The applicable hospital-specific cost per discharge is multiplied by the appropriate DRG weighting factor to determine the hospital-specific base payment amount (target amount) for a particular covered discharge.

(f) Notice of hospital-specific rate. The intermediary furnishes the hospital a notice of its hospital-specific rate, which contains a statement of the hospital’s Medicare part A allowable inpatient operating costs, number of Medicare discharges, and case-mix index adjustment factor used to determine the hospital’s cost per discharge for the Federal fiscal year 1987 base period.

(g) Right to administrative and judicial review. An intermediary’s determination of the hospital-specific rate for a hospital is subject to administrative and judicial review. Review is available to a hospital upon receipt of the notice of the hospital-specific rate. This notice is treated as a final intermediary determination of the amount of program reimbursement for purposes of subpart R of part 405 of this chapter, governing provider reimbursement determinations and appeals.

(h) Modification of hospital-specific rate. (1) The intermediary recalculates the hospital-specific rate to reflect the following:

(i) Any modifications that are determined as a result of administrative or judicial review of the hospital-specific rate determinations; or

(ii) Any additional costs that are recognized as allowable costs for the hospital’s base period as a result of administrative or judicial review of the base-period notice of amount of program reimbursement.

(2) With respect to either the hospital-specific rate determination or the amount of program reimbursement determination, the actions taken on administrative or judicial review that provide a basis for recalculations of the hospital-specific rate include the following:

(i) A reopening and revision of the hospital’s base-period notice of amount of program reimbursement under §§405.1885 through 405.1889 of this chapter.

(ii) A prehearing order or finding issued during the provider payment appeals process by the appropriate reviewing authority under §405.1821 or §405.1853 of this chapter that resolved a matter at issue in the hospital’s base-
§ 412.77 Determination of the hospital-specific rate for inpatient operating costs for certain sole community hospitals based on a Federal fiscal year 1996 base period.

(a) Applicability. (1) This section applies to a hospital that has been designated as a sole community hospital, as described in §412.92, that received payment for its cost reporting period beginning during 1999 based on its hospital-specific rate for either fiscal year 1982 under §412.73 or fiscal year 1987 under §412.75, and that elects under paragraph (a)(2) of this section to be paid based on a fiscal year 1996 base period. If the 1996 hospital-specific rate exceeds the hospital-specific rates for either fiscal year 1982 or 1987, unless the hospital elects to the contrary, this rate will be used in the payment formula set forth under §412.92(d)(1).

(2) Hospitals that are otherwise eligible for but elect not to receive payment on the basis of their Federal fiscal year 1996 updated costs per case must notify their fiscal intermediary of this decision prior to the end of their cost reporting period beginning on or after October 1, 2000, for which such payments would otherwise be made. If a hospital does not make the notification to its fiscal intermediary before the end of the cost reporting period, the hospital is deemed to have elected to have section 1886(b)(3)(I) of the Act apply to the hospital.

(3) This section applies only to cost reporting periods beginning on or after October 1, 2000.

(4) The formula for determining the hospital-specific costs for hospitals described under paragraph (a)(1) of this section is set forth in paragraph (f) of this section.

(b) Based costs for hospitals subject to fiscal year 1996 rebasing. (1) General rule. Except as provided in paragraph (b)(2) of this section, for each hospital eligible under paragraph (a) of this section, the intermediary determines the hospital's Medicare Part A allowable inpatient operating costs, as described in §412.2(c), for the 12-month or longer cost reporting period ending on or after September 30, 1996 and before September 30, 1997, and computes the hospital-specific rate for purposes of determining prospective payment rates for inpatient operating costs as determined under §412.92(d).

(2) Exceptions. (i) If the hospital's last cost reporting period ending before September 30, 1997 is for less than 12 months, the base period is the hospital's most recent 12-month or longer cost reporting period ending before the short period report.

(ii) If the hospital does not have a cost reporting period ending on or after September 30, 1996 and before September 30, 1997, and does have a cost reporting period beginning on or after October 1, 1995 and before October 1, 1996, that cost reporting period is the base period unless the cost reporting period is for less than 12 months. If that cost reporting period is for less than 12 months, the base period is the hospital's most recent 12-month or longer cost reporting period ending before the short period report.

(iii) If the hospital does not have a cost reporting period ending on or after September 30, 1996 and before September 30, 1997, and does have a cost reporting period beginning on or after October 1, 1995 and before October 1, 1996, that cost reporting period is the base period unless the cost reporting period is for less than 12 months. If that cost reporting period is for less than 12 months, the base period is the hospital's most recent 12-month or longer cost reporting period ending before the short period report. If a hospital has no cost reporting period beginning in fiscal year 1996, the hospital will not have a hospital-specific rate based on fiscal year 1996.
(c) Costs on a per discharge basis. The intermediary determines the hospital’s average base-period operating cost per discharge by dividing the total operating costs by the number of discharges in the base period. For purposes of this section, a transfer as defined in §412.4(b) is considered to be a discharge.

(d) Case-mix adjustment. The intermediary divides the average base-period cost per discharge by the hospital’s case-mix index for the base period.

(e) Updating base-period costs. For purposes of determining the updated base-period costs for cost reporting periods beginning in Federal fiscal year 1996, the update factor is determined using the methodology set forth in §412.73(c)(12) through (c)(15).

(f) DRG adjustment. The applicable hospital-specific cost per discharge is multiplied by the appropriate DRG weighting factor to determine the hospital-specific base payment amount (target amount) for a particular covered discharge.

(g) Notice of hospital-specific rates. The intermediary furnishes a hospital eligible for rebasing a notice of the hospital-specific rate as computed in accordance with this section. The notice will contain a statement of the hospital’s Medicare Part A allowable inpatient operating costs, the number of Medicare discharges, and the case-mix index adjustment factor used to determine the hospital’s cost per discharge for the Federal fiscal year 1996 base period.

(h) Right to administrative and judicial review. An intermediary’s determination of the hospital-specific rate for a hospital is subject to administrative and judicial review. Review is available to a hospital upon receipt of the notice of the hospital-specific rate. This notice is treated as a final intermediary determination of the amount of program reimbursement for purposes of subpart R of part 405 of this chapter.

(i) Modification of hospital-specific rate. (1) The intermediary recalculates the hospital-specific rate to reflect the following:

(i) Any modifications that are determined as a result of administrative or judicial review of the hospital-specific rate determinations; or

(ii) Any additional costs that are recognized as allowable costs for the hospital’s base period as a result of administrative or judicial review of the base-period notice of amount of program reimbursement.

(2) With respect to either the hospital-specific rate determination or the amount of program reimbursement determination, the actions taken on administrative or judicial review that provide a basis for the recalculations of the hospital-specific rate include the following:

(i) A reopening and revision of the hospital’s base-period notice of amount of program reimbursement under §§405.1865 through 405.1889 of this chapter.

(ii) A prehearing order or finding issued during the provider payment appeals process by the appropriate reviewing authority under §405.1821 or §405.1853 of this chapter that resolved a matter at issue in the hospital’s base-period notice of amount of program reimbursement.

(iii) An affirmation, modification, or reversal of a Provider Reimbursement Review Board decision by the Administrator of HCFA under §405.1875 of this chapter that resolved a matter at issue in the hospital’s base-period notice of amount of program reimbursement.

(iv) An administrative or judicial review decision under §405.1831, §405.1871, or §405.1877 of this chapter that is final and no longer subject to review under applicable law or regulations by a higher reviewing authority, and that resolved a matter at issue in the hospital’s base-period notice of amount of program reimbursement.

(v) A final, nonappealable court judgment relating to the base-period costs.

(3) The adjustments to the hospital-specific rate made under paragraphs (i)(1) and (i)(2) of this section are effective retroactively to the time of the intermediary’s initial determination of the rate.

[65 FR 47106, Aug. 1, 2000]
§ 412.78 Recovery of excess transition period payment amounts resulting from unlawful claims.

If a hospital's base-year costs, as estimated for purposes of determining the hospital-specific portion, are determined, by criminal conviction or imposition of a civil money penalty or assessment, to include costs that were unlawfully claimed, the hospital's base-period costs are adjusted to remove the effect of the excess costs, and HCFA recovers both the excess costs reimbursed for the base period and the additional amounts paid due to the inappropriate increase of the hospital-specific portion of the hospital's transition payment rates.


Subpart F—Payment for Outlier Cases

412.80 General provisions.

(a) Basic rule—(1) Discharges occurring on or after October 1, 1994 and before October 1, 1997. For discharges occurring on or after October 1, 1994, and before October 1, 1997, except as provided in paragraph (b) of this section concerning transferring hospitals, HCFA provides for additional payment, beyond standard DRG payments, to a hospital for covered inpatient hospital services furnished to a Medicare beneficiary if either of the following conditions is met:

(i) The beneficiary's length-of-stay (including days at the SNF level of care if a SNF bed is not available in the area) exceeds the mean length-of-stay for the applicable DRG by the lesser of the following:

(A) A fixed number of days, as specified by HCFA; or

(B) A fixed number of standard deviations, as specified by HCFA.

(ii) The beneficiary's length-of-stay does not exceed criteria established under paragraph (a)(1)(i) of this section, but the hospital's charges for covered services furnished to the beneficiary, adjusted to operating costs and capital costs by applying cost-to-charge ratios as described in §412.84(h), exceed the DRG payment for the case plus a fixed dollar amount (adjusted for geographic variation in costs) as specified by HCFA.

(2) Discharges occurring on or after October 1, 1997. For discharges occurring on or after October 1, 1997, except as provided in paragraph (b) of this section concerning transfers, HCFA provides for additional payment, beyond standard DRG payments, to a hospital for covered inpatient hospital services furnished to a Medicare beneficiary if the hospital's charges for covered services, adjusted to operating costs and capital costs by applying cost-to-charge ratios as described in §412.84(h), exceed the DRG payment for the case plus a fixed dollar amount (adjusted for geographic variation in costs) as specified by HCFA.

(b) Outlier cases in transferring hospitals. HCFA provides cost outlier payments to a transferring hospital for cases paid in accordance with §412.4(f), if the hospital's charges for covered services furnished to the beneficiary, adjusted to costs by applying cost-to-charge ratios as described in §412.84(h), exceed the DRG payment for the case plus a fixed dollar amount (adjusted for geographic variation in costs) as specified by HCFA, divided by the geometric mean length of stay for the DRG, and multiplied by an applicable factor determined as follows:

(1) For transfer cases paid in accordance with §412.4(f)(1), the applicable factor is equal to the length of stay plus 1 day.

(2) For transfer cases paid in accordance with §412.4(f)(2), the applicable factor is equal to 0.5 plus the product of the length of stay plus 1 day multiplied by 0.5.

(c) Publication and revision of outlier criteria. HCFA will issue threshold criteria for determining outlier payment in the annual notice of the prospective payment rates published in accordance with §412.8(b).


§ 412.82 Payment for extended length-of-stay cases (day outliers).

(a) For discharges occurring before October 1, 1997, if the hospital stay reflected by a discharge includes covered days of care beyond the applicable
threshold criterion, the intermediary will make an additional payment, on a per diem basis, to the discharging hospital for those days. A special request or submission by the hospital is not necessary to initiate this payment. However, a hospital may request payment for day outliers before the medical review required in paragraph (b) of this section.

(b) The PRO must review and approve to the extent required by HCFA—
(1) The medical necessity and appropriateness of the admission and outlier services in the context of the entire stay;
(2) The validity of the diagnostic and procedural coding; and
(3) The granting of grace days.

(c) Except as provided in §412.86, the per diem payment made under paragraph (a) of this section is derived by taking a percentage of the average per diem payment for the applicable DRG, as calculated by dividing the Federal prospective payment rate for inpatient operating costs and inpatient capital-related costs determined under subpart D of this part, by the arithmetic mean length of stay for that DRG. HCFA issues the applicable percentage of the average per diem payment in the annual publication of the prospective payment rates in accordance with §412.8(b).

(d) Any days in a covered stay identified as noncovered reduce the number of days reimbursed at the day outlier rate but not to exceed the number of days that occur after the day outlier threshold.

§412.84 Payment for extraordinarily high-cost cases (cost outliers).

(a) A hospital may request its intermediary to make an additional payment for inpatient hospital services that meet the criteria established in accordance with §412.80(a).

(b) The hospital must request additional payment—
(1) With initial submission of the bill; or
(2) Within 60 days of receipt of the intermediary's initial determination.

(c) Except as specified in paragraph (e) of this section, an additional payment for a cost outlier case is made prior to medical review.

(d) As described in paragraph (f) of this section, the PRO reviews a sample of cost outlier cases after payment. The charges for any services identified as uncovered through this review are denied and any outlier payment made for these services are recovered, as appropriate, after a determination as to the provider's liability has been made.

(e) If the PRO finds a pattern of inappropriate utilization by a hospital, all cost outlier cases from that hospital are subject to medical review, and this review may be conducted prior to payment until the PRO determines that appropriate corrective actions have been taken.

(f) The PRO reviews the cost outlier cases, using the medical records and itemized charges, to verify the following:
(1) The admission was medically necessary and appropriate.
(2) Services were medically necessary and delivered in the most appropriate setting.
(3) Services were ordered by the physician, actually furnished, and not duplicatively billed.
(4) The diagnostic and procedural codings are correct.

(g) The intermediary bases the operating and capital costs of the discharge on the billed charges for covered inpatient services adjusted by the cost to charge ratios applicable to operating and capital costs, respectively, as described in paragraph (h) of this section.

(h) The operating cost-to-charge ratio and, effective with cost reporting periods beginning on or after October 1, 1991, the capital cost-to-charge ratio used to adjust covered charges are computed annually by the intermediary for each hospital based on the latest available settled cost report for that hospital and charge data for the same time period as that covered by the cost report. Statewide cost-to-charge ratios are used in those instances in which a hospital's operating or capital cost-to-charge ratios fall outside reasonable parameters. HCFA
§ 412.86 Payment for extraordinarily high-cost day outliers.

For discharges occurring before October 1, 1997, if a discharge that qualifies for an additional payment under the provisions of §412.82 has charges adjusted to costs that exceed the cost outlier threshold criteria for an extraordinarily high-cost case as set forth in §412.80(a)(1)(ii), the additional payment made for the discharge is the greater of—

(a) The applicable per diem payment computed under §412.82(c) or (d); or

(b) The payment that would be made under §412.84(i) or (j) if the case had not met the day outlier criteria threshold set forth in §412.80(a)(1)(i).


Subpart G—Special Treatment of Certain Facilities Under the Prospective Payment System for Inpatient Operating Costs

§ 412.90 General rules.

(a) Sole community hospitals. HCFA may adjust the prospective payment rates for inpatient operating costs determined under subpart D or E of this part if a hospital, by reason of factors such as isolated location, weather conditions, travel conditions, or absence of other hospitals, is the sole source of inpatient hospital services reasonably available in a geographic area to Medicare beneficiaries. If a hospital meets the criteria for such an exception under §412.92(a), its prospective payment rates for inpatient operating costs are determined under §412.92(d).

(b) Referral center. HCFA may adjust the prospective payment rates for inpatient operating costs determined under subpart D or E of this part if a hospital acts as a referral center for patients transferred from other hospitals. Criteria for identifying such referral centers are set forth in §412.96.

(c) [Reserved]

(d) Kidney acquisition costs incurred by hospitals approved as renal transplantation centers. HCFA pays for kidney acquisition costs incurred by renal transplantation centers on a reasonable cost basis. The criteria for this special payment provision are set forth in §412.100.

(e) Hospitals located in areas that are reclassified from urban to rural. (1) HCFA adjusts the rural Federal payment amounts for inpatient operating costs for hospitals located in geographic areas that are reclassified from urban
§ 412.92 Special treatment: Sole community hospitals.

(a) Criteria for classification as a sole community hospital. HCFA classifies a hospital as a sole community hospital if it is located more than 35 miles from other like hospitals, or it is located in a rural area as defined in §412.103.

(b) Method of determining the prospective payment rate. HCFA determines the prospective payment rate for a sole community hospital, as it does for other hospitals, under §412.92(d).

(c) Additional update for FYs 1998 and 1999. For FYs 1998 and 1999, HCFA makes an upward adjustment to the standardized amounts for certain hospitals that do not receive indirect medical education or disproportionate share payments and are not Medicare-dependent, small rural hospitals. The criteria for identifying these hospitals are set forth in §412.107.

(d) Medicare-dependent, small rural hospitals. For cost reporting periods beginning on or after April 1, 1990 and before October 1, 1994, or beginning on or after October 1, 1997 and before October 1, 2006, HCFA adjusts the prospective payment rates for inpatient operating costs determined under subparts D and E of this part if a hospital is classified as a Medicare-dependent, small rural hospital.

(e) Essential access community hospitals (EACHs). If a hospital was designated as an EACH by HCFA as described in §412.109(a) and is located in a rural area as defined in §412.103, HCFA determines the prospective payment rate for that hospital, as it does for other hospitals, under §412.92(d).

§ 412.92 Special treatment: Sole community hospitals.

(a) Criteria for classification as a sole community hospital. HCFA classifies a hospital as a sole community hospital if it is located more than 35 miles from other like hospitals, or it is located in a rural area as defined in §412.103.

(b) Method of determining the prospective payment rate. HCFA determines the prospective payment rate for a sole community hospital, as it does for other hospitals, under §412.92(d).

(c) Additional update for FYs 1998 and 1999. For FYs 1998 and 1999, HCFA makes an upward adjustment to the standardized amounts for certain hospitals that do not receive indirect medical education or disproportionate share payments and are not Medicare-dependent, small rural hospitals. The criteria for identifying these hospitals are set forth in §412.107.

(d) Medicare-dependent, small rural hospitals. For cost reporting periods beginning on or after April 1, 1990 and before October 1, 1994, or beginning on or after October 1, 1997 and before October 1, 2006, HCFA adjusts the prospective payment rates for inpatient operating costs determined under subparts D and E of this part if a hospital is classified as a Medicare-dependent, small rural hospital.

(e) Essential access community hospitals (EACHs). If a hospital was designated as an EACH by HCFA as described in §412.109(a) and is located in a rural area as defined in §412.103, HCFA determines the prospective payment rate for that hospital, as it does for other hospitals, under §412.92(d).
but because of local topography or periods of prolonged severe weather conditions, the other like hospitals are inaccessible for at least 30 days in each 2 out of 3 years.

(3) Because of distance, posted speed limits, and predictable weather conditions, the travel time between the hospital and the nearest like hospital is at least 45 minutes.

(b) Classification procedures. (1) Request for classification as sole community hospital. (i) The hospital must make its request to its fiscal intermediary.

(ii) If a hospital is seeking sole community hospital classification under paragraph (a)(1)(i) or (a)(1)(ii) of this section, the hospital must include the following information with its request:

(A) The hospital must provide patient origin data (for example, the number of patients from each zip code from which the hospital draws inpatients) for all inpatient discharges to document the boundaries of its service area.

(B) The hospital must provide patient origin data from all other hospitals located within a 35 mile radius of it or, if larger, within its service area, to document that no more than 25 percent of either all of the population or the Medicare beneficiaries residing in the hospital's service area and hospitalized for inpatient care were admitted to other like hospitals for care.

(iii)(A) If the hospital is unable to obtain the information required under paragraph (b)(1)(iii)(A) of this section concerning the residences of Medicare beneficiaries who were inpatients in other hospitals located within a 50 mile radius of the hospital or, if larger, within the hospital's service area, the hospital may request that HCFA provide this information.

(B) If a hospital obtains the information as requested under paragraph (b)(1)(iii)(A) of this section, that information is used by both the intermediary and HCFA in making the determination of the residences of Medicare beneficiaries under paragraphs (b)(1)(iii) and (b)(1)(iv) of this section, regardless of any other information concerning the residences of Medicare beneficiaries submitted by the hospital.

(iv) The intermediary reviews the request and send the request, with its recommendation, to HCFA.

(v) HCFA reviews the request and the intermediary’s recommendation and forward its approval or disapproval to the intermediary.

(2) Effective dates of classification. (i) Sole community hospital status is effective 30 days after the date of HCFA’s written notification of approval.

(ii) When a court order or a determination by the Provider Reimbursement Review Board (PRRB) reverses an HCFA denial of sole community hospital status and no further appeal is made, the sole community hospital status is effective as follows:

(A) If the hospital’s application was submitted prior to October 1, 1983, its status as a sole community hospital is effective at the start of the cost reporting period for which it sought exemption from the cost limits.

(B) If the hospital’s application for sole community hospital status was filed on or after October 1, 1983, the effective date is 30 days after the date of HCFA’s original written notification of denial.

(iii) When a hospital is granted retroactive approval of sole community hospital status by a court order or a PRRB decision and the hospital wishes its sole community hospital status terminated before the date of the court order or PRRB determination, it must submit written notice to the HCFA regional office within 90 days of the court order or PRRB decision. A written request received after the 90-day period is effective no later than 30 days after the request is submitted.

(iv) A hospital classified as a sole community hospital receives a payment adjustment, as described in paragraph (d) of this section, effective with discharges occurring on or after 30 days after the request is submitted.

(3) Duration of classification. An approved classification as a sole community hospital remains in effect without need for reapproval unless there is a change in the circumstances under which the classification was approved.

(4) Cancellation of classification. (i) A hospital may at any time request cancellation of its classification as a sole community hospital and must do so in writing to its fiscal intermediary. The request must be received by the intermediary within 90 days of the date on which the hospital learns that the circumstances under which its classification was approved have changed and the hospital no longer meets the criteria for the classification as a sole community hospital.

(ii) If the intermediary approves the request, it must forward the request, along with its recommendation, to HCFA. HCFA must issue a written determination of approval or disapproval within 30 days of receiving the request.

(iii) If a hospital requests cancellation of its classification as a sole community hospital by sending a written notice to its regional HCFA office within 90 days of the date on which the hospital learns that the circumstances under which its classification was approved have changed and the hospital no longer meets the criteria for the classification as a sole community hospital, the intermediary may cancel such classification.

(iv) A hospital classified as a sole community hospital receives a payment adjustment, as described in paragraph (d) of this section, effective with discharges occurring on or after 30 days after the request is submitted.

(v) When a court order or a determination by the Provider Reimbursement Review Board (PRRB) reverses an HCFA denial of sole community hospital status and no further appeal is made, the sole community hospital status is effective as follows:

(A) If the hospital’s application was submitted prior to October 1, 1983, its status as a sole community hospital is effective at the start of the cost reporting period for which it sought exemption from the cost limits.

(B) If the hospital’s application for sole community hospital status was filed on or after October 1, 1983, the effective date is 30 days after the date of HCFA’s original written notification of denial.

(iii) When a hospital is granted retroactive approval of sole community hospital status by a court order or a PRRB decision and the hospital wishes its sole community hospital status terminated before the date of the court order or PRRB determination, it must submit written notice to the HCFA regional office within 90 days of the court order or PRRB decision. A written request received after the 90-day period is effective no later than 30 days after the request is submitted.

(iv) A hospital classified as a sole community hospital receives a payment adjustment, as described in paragraph (d) of this section, effective with discharges occurring on or after 30 days after the request is submitted.

(3) Duration of classification. An approved classification as a sole community hospital remains in effect without need for reapproval unless there is a change in the circumstances under which the classification was approved.

(4) Cancellation of classification. (i) A hospital may at any time request cancellation of its classification as a sole community hospital and must do so in writing to its fiscal intermediary. The request must be received by the intermediary within 90 days of the date on which the hospital learns that the circumstances under which its classification was approved have changed and the hospital no longer meets the criteria for the classification as a sole community hospital.

(ii) If the intermediary approves the request, it must forward the request, along with its recommendation, to HCFA. HCFA must issue a written determination of approval or disapproval within 30 days of receiving the request.

(iii) If a hospital requests cancellation of its classification as a sole community hospital by sending a written notice to its regional HCFA office within 90 days of the date on which the hospital learns that the circumstances under which its classification was approved have changed and the hospital no longer meets the criteria for the classification as a sole community hospital, the intermediary may cancel such classification.

(iv) A hospital classified as a sole community hospital receives a payment adjustment, as described in paragraph (d) of this section, effective with discharges occurring on or after 30 days after the request is submitted.

(v) When a court order or a determination by the Provider Reimbursement Review Board (PRRB) reverses an HCFA denial of sole community hospital status and no further appeal is made, the sole community hospital status is effective as follows:

(A) If the hospital’s application was submitted prior to October 1, 1983, its status as a sole community hospital is effective at the start of the cost reporting period for which it sought exemption from the cost limits.

(B) If the hospital’s application for sole community hospital status was filed on or after October 1, 1983, the effective date is 30 days after the date of HCFA’s original written notification of denial.

(iii) When a hospital is granted retroactive approval of sole community hospital status by a court order or a PRRB decision and the hospital wishes its sole community hospital status terminated before the date of the court order or PRRB determination, it must submit written notice to the HCFA regional office within 90 days of the court order or PRRB decision. A written request received after the 90-day period is effective no later than 30 days after the request is submitted.

(iv) A hospital classified as a sole community hospital receives a payment adjustment, as described in paragraph (d) of this section, effective with discharges occurring on or after 30 days after the request is submitted.

(3) Duration of classification. An approved classification as a sole community hospital remains in effect without need for reapproval unless there is a change in the circumstances under which the classification was approved.

(4) Cancellation of classification. (i) A hospital may at any time request cancellation of its classification as a sole community hospital and must do so in writing to its fiscal intermediary. The request must be received by the intermediary within 90 days of the date on which the hospital learns that the circumstances under which its classification was approved have changed and the hospital no longer meets the criteria for the classification as a sole community hospital.

(ii) If the intermediary approves the request, it must forward the request, along with its recommendation, to HCFA. HCFA must issue a written determination of approval or disapproval within 30 days of receiving the request.

(iii) If a hospital requests cancellation of its classification as a sole community hospital by sending a written notice to its regional HCFA office within 90 days of the date on which the hospital learns that the circumstances under which its classification was approved have changed and the hospital no longer meets the criteria for the classification as a sole community hospital, the intermediary may cancel such classification.

(iv) A hospital classified as a sole community hospital receives a payment adjustment, as described in paragraph (d) of this section, effective with discharges occurring on or after 30 days after the request is submitted.

(v) When a court order or a determination by the Provider Reimbursement Review Board (PRRB) reverses an HCFA denial of sole community hospital status and no further appeal is made, the sole community hospital status is effective as follows:

(A) If the hospital’s application was submitted prior to October 1, 1983, its status as a sole community hospital is effective at the start of the cost reporting period for which it sought exemption from the cost limits.

(B) If the hospital’s application for sole community hospital status was filed on or after October 1, 1983, the effective date is 30 days after the date of HCFA’s original written notification of denial.

(iii) When a hospital is granted retroactive approval of sole community hospital status by a court order or a PRRB decision and the hospital wishes its sole community hospital status terminated before the date of the court order or PRRB determination, it must submit written notice to the HCFA regional office within 90 days of the court order or PRRB decision. A written request received after the 90-day period is effective no later than 30 days after the request is submitted.

(iv) A hospital classified as a sole community hospital receives a payment adjustment, as described in paragraph (d) of this section, effective with discharges occurring on or after 30 days after the request is submitted.
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Community hospital, and be paid at rates determined under subparts D and E of this part, as appropriate.

(ii) The cancellation becomes effective no later than 30 days after the date the hospital submits its request.

(iii) If a hospital requests that its sole community hospital classification be cancelled, it may not be reclassified as a sole community hospital unless it meets the following conditions:

(A) At least one full year has passed since the effective date of its cancellation.

(B) The hospital meets the qualifying criteria set forth in paragraph (a) of this section in effect at the time it re-applies.

(5) Automatic classification as a sole community hospital.

A hospital that has been granted an exemption from the hospital cost limits before October 1, 1983, or whose request for the exemption was received by the appropriate intermediary before October 1, 1983, and was subsequently approved, is automatically classified as a sole community hospital unless that classification has been cancelled under paragraph (b)(3) of this section, or there is a change in the circumstances under which the classification was approved.

(c) Terminology. As used in this section—

(1) The term miles means the shortest distance in miles measured over improved roads. An improved road for this purpose is any road which is maintained by a local, State, or Federal government entity and which is available for use by the general public.

(2) The term like hospital means a hospital furnishing short-term, acute care. HCFA will not evaluate comparability of specialty services in making determinations on classifications as sole community hospitals.

(3) The term service area means the area from which a hospital draws at least 75 percent of its inpatients during the most recent 12-month cost reporting period ending before it applies for classification as a sole community hospital.

(d) Determining prospective payment rates for inpatient operating costs for sole community hospitals. (1) General rule. For cost reporting periods beginning on or after April 1, 1990, a sole community hospital is paid based on whichever of the following amounts yields the greatest aggregate payment for the cost reporting period:

(i) The Federal payment rate applicable to the hospitals as determined under §412.63.

(ii) The hospital-specific rate as determined under §412.73.

(iii) The hospital-specific rate as determined under §412.75.

(iv) For cost reporting periods beginning on or after October 1, 2000, the hospital-specific rate as determined under §412.77 (calculated under the transition schedule set forth in paragraph (d)(2) of this section), if the sole community hospital was paid for its cost reporting period beginning during 1999 on the basis of the hospital-specific rate specified in paragraph (d)(1)(ii) or (d)(1)(iii) of this section, unless the hospital elects otherwise under §412.77(a)(1).

(2) Transition of FY 1996 hospital-specific rate. The intermediary calculates the hospital-specific rate determined on the basis of the fiscal year 1996 base period rate as follows:

(i) For Federal fiscal year 2001, the hospital-specific rate is the sum of 75 percent of the greater of the hospital-specific rates specified in paragraph (d)(1)(ii) or (d)(1)(iii) of this section, plus 25 percent of the hospital-specific rate specified in paragraph (d)(1)(iv) of this section.

(ii) For Federal fiscal year 2002, the hospital-specific rate is the sum of 50 percent of the greater of the hospital-specific rates specified in paragraph (d)(1)(ii) or (d)(1)(iii) of this section, plus 50 percent of the hospital-specific rate specified in paragraph (d)(1)(iv) of this section.

(iii) For Federal fiscal year 2003, the hospital-specific rate is the sum of 25 percent of the greater of the hospital-specific rates specified in paragraph (d)(1)(ii) or (d)(1)(iii) of this section, plus 75 percent of the hospital-specific rate specified in paragraph (d)(1)(iv) of this section.

(iv) For Federal fiscal year 2004 and any subsequent fiscal years, the hospital-specific rate is 100 percent of the hospital-specific rate specified in paragraph (d)(1)(iv) of this section.
(3) Adjustments to payments. A sole community hospital may receive an adjustment to its payments to take into account a significant decrease in number of discharges or a significant increase in inpatient operating costs, as described in paragraphs (e) and (g) of this section respectively.

(e) Additional payments to sole community hospitals experiencing a significant volume decrease. (1) For cost reporting periods beginning on or after October 1, 1983, the intermediary provides for a payment adjustment for a sole community hospital for any cost reporting period during which the hospital experiences, due to circumstances as described in paragraph (e)(2) of this section a more than five percent decrease in its total discharges of inpatients as compared to its immediately preceding cost reporting period. If either the cost reporting period in question or the immediately preceding cost reporting period is other than a 12-month cost reporting period, the intermediary must convert the discharges to a monthly figure and multiply this figure by 12 to estimate the total number of discharges for a 12-month cost reporting period.

(2) To qualify for a payment adjustment on the basis of a decrease in discharges, a sole community hospital must submit its request no later than 180 days after the date on the intermediary’s Notice of Amount of Program Reimbursement—

(i) Submit to the intermediary documentation demonstrating the size of the decrease in discharges, and the resulting effect on per discharge costs; and

(ii) Show that the decrease is due to circumstances beyond the hospital’s control.

(3) The intermediary determines a lump sum adjustment amount not to exceed the difference between the hospital’s Medicare inpatient operating costs and the hospital’s total DRG revenue for inpatient operating costs based on DRG-adjusted prospective payment rates for inpatient operating costs (including outlier payments for inpatient operating costs determined under subpart F of this part and additional payments made for inpatient operating costs for hospitals that serve a disproportionate share of low-income patients as determined under §412.106 and for indirect medical education costs as determined under §412.105).

(i) In determining the adjustment amount, the intermediary considers—

(A) The individual hospital’s needs and circumstances, including the reasonable cost of maintaining necessary core staff and services in view of minimum staffing requirements imposed by State agencies;

(B) The hospital’s fixed (and semi-fixed) costs, other than those costs paid on a reasonable cost basis under part 413 of this chapter; and

(C) The length of time the hospital has experienced a decrease in utilization.

(ii) The intermediary makes its determination within 180 days from the date it receives the hospital’s request and all other necessary information.

(iii) The intermediary determination is subject to review under subpart R of part 405 of this chapter.

§412.96 Special treatment: Referral centers.

(a) Criteria for classification as a referral center. Basic rule. HCFA classifies a hospital as a referral center only if the hospital is a Medicare participating acute care hospital and meets the applicable criteria of paragraph (b) or (c) of this section.

(b) Criteria for cost reporting periods beginning on or after October 1, 1983. The hospital meets either of the following criteria:

(1) The hospital is located in a rural area (as defined in §412.63(b)) and has the following number of beds, as determined under the provisions of §412.105(b), available for use:

(i) Effective for discharges occurring before April 1, 1988, the hospital has 500 or more beds.

(ii) Effective for discharges occurring on or after April 1, 1988, the hospital...
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has 275 or more beds during its most recently completed cost reporting period unless the hospital submits written documentation with its application that its bed count has changed since the close of its most recently completed cost reporting period for one or more of the following reasons:

(A) Merger of two or more hospitals.

(B) Reopening of acute care beds previously closed for renovation.

(C) Transfer to the prospective payment system of acute care beds previously classified as part of an excluded unit.

(D) Expansion of acute care beds available for use and permanently maintained for lodging inpatients, excluding beds in corridors and other temporary beds.

(2) The hospital shows that—

(i) At least 50 percent of its Medicare patients are referred from other hospitals or from physicians not on the staff of the hospital; and

(ii) At least 60 percent of the hospital’s Medicare patients live more than 25 miles from the hospital, and at least 60 percent of all the services that the hospital furnishes to Medicare beneficiaries are furnished to beneficiaries who live more than 25 miles from the hospital.

(c) Alternative criteria. For cost reporting periods beginning on or after October 1, 1985, a hospital that does not meet the criteria of paragraph (b) of this section is classified as a referral center if it is located in a rural area (as defined in § 412.62(f)) and meets the criteria specified in paragraphs (c)(1) and (c)(2) of this section and at least one of the three criteria specified in paragraphs (c)(3), (c)(4), and (c)(5) of this section.

(1) Case-mix index. HCFA sets forth national and regional case-mix index values in each year’s annual notice of prospective payment rates published under § 412.8(b). The methodology HCFA uses to calculate these criteria is described in paragraph (h) of this section. Except as provided in paragraph (c)(2)(ii) of this section for an osteopathic hospital, for the hospital’s most recently completed cost reporting period, its number of discharges (not including discharges from units excluded from the prospective payment system under subpart B of this part or from newborn units) is at least equal to—

(A) For hospitals applying for rural referral center status for cost reporting periods beginning on or after October 1, 1985 and before October 1, 1986, the number of discharges under either the national or regional criterion; or

(B) For hospitals applying for rural referral center status for cost reporting periods beginning on or after October 1, 1986, 5,000 discharges or, if less, the median number of discharges for urban hospitals located in each region.

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(ii) For cost reporting periods beginning on or after January 1, 1986, an osteopathic hospital, recognized by the American Osteopathic Healthcare Association (or any successor organization), that is located in a rural area must have at least 3,000 discharges during its most recently completed cost reporting period to meet the number of discharges criterion. The 3,000 discharges benchmark is also used in evaluating an osteopathic hospital for purposes of the triennial review.

(3) Medical staff. More than 50 percent of the hospital’s active medical staff are specialists who meet one of the following conditions:

(i) Are certified as specialists by one of the Member Boards of the American Board of Medical Specialties or the Advisory Board of Osteopathic Specialists.

(ii) Have completed the current training requirements for admission to the certification examination of one of the Member Boards of the American Board of Medical Specialties or the Advisory Board of Osteopathic Specialists.

(iii) Have successfully completed a residency program in a medical specialty accredited by the Accreditation Council of Graduate Medical Education or the American Osteopathic Association.

(4) Source of inpatients. At least 60 percent of all its discharges are for inpatients who reside more than 25 miles from the hospital.

(5) Volume of referrals. At least 40 percent of all inpatients treated at the hospital are referred from other hospitals or from physicians not on the hospital’s staff.

(d) Payment to rural referral centers. Effective for discharges occurring on or after April 1, 1988, and before October 1, 1994, a hospital that is located in a rural area and meets the criteria of paragraphs (b)(1), (b)(2) or (c) of this section is paid prospective payments for inpatient operating costs per discharge based on the applicable other urban payment rates as determined in accordance with §412.63, as adjusted by the hospital’s area wage index value.

(e) (f) [Reserved]

(g) Hospital cancellation of referral center status. (1) A hospital may at any time request cancellation of its status as a referral center and be paid prospective payments per discharge based on the applicable rural rate as determined in accordance with §412.63, as adjusted by the hospital’s area wage index value.

(2) The cancellation becomes effective no later than 30 days after the date the hospital submits its request.

(3) If a hospital requests that its referral center status be canceled, it may not be reclassified as a referral center unless it meets the qualifying criteria set forth in paragraph (a) of this section in effect at the time it reapplies.

(h) Methodology for calculating case-mix index criteria. HCFA calculates the national and regional case-mix index standards using the latest available data from hospitals subject to the prospective payment system for the Federal fiscal year.

(1) Updating process. HCFA updates the national and regional case-mix index standards using the latest available data from hospitals subject to the prospective payment system for the Federal fiscal year.

(2) Source of data. In making the calculations described in paragraph (g)(1) of this section, HCFA uses all inpatient hospital bills received for discharges subject to prospective payment during the Federal fiscal year being monitored.

(3) Effective date. HCFA sets forth the national and regional criteria in the annual notice of prospective payment rates published under §412.8(b). These criteria are used to determine if a hospital qualifies for referral center status for cost reporting periods beginning on or after October 1 of the Federal fiscal year to which the notice applies.

(4) Applicability of criteria to HCFA review of referral center status. For purposes of the triennial HCFA review of a referral center’s status as described in paragraph (f) of this section, the referral center’s case-mix index value for a Federal fiscal year is evaluated using the appropriate case-mix value criteria published in the annual notice of prospective payment rates.

(i) Methodology for calculating number of discharges criteria. For purposes of determining compliance with the national or regional number of discharges criterion under paragraph (c)(2) of this
section, HCFA calculates the criteria as follows:

(1) Updating process. HCFA updates the national and regional number of discharges using the latest available data for levels of admissions or discharges or both.

(2) Source of data. In making the calculations described in paragraph (h)(1) of this section, HCFA uses the most recent hospital admissions or discharge data available.

(3) Annual notice. HCFA sets forth the national and regional criteria in the annual notice of prospective payment rates published under §412.8(b). These criteria are compared to an applying hospital’s number of discharges for its most recently completed cost reporting period in determining if the hospital qualifies for referral center status for cost reporting periods beginning on or after October 1 of the Federal fiscal year to which the notice applies.

(4) Applicability of criteria to HCFA review of referral center status. For purposes of the triennial review of a referral center’s status as described in paragraph (f) of this section, the referral center’s number of discharges for its most recently completed cost reporting period is evaluated using the appropriate discharge criteria published in the annual notice of prospective payment rates.

[50 FR 12741, Mar. 29, 1985]

EDITORIAL NOTE: For FEDERAL REGISTER citations affecting §412.102, see the List of Sections Affected in the Finding Aids section of this volume.

§ 412.102 Special treatment: Hospitals located in areas that are reclassified from urban to rural as a result of a geographic redesignation.

Effective on or after October 1, 1983, a hospital reclassified as rural, as defined in §412.62(f), may receive an adjustment to its rural Federal payment amount for operating costs for two successive fiscal years.
§ 412.103 Special treatment: Hospitals located in urban areas and that apply for reclassification as rural.

(a) General criteria. A prospective payment hospital that is located in an urban area (as defined in §412.62(f)(1)(ii)) may be reclassified as a rural hospital if it submits an application in accordance with paragraph (b) of this section and meets any of the following conditions:

(1) The hospital is located in a rural census tract of a Metropolitan Statistical Area (MSA) as determined under the most recent version of the Goldsmith Modification as determined by the Office of Rural Health Policy (ORHP) of the Health Resources and Services Administration which is available via the ORHP website at http://www.nal.usda.gov/orph or from the U.S. Department of Health and Human Services, Health Resources and Services Administration, Office of Rural Health Policy, 5600 Fishers Lane, Rockville, MD 20857.

(2) The hospital is located in an area designated by any law or regulation of the State in which it is located as a rural area, or the hospital is designated as a rural hospital by State law or regulation.

(3) The hospital would qualify as a rural referral center as set forth in §412.96, or as a sole community hospital as set forth in §412.92, if the hospital were located in a rural area.

(b) Application requirements. (1) Written application. A hospital seeking reclassification under this section must submit a complete application in writing to HCFA in accordance with paragraphs (b)(2) and (b)(3) of this section.

(2) Contents of application. An application is complete if it contains an explanation of how the hospital meets the condition that constitutes the basis of the request for reclassification set forth in paragraph (a) of this section, including data and documentation necessary to support the request.

(3) Mailing of application. An application must be mailed to the HCFA Regional Office by the requesting hospital and may not be submitted by facsimile or other electronic means.

(4) Notification by HCFA. Within 5 business days after receiving the hospital’s application, the HCFA Regional Office will send the hospital a letter acknowledging receipt, with a copy to the HCFA Central Office.

(5) Filing date. The filing date of the application is the date HCFA receives the application.

(c) HCFA review. The HCFA Regional Office will review the application and notify the hospital of its approval or disapproval of the request within 60 days of the filing date.

(d) Effective dates of reclassification. (1) Except as specified in paragraph (d)(2) of this section, HCFA will consider a hospital that satisfies any of the criteria set forth in paragraph (a) of this section as being located in the rural area of the State in which the hospital is located as of that filing date.

(2) If a hospital’s complete application is received by HCFA by September 1, 2000, and satisfies any of the criteria set forth in paragraph (a) of this section, HCFA will consider the filing date to be January 1, 2000.
§ 412.105 Special treatment: Hospitals with high percentage of ESRD discharges.

(a) Criteria for classification. Effective with cost reporting periods that begin on or after October 1, 1984, HCFA provides an additional payment to a hospital for inpatient dialysis provided to ESRD beneficiaries if the hospital has established that ESRD beneficiary discharges, excluding discharges classified into DRG No. 302 (Kidney Transplant), DRG No. 316 (Renal Failure) or DRG No. 317 (Admit for Renal Dialysis), constitute ten percent or more of its total Medicare discharges.

(b) Additional payment. A hospital that meets the criteria of paragraph (a) of this section is paid an additional payment for each ESRD beneficiary discharge except those excluded under paragraph (a) of this section.

(1) The payment is based on the estimated weekly cost of dialysis and the average length of stay of ESRD beneficiaries for the hospital.

(2) The estimated weekly cost of dialysis is the average number of dialysis sessions furnished per week during the 12-month period that ended June 30, 1983 multiplied by the average cost of dialysis for the same period.

(3) The average cost of dialysis includes only those costs determined to be directly related to the dialysis service. (These costs include salary, employee health and welfare, drugs, supplies, and laboratory services.)

(4) The average cost of dialysis is reviewed and adjusted, if appropriate, at the time the composite rate reimbursement for outpatient dialysis is reviewed.

(5) The payment to a hospital equals the average length of stay of ESRD beneficiaries in the hospital, expressed as a ratio to one week, times the estimated weekly cost of dialysis multiplied by the number of ESRD beneficiary discharges except for those excluded under paragraph (a) of this section. This payment is made only on the Federal portion of the payment rate.

[50 FR 12741, Mar. 29, 1985, as amended at 57 FR 39824, Sept. 1, 1992]

§ 412.105 Special treatment: Hospitals that incur indirect costs for graduate medical education programs.

HCFA makes an additional payment to hospitals for indirect medical education costs using the following procedures:

(a) Basic data. HCFA determines the following for each hospital:

(1) The hospital's ratio of full-time equivalent residents, except as limited under paragraph (f) of this section, to the number of beds (as determined in paragraph (b) of this section). Except for the special circumstances for affiliated groups and new programs described in paragraphs (f)(1)(vi) and (f)(1)(vii) of this section, for a hospital's cost reporting periods beginning on or after October 1, 1997, this ratio may not exceed the ratio for the hospital's most recent prior cost reporting period.

(2) The hospital's DRG revenue for inpatient operating costs based on DRG-adjusted prospective payment rates for inpatient operating costs, excluding outlier payments for inpatient operating costs determined under subpart F of this part and additional payments made under the provisions of § 412.106.

(b) Determination of number of beds. For purposes of this section, the number of beds in a hospital is determined by counting the number of available bed days during the cost reporting period, not including beds or bassinets in
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the healthy newborn nursery, custodial care beds, or beds in excluded distinct part hospital units, and dividing that number by the number of days in the cost reporting period.

(c) Measurement for teaching activity. The factor representing the effect of teaching activity on inpatient operating costs equals .405 for discharges occurring on or after May 1, 1986.

(d) Determination of education adjustment factor. Each hospital's education adjustment factor is calculated as follows:

(1) Step one. A factor representing the sum of 1.00 plus the hospital's ratio of full-time equivalent residents to beds, as determined under paragraph (a)(1) of this section, is raised to an exponential power equal to the factor set forth in paragraph (c) of this section.

(2) Step two. The factor derived from step one is reduced by 1.00.

(3) Step three. The factor derived from completing steps one and two is multiplied by 'c', and where 'c' is equal to the following:

(i) For discharges occurring on or after October 1, 1988, and before October 1, 1997, 1.89.

(ii) For discharges occurring during fiscal year 1998, 1.72.

(iii) For discharges occurring during fiscal year 1999, 1.6.

(iv) For discharges occurring during fiscal year 2000, 1.47.

(A) Each hospital receives an amount that is equal in the aggregate to the difference between the amount of payments made to the hospital if 'c' equaled 1.6, rather than 1.47.

(B) The payment of this amount will not affect any other payments, determinations, or budget neutrality adjustments.

(v) For discharges occurring during fiscal year 2001, 1.54.

(vi) For discharges occurring on or after October 1, 2001, 1.35.

(e) Determination of payment amount. Each hospital's indirect medical education payment under the prospective payment system for inpatient operating costs is determined by multiplying the total DRG revenue for inpatient operating costs, as determined under paragraph (a)(2) of this section, by the applicable education adjustment factor derived in paragraph (d) of this section.

C. Adding and reserving paragraphs (f)(1)(viii) and (ix).

D. Adding new paragraphs (f)(1)(x), (f)(1)(xi), and (f)(1)(xii).

(f) Determining the total number of full-time equivalent residents for cost reporting periods beginning on or after July 1, 1991. (1) For cost reporting periods beginning on or after July 1, 1991, the count of full-time equivalent residents for the purpose of determining the indirect medical education adjustment is determined as follows:

(i) The resident must be enrolled in an approved teaching program. An approved teaching program is one that meets one of the following requirements:

(A) Is approved by one of the national organizations listed in §415.200(a) of this chapter.

(B) May count towards certification of the participant in a specialty or subspecialty listed in the current edition of either of the following publications:

(1) The Directory of Graduate Medical Education Programs published by the American Medical Association.

(2) The Annual Report and Reference Handbook published by the American Board of Medical Specialties.

(C) Is approved by the Accreditation Council for Graduate Medical Education (ACGME) as a fellowship program in geriatric medicine.

(D) Is a program that would be accredited except for the accrediting agency's reliance upon an accreditation standard that requires an entity to perform an induced abortion or require, provide, or refer for training in the performance of induced abortions, or make arrangements for such training, regardless of whether the standard provides exceptions or exemptions.

(ii) In order to be counted, the resident must be assigned to one of the following areas:

(A) The portion of the hospital subject to the prospective payment system.

(B) The outpatient department of the hospital.

(C) Effective for discharges occurring on or after October 1, 1997, the time spent by a resident in a nonhospital setting in patient care activities under
an approved medical residency training program is counted towards the determination of full-time equivalency if the criteria set forth at §413.86(f)(4) are met.

(iii) Full-time equivalent status is based on the total time necessary to fill a residency slot. No individual may be counted as more than one full-time equivalent. If a resident is assigned to more than one hospital, the resident counts as a partial full-time equivalent based on the proportion of time worked in any of the areas of the hospital listed in paragraph (f)(1)(ii) of this section, to the total time worked by the resident. A part-time resident or one working in an area of the hospital other than those listed under paragraph (f)(1)(ii) of this section (such as a free-standing family practice center or an excluded hospital unit) would be counted as a partial full-time equivalent based on the proportion of time assigned to an area of the hospital listed in paragraph (f)(1)(ii) of this section, compared to the total time necessary to fill a full-time internship or residency slot.

(iv) Effective for discharges occurring on or after October 1, 1997, the total number of FTE residents in the fields of allopathic and osteopathic medicine in either a hospital or a non-hospital setting that meets the criteria listed in paragraph (f)(1)(ii) of this section (such as a family practice center or an excluded hospital unit) would be counted as a partial full-time equivalent based on the proportion of time assigned to an area of the hospital listed in paragraph (f)(1)(ii) of this section, compared to the total time necessary to fill a full-time internship or residency slot.

(v) Effective for discharges occurring on or after April 1, 1997, the total number of FTE residents in the fields of allopathic and osteopathic medicine in either a hospital or a non-hospital setting that meets the criteria listed in paragraph (f)(1)(ii) of this section (such as a free-standing family practice center or an excluded hospital unit) would be counted as a partial full-time equivalent based on the proportion of time assigned to an area of the hospital listed in paragraph (f)(1)(ii) of this section, compared to the total time necessary to fill a full-time internship or residency slot.

(x) Effective for discharges occurring on or after April 1, 2000, an urban hospital that establishes a new residency program (as defined in §413.86(g)(12) of this subchapter), or has an existing residency program, with a rural track (or an integrated rural track) may include in its FTE count residents in those rural tracks in accordance with the provisions of §§413.86(g)(11) of this subchapter.

(xi) Effective for discharges occurring on or after November 29, 1999, a hospital may receive an adjustment to its FTE cap of up to three additional FTEs to the extent that the additional residents would have been counted as
primary care residents for purposes of the hospital’s FTE cap but for the fact that the additional residents were on maternity or disability leave or a similar approved leave of absence, in accordance with the provisions of §413.86(g)(9) of this subchapter.

(xii) For discharges occurring on or after October 1, 1997, a non-Veterans Affairs (VA) hospital may receive a temporary adjustment to its FTE cap to reflect residents who had been previously trained at a VA hospital and were subsequently transferred to the non-VA hospital, if the hospital meets the criteria and other provisions of §413.86(g)(10) of this subchapter.

(2) To include a resident in the full-time equivalent count for a particular cost reporting period, the hospital must furnish the following information. The information must be certified by an official of the hospital and, if different, an official responsible for administering the residency program.

(i) A listing, by specialty, of all residents assigned to the hospital and providing services to the hospital during the cost reporting period.

(ii) The name and social security number of each resident.

(iii) The dates the resident is assigned to the hospital and providing services to the hospital during the cost reporting period.

(iv) The dates the resident is assigned to other hospitals or other freestanding providers and any nonprovider setting during the cost reporting period.

(v) The proportion of the total time necessary to fill a residency slot that the resident is assigned to an area of the hospital listed under paragraph (f)(1)(ii) of this section.

(3) Fiscal intermediaries must verify the correct count of residents.

(g) Indirect medical education payment for managed care enrollees. For portions of cost reporting periods occurring on or after January 1, 1998, a payment is made to a hospital for indirect medical education costs, as determined under paragraph (e) of this section, for discharges associated with individuals who are enrolled under a risk-sharing contract with an eligible organization under section 1876 of the Act or with a Medicare+Choice organization under title XVIII, Part C of the Act during the period, according to the applicable payment percentages described in §§413.86(d)(3)(i) through (d)(3)(v) of this subchapter.


EDITORIAL NOTE: For Federal Register citations affecting §412.105, see the List of Sections Affected in the Finding Aids section of this volume.

§412.106 Special treatment: Hospitals that serve a disproportionate share of low-income patients.

(a) General considerations. (1) The factors considered in determining whether a hospital qualifies for a payment adjustment include the number of beds, the number of patient days, and the hospital’s location.

(i) The number of beds in a hospital is determined in accordance with §412.105(b).

(ii) The number of patient days includes only those days attributable to areas of the hospital that are subject to the prospective payment system and excludes all others.

(iii) The hospital’s location, in an urban or rural area, is determined in accordance with the definitions in §412.62(f).

(2) The payment adjustment is applied to the hospital’s DRG revenue for inpatient operating costs based on DRG-adjusted prospective payment rates for inpatient operating costs, excluding outlier payments for inpatient operating costs under subpart F of this part and additional payments made under the provisions of §412.105.

(b) Determination of a hospital’s disproportionate patient percentage. (1) General rule. A hospital’s disproportionate patient percentage is determined by adding the results of two computations and expressing that sum as a percentage.

(2) First computation: Federal fiscal year. For each month of the Federal fiscal year in which the hospital’s cost reporting period begins, HCFA—

(i) Determines the number of covered patient days that—

(A) Are associated with discharges occurring during each month; and

(B) Are furnished to patients who during that month were entitled to both Medicare Part A and SSI, excluding those patients who received only State supplementation;

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(ii) Adds the results for the whole period; and
(iii) Divides the number determined under paragraph (b)(2)(ii) of this section by the total number of patient days that—
(A) Are associated with discharges that occur during that period; and
(B) Are furnished to patients entitled to Medicare Part A.

(3) First computation: Cost reporting period. If a hospital prefers that HCFA use its cost reporting period instead of the Federal fiscal year, it must furnish to HCFA, through its intermediary, a written request including the hospital’s name, provider number, and cost reporting period end date. This exception will be performed once per hospital per cost reporting period, and the resulting percentage becomes the hospital’s official Medicare Part A/SSI percentage for that period.

(4) Second computation. The fiscal intermediary determines, for the same cost reporting period used for the first computation, the number of the hospital’s patient days of service for which patients were eligible for Medicaid but not entitled to Medicare Part A, and divides that number by the total number of patient days in the same period. For purposes of this second computation, the following requirements apply:
(i) A patient is deemed eligible for Medicaid on a given day if the patient is eligible for medical assistance under an approved State Medicaid plan on such day, regardless of whether particular items or services were covered or paid under the State plan.
(ii) Effective with discharges occurring on or after January 20, 2000, for purposes of counting days under paragraph (b)(4)(i) of this section, hospitals may include all days attributable to populations eligible for Title XIX matching payments through a waiver approved under section 1115 of the Social Security Act.
(iii) The hospital has the burden of furnishing data adequate to prove eligibility for each Medicaid patient day claimed under this paragraph, and of verifying with the State that a patient was eligible for Medicaid during each claimed patient hospital day.

(5) Disproportionate patient percentage. The intermediary adds the results of the first computation made under either paragraph (b)(2) or (b)(3) of this section and the second computation made under paragraph (b)(4) of this section and expresses that sum as a percentage. This is the hospital’s disproportionate patient percentage, and is used in paragraph (c) of this section.

(c) Criteria for classification. A hospital is classified as a “disproportionate share” hospital under any of the following circumstances:
(1) The hospital’s disproportionate patient percentage, as determined under paragraph (b)(5) of this section, is at least equal to one of the following:
(i) 15 percent, if the hospital is located in an urban area and has 100 or more beds, or is located in a rural area and has 500 or more beds.
(ii) 30 percent, if the hospital is located in a rural area and either has more than 100 beds and fewer than 500 beds or is classified as a sole community hospital under §412.92 of this subpart.
(iii) 40 percent, if the hospital is located in an urban area and has fewer than 100 beds.
(iv) 45 percent, if the hospital is located in a rural area and has 100 or fewer beds.
(2) The hospital is located in an urban area, has 100 or more beds, and can demonstrate that, during its cost reporting period, more than 30 percent of its net inpatient care revenues are derived from State and local government payments for care furnished to indigent patients.

(d) Payment adjustment. (1) Method of adjustment. Subject to the reduction factor set forth in paragraph (e) of this section, if a hospital serves a disproportionate number of low-income patients, its DRG revenues for inpatient operating costs are increased by an adjustment factor as specified in paragraph (d)(2) of this section.
(2) Payment adjustment factors. (i) If the hospital meets the criteria of paragraph (c)(1)(i) of this section, the payment adjustment factor is equal to one of the following:
(A) If the hospital’s disproportionate patient percentage is greater than 20.2 percent, the applicable payment adjustment factor is as follows:

(a) Additional payment update. A hospital that meets the criteria set forth in paragraph (b) of this section receives the following increase to its applicable percentage amount set forth in §412.63 (p) and (q):

(1) For FY 1998, 0.5 percent.
(2) For FY 1999, 0.3 percent.

(b) Criteria for classification. A hospital is eligible for the additional payment update set forth in paragraph (a) of this section if it meets all of the following criteria:
§ 412.108 Special treatment: Medicare-dependent, small rural hospitals.

(a) Criteria for classification as a Medicare-dependent, small rural hospital. (1) General considerations. For cost reporting periods beginning on or after April 1, 1990 and ending before October 1, 1994, or beginning on or after October 1, 1997 and ending before October 1, 2006, a hospital is classified as a Medicare-dependent, small rural hospital if it is located in a rural area (as defined in §412.63(b)) and meets all of the following conditions:

(i) The hospital has 100 or fewer beds as defined in §412.105(b) during the cost reporting period.

(ii) The hospital is not also classified as a sole community hospital under §412.92.

(iii) At least 60 percent of the hospital’s inpatient days or discharges were attributable to individuals receiving Medicare part A benefits during the hospital’s cost reporting period as follows, subject to the provisions of paragraph (a)(1)(iv) of this section:

(A) The hospital’s cost reporting period ending on or after September 30, 1987 and before September 30, 1988.

(B) If the hospital does not have a cost reporting period that meets the criterion set forth in paragraph (a)(1)(iii)(A) of this section, the hospital’s cost reporting period beginning on or after October 1, 1986, and before October 1, 1987.

(iv) If the cost reporting period determined under paragraph (a)(1)(iii) of this section is for less than 12 months, the hospital’s most recent 12-month or longer cost reporting period before the short period is used.

(2) Counting days and discharges. In counting inpatient days and discharges for purposes of meeting the criteria in paragraph (a)(1)(iii) of this section, only days and discharges from acute care inpatient hospital stays are counted (including days and discharges from swing beds when used for acute care inpatient hospital services), but not including days and discharges from units excluded from the prospective payment system under §§412.25 through 412.30 or from newborn nursery units. For purposes of this section, a transfer as defined in §412.4(b) is considered to be a discharge.

(b) Classification procedures. The fiscal intermediary determines whether a hospital meets the criteria in paragraph (a) of this section. If a hospital disagrees with an intermediary’s decision, it should notify its intermediary and submit documentable evidence that it meets the criteria.

(c) Payment methodology. A hospital that meets the criteria in paragraph (a) of this section is paid for its inpatient operating costs the sum of paragraphs (c)(1) and (c)(2) of this section.

(1) The Federal payment rate applicable to the hospital as determined under §412.63, subject to the regional floor defined in §412.70(c)(6).

(2) The amount, if any, determined as follows:

(i) For discharges occurring during the first three 12-month cost reporting periods that begin on or after April 1, 1990, 100 percent of the amount that the Federal rate determined under paragraph (c)(1) of this section is exceeded by the higher of the following:

(A) The hospital-specific rate as determined under §412.73.

(B) The hospital-specific rate as determined under §412.75.

§ 412.108 Special treatment: Medicare-dependent, small rural hospitals.

(1) Definition. The hospital is not a Medicare-dependent, small rural hospital as defined in §412.108(a) and does not receive any additional payment under the following provisions:

(i) The indirect medical education adjustment made under §412.105.

(ii) The disproportionate share adjustment made under §412.106.

(2) State criteria. The hospital is located in a State in which the aggregate payment made under §412.112 (a) and (c) for hospitals described in paragraph (b)(1) of this section for their cost reporting periods beginning in FY 1995 is less than the allowable operating costs described in §412.2(c) for those hospitals.

(3) Hospital criteria. The aggregate payment made to the hospital under §412.112 (a) and (c) for the hospital’s cost reporting period beginning in the fiscal year in which the additional payment update described in paragraph (a) of this section is made is less than the allowable operating cost described in §412.2(c) for that hospital.

(ii) For discharges occurring during any subsequent cost reporting period (or portion thereof) and before October 1, 1994, and for discharges occurring or after October 1, 1997 and before October 1, 2006, 50 percent of the amount that the Federal rate determined under paragraph (c)(1) of this section is exceeded by the higher of the following:

(A) The hospital-specific rate as determined under §412.73.

(B) The hospital-specific rate as determined under §412.75.

d) Additional payments to hospitals experiencing a significant volume decrease.

(1) HCFA provides for a payment adjustment for a Medicare-dependent, small rural hospital for any cost reporting period during which the hospital experiences, due to circumstances as described in paragraph (d)(2) of this section, a more than 5 percent decrease in its total inpatient discharges as compared to its immediately preceding cost reporting period. If either the cost reporting period in question or the immediately preceding cost reporting period is other than a 12-month cost reporting period, the intermediary must convert the discharges to a monthly figure and multiply this figure by 12 to estimate the total number of discharges for a 12-month cost reporting period.

(2) To qualify for a payment adjustment on the basis of a decrease in discharges, a Medicare-dependent, small rural hospital must submit its request no later than 180 days after the date on the intermediary’s Notice of Amount of Program Reimbursement and it must—

(i) Submit to the intermediary documentation demonstrating the size of the decrease in discharges and the resulting effect on per discharge costs; and

(ii) Show that the decrease is due to circumstances beyond the hospital’s control.

(3) The intermediary determines a lump sum adjustment amount not to exceed the difference between the hospital’s Medicare inpatient operating costs and the hospital’s total DRG revenue for inpatient operating costs based on DRG-adjusted prospective payment rates for inpatient operating costs determined under subpart F of this part and additional payments made for inpatient operating costs hospitals that serve a disproportionate share of low-income patients as determined under §412.106 and for indirect medical education costs as determined under §412.105.

(i) In determining the adjustment amount, the intermediary considers—

(A) The individual hospital’s needs and circumstances, including the reasonable cost of maintaining necessary core staff and services in view of minimum staffing requirements imposed by State agencies;

(B) The hospital’s fixed (and semifixed) costs, other than those costs paid on a reasonable cost basis under part 413 of this chapter; and

(C) The length of time the hospital has experienced a decrease in utilization.

(ii) The intermediary makes its determination within 180 days from the date it receives the hospital’s request and all other necessary information.

(iii) The intermediary determination is subject to review under subpart R of part 405 of this chapter.

§412.109 Special treatment: Essential access community hospitals (EACHs).

(a) General rule. For payment purposes, HCFA treats as a sole community hospital any hospital that is located in a rural area as described in paragraph (b) of this section and that HCFA designated as an EACH under section 1820(i)(1) of the Act as in effect on September 30, 1997, for as long as the hospital continues to comply with the terms, conditions, and limitations that were applicable at the time HCFA designated the hospital as an EACH. The payment methodology for sole community hospitals is set forth at §412.92(d).

(b) Location in a rural area. For purposes of this section, a hospital is located in a rural area if it—
Health Care Financing Administration, HHS

§ 412.110  Total Medicare payment.

Under the prospective payment systems, Medicare's total payment for inpatient hospital services furnished to a Medicare beneficiary by a hospital will equal the sum of the payments listed in

(1) Is located outside any area that is a Metropolitan Statistical Area as defined by the Office of Management and Budget or that has been recognized as urban under §412.62;

(2) Is not deemed to be located in an urban area under §412.63;

(3) Is not classified as an urban hospital for purposes of the standardized payment amount by HCFA or the Medicare Geographic Classification Review Board; or

(4) Is not located in a rural county that has been redesignated to an adjacent urban area under §412.232.

(c) Adjustment to the hospital-specific rate for rural EACH's experiencing increased costs.

(1) General rule. HCFA increases the applicable hospital-specific rate of an EACH that it treats as a sole community hospital if, during a cost reporting period, the hospital experiences an increase in its Medicare inpatient operating costs per discharge that is directly attributable to activities related to its membership in a rural health network.

(2) Request and documentation. In order for a hospital to qualify for an increase in its hospital-specific rate, it must meet the following criteria:

(i) The hospital must submit its request to its intermediary no later than 180 days after the date on the intermediary's notice of program reimbursement.

(ii) The request must include documentation specifically identifying the increased costs resulting from the hospital's participation in a rural health network and show that the increased costs during the cost reporting period will result in increased costs in subsequent cost reporting periods that are not already accounted for under the prospective payment system.

(iii) The hospital must show that the cost increases are incremental costs that would not have been incurred in the absence of the hospital's membership in a rural health network.

(iv) The hospital must show that the cost increases do not include amounts for start-up and one-time, nonrecurring costs attributable to its membership in a rural health network.

(3) Intermediary recommendation. The intermediary forwards the following material to HCFA within 60 days of receipt from the hospital:

(i) The hospital's documentation and the intermediary's verification of that documentation.

(ii) The intermediary's analysis and recommendation of the request.

(3) The hospital's Medicare cost report for the year in which the increase in costs occurred and the prior year.

(4) HCFA determination. HCFA determines, within 120 days of receiving all necessary information from the intermediary, whether an increase in the hospital-specific rate is warranted and, if it is, the amount of the increase. HCFA grants an adjustment only if a hospital's Medicare inpatient operating costs per discharge exceed the hospital's hospital-specific rate. The adjusted hospital-specific rate cannot exceed the hospital's Medicare inpatient operating costs per discharge for the cost reporting period.

(d) Termination of EACH designation.

If HCFA determines that a hospital no longer complies with the terms, conditions, and limitations that were applicable at the time HCFA designated the hospital as an EACH, HCFA will terminate the EACH designation of the hospital, effective with discharges occurring on or after 30 days after the date of the determination.

(e) Review of HCFA determination. A determination by HCFA that a hospital's EACH designation should be terminated, is subject to review under part 405, subpart R of this chapter, including the time limits for filing requests for hearings as specified in §§405.1811(a) and 405.1841(a)(1) and (b) of this chapter.


Subpart H—Payments to Hospitals Under the Prospective Payment Systems

§ 412.110  Total Medicare payment.

Under the prospective payment systems, Medicare's total payment for inpatient hospital services furnished to a Medicare beneficiary by a hospital will equal the sum of the payments listed in
§ 412.112 Payments determined on a per case basis.

A hospital is paid the following amounts on a per case basis:

(a) The appropriate prospective payment rate for inpatient operating costs for each discharge as determined in accordance with subparts D, E, and G of this part.

(b) Effective for cost reporting periods beginning on or after October 1, 1991, the appropriate prospective payment rate for capital-related costs for each discharge as determined in accordance with subpart M of this part.

(c) The appropriate outlier payment amounts determined under subpart F of this part.

§ 412.113 Other payments.

(a) Capital-related costs.

(1) Payment.

Subject to the reductions described in paragraph (a)(2) of this section, payment for capital-related costs (as described in § 413.130 of this chapter) for cost reporting periods beginning before October 1, 1991 is determined on a reasonable cost basis.

(2) Reduction to capital-related payments.

(i) Except for sole community hospitals as defined in § 412.92, the amount of capital-related payments for cost reporting periods beginning before October 1, 1991 (including a return on equity capital as provided under § 413.157 of this chapter) is reduced by—

(A) Three and one-half percent for payments attributable to portions of cost reporting periods occurring during Federal FY 1987;

(B) Seven percent for payments attributable to portions of cost reporting periods occurring during Federal FY 1988 and before January 1, 1989;

(C) Twelve percent for payments attributable to portions of cost reporting periods or discharges (as the case may be) occurring during fiscal year 1988 and before January 1, 1989;

(D) Fifteen percent for payments attributable to portions of cost reporting periods or discharges (as the case may be) occurring during fiscal year 1989 and beginning on or after January 1, 1990 and ending on or before September 30, 1991; and

(E) Ten percent for payments attributable to portions of cost reporting periods occurring on or after October 1, 1991 and before the beginning of the hospital’s first cost-reporting period beginning on or after October 1, 1991.

(ii) If a hospital’s cost reporting period encompasses more than one Federal fiscal year, the reductions to capital-related payments are determined on a prorated monthly basis.

(3) For cost-reporting periods beginning on or after October 1, 1991, a hospital with a hospital-specific rate above the Federal capital rate is paid a hold-harmless payment for old capital determined in accordance with subpart M of this part.

(b) Direct medical education costs.

(1) Payment for the direct medical education costs of interns and residents in approved programs for cost reporting periods beginning prior to July 1, 1985, and for approved education activities of nurses and paramedical health professionals is made as described in § 413.85 of this chapter.

(2) For cost reporting periods beginning on or after July 1, 1985, payment for the direct medical education costs of interns and residents in approved programs is made as described in § 413.86 of this chapter.

(3) Except as provided in § 413.86(c) of this chapter, for cost reporting periods during the prospective payment transition period, the costs of medical education must be determined in a manner that is consistent with the treatment of these costs for purposes of determining the hospital-specific portion of the payment rate as provided in subpart E of this part.

(c) Anesthesia services furnished by hospital employed nonphysician anesthetists or obtained under arrangements.

(1) For cost reporting periods beginning on or after October 1, 1984 through any part of a cost reporting period occurring before January 1, 1989, payment is determined on a reasonable cost basis for anesthesia services provided in the hospital by qualified nonphysician anesthetists (certified registered nurse
anesthetists and anesthesiologist’s assistants) employed by the hospital or obtained under arrangements.

(2)(i) For cost reporting periods, or any part of a cost reporting period, beginning on or after January 1, 1989, through any part of a cost reporting period occurring before January 1, 1990, payment is determined on a reasonable cost basis for anesthesia services provided in a hospital by qualified nonphysician anesthetists employed by the hospital or obtained under arrangement, if the hospital demonstrates to its intermediary prior to April 1, 1989 that it meets the following criteria:

(A) The hospital is located in a rural area as defined in §412.62(f) and is not deemed to be located in an urban area under the provisions of §412.64(b)(3).

(B) The hospital must have employed or contracted with a qualified nonphysician anesthetist, as defined in §410.66 of this chapter, as of January 1, 1988 to perform anesthesia services in that hospital. The hospital may employ or contract with more than one anesthetist; however, the total number of hours of service furnished by the anesthetists may not exceed 2,080 hours per year.

(C) The hospital must provide data for its entire patient population to demonstrate that, during calendar year 1987, its volume of surgical procedures (inpatient and outpatient) requiring anesthesia services did not exceed 250 procedures. For purposes of this section, a surgical procedure requiring anesthesia services means a surgical procedure in which the anesthesia is administered and monitored by a qualified nonphysician anesthetist, a physician other than the primary surgeon, or an intern or resident.

(D) Each qualified nonphysician anesthetist employed by or under contract with the hospital has agreed in writing not to bill on a reasonable charge basis for his or her patient care in that hospital.

(ii) To maintain its eligibility for reasonable cost payment under paragraph (c)(2)(i) of this section in calendar years after 1989 and the year immediately preceding its election of reasonable cost payment, its volume of surgical procedures (inpatient and outpatient) requiring anesthesia services did not exceed 500 procedures.

(iii) A hospital that did not qualify for reasonable cost payment for non-physician anesthetist services furnished in calendar year 1989 can qualify for reasonable cost payment in subsequent calendar years, if it meets the criteria in §412.113(c)(2)(i) (A), (B) and (D) above, and demonstrates to its intermediary prior to the start of the calendar year that it met these criteria. The hospital must provide data for its entire patient population to demonstrate that, during calendar year 1987 and the year immediately preceding its election of reasonable cost payment, its volume of surgical procedures did not exceed 500 procedures.

(iv) For administrative purposes, the volume of surgical procedures for the immediately preceding year is the sum of the surgical procedures for the nine month period ending September 30, annualized for the twelve month period.

(d) Organ acquisition. Payment for organ acquisition costs incurred by hospitals with approved transplantation centers is made on a reasonable cost basis. The term “Organs” is defined in §486.302 of this chapter.

§412.115 Additional payments.

(a) Bad debts. An additional payment is made to each hospital in accordance with §413.80 of this chapter for bad debts attributable to deductible and coinurance amounts related to covered services received by beneficiaries.

(b) Administration of blood clotting factor. For discharges occurring on or after June 19, 1990, and before October 1, 1994, and for discharges occurring on or after October 1, 1997, an additional payment is made to a hospital for each unit of blood clotting factor furnished to a Medicare inpatient who is a hemophiliac.

(c) PRO photocopy and mailing costs. An additional payment is made to a
hospital in accordance with §466.78 of this chapter for the costs of photocopying and mailing medical records requested by a PRO.

§ 412.116 Method of payment.

(a) General rule. Unless the provisions of paragraphs (b) and (c) of this section apply, hospitals are paid for hospital inpatient operating costs and capital-related costs for each discharge based on the submission of a discharge bill. Payments for inpatient hospital services furnished by an excluded psychiatric or rehabilitation unit of a hospital are made as described in §413.64 (a), (c), (d), and (e) of this chapter.

(b) Periodic interim payments—(1) Criteria for receiving periodic interim payments. Effective with claims received on or after July 1, 1987, a hospital that meets the criteria in §413.64(h) of this chapter may request in writing to receive periodic interim payments as described in this paragraph. A hospital that is receiving periodic interim payments also receives payment on this basis for inpatient hospital services furnished by its excluded psychiatric or rehabilitation unit.

(i) Failure of intermediary to make prompt payment. Beginning with claims received in April 1987, the hospital's fiscal intermediary does not meet the requirements of section 1816(c)(2) of the Act, which provides for prompt payment of claims under Medicare Part A, for three consecutive calendar months. The hospital may continue to receive periodic interim payments until the intermediary meets the requirements of section 1816(c)(2) of the Act for three consecutive calendar months beginning with April 1987.

(ii) Hospitals that serve a disproportionate share of low-income patients. The hospital is receiving periodic interim payments as of June 30, 1987 and has a disproportionate share payment adjustment factor of at least 5.1 percent as determined under §412.106(c) for purposes of establishing the average standardized amounts for discharges occurring on or after October 1, 1986 and before October 1, 1987. The hospital's request must be made by a date prior to July 1, 1987, specified by the intermediary.

(iii) Small rural hospitals. The hospital is receiving periodic interim payments as of June 30, 1987 and has a disproportionate share payment adjustment factor of at least 5.1 percent as determined under §412.106(c) for purposes of establishing the average standardized amounts for discharges occurring on or after October 1, 1986 and before October 1, 1987. The hospital may continue to receive periodic interim payments until the hospital's intermediary meets the requirements of section 1816(c)(2) of the Act for three consecutive calendar months beginning with April 1987.

(b)(i) Periodic interim payments. Effective with claims received on or after July 1, 1987, a hospital that meets the criteria in §413.64(h) of this chapter may request in writing to receive periodic interim payments as described in this paragraph. A hospital that is receiving periodic interim payments also receives payment on this basis for inpatient hospital services furnished by its excluded psychiatric or rehabilitation unit.

(ii) Frequency of payment. The intermediary estimates a hospital’s prospective payments as described in paragraph (b)(3) of this section and makes biweekly payments equal to 1/26 of the total estimated amount of payment for the year. Each payment is made two weeks after the end of a biweekly period of service, as described in §413.64(h)(5) of this chapter. These payments are subject to final settlement.

(iii) Amount of payment. (i) The biweekly interim payment amount is based on the total estimated Medicare discharges for the reporting period multiplied by the hospital’s estimated average prospective payment amount as described in paragraph (b)(3)(ii) of this paragraph. These interim payments are reviewed at least twice during the reporting period and adjusted if necessary. Fewer reviews may be necessary if a hospital receives interim payments for less than a full reporting period.

(ii) For purposes of determining periodic interim payments under this paragraph, a hospital’s estimated average prospective payment amount is computed as follows:

(A) If a hospital has no payment experience under the prospective payment system for operating costs, the intermediary computes the hospital’s
estimated average prospective payment amount for operating costs by multiplying its payment rates as determined under §412.70(c), but without adjustment by a DRG weighting factor, by the hospital's case-mix index, and subtracting from this amount estimated deductibles and coinsurance.

(B) Effective for cost-reporting periods beginning on or after October 1, 1991, the intermediary computes a hospital's estimated average prospective payment amount for capital-related costs by multiplying its prospective payment rate as determined under §412.340 or §412.344(a), as applicable, and under §412.308 for cost reporting periods beginning on or after October 1, 2001 but without adjustment by a DRG weighting factor, by the hospital's case-mix index. The intermediary may take into account estimated additional payments per discharge under §412.348. If the hospital is paid under §412.344(a)(1), the intermediary includes an estimated payment for old capital costs per discharge.

(C) If a hospital has payment experience under the prospective payment system for operating costs, and, for cost reporting periods beginning on or after October 1, 1991, for inpatient capital-related costs, the intermediary computes a hospital's estimated average prospective payment amount for operating costs and capital-related costs based on that payment experience, adjusted for projected changes, and subtracts from this amount estimated deductibles and coinsurance.

(4) Termination of periodic interim payments—(i) Request by the hospital. A hospital receiving periodic interim payments may convert to payments on a per discharge basis at any time.

(ii) Removal by the intermediary. An intermediary terminates periodic interim payments if—

(A) A hospital no longer meets the requirements of §413.64(h);

(B) A hospital is receiving payment under the criterion in paragraph (b)(1)(i) of this section and the intermediary meets the prompt payment requirements of section 1816(c)(2) of the Act for three consecutive calendar months;

(C) A hospital that is receiving payment under the criterion set forth in paragraph (b)(1)(iii) of this section no longer meets the criterion.

(iii) Limitation on reelection. If a hospital that is receiving periodic interim payments under the criterion set forth in paragraph (b)(1)(ii) or (b)(1)(iii) of this section is removed from that method of payment at its own request, it may reelect to receive periodic interim payments only under the criterion set forth in paragraph (b)(1)(i) of this section. However, if the hospital is removed from that method of payment by its intermediary because it no longer meets the requirements of §413.64(h) of this chapter, that hospital may subsequently reelect to receive periodic interim payments if it qualifies under the provisions of paragraph (b)(1)(iii) or (b)(1)(iii) of this section, subject to the requirements in §413.64(h) of this chapter.

(c) Special interim payments for certain costs. For capital-related costs for cost-reporting periods beginning before October 1, 1991 and the direct costs of medical education, which are not included in prospective payments but are reimbursed as specified in §§413.130 and 413.85 of this chapter, respectively, interim payments are made subject to final cost settlement. Interim payments for capital-related items for cost-reporting periods beginning before October 1, 1991 and the estimated cost of approved medical education programs (applicable to inpatient costs payable under Medicare Part A and for kidney acquisition costs in hospitals approved as renal transplantation centers) are determined by estimating the reimbursable amount for the year based on the previous year's experience and on substantiated information for the current year and divided into 26 equal biweekly payments. Each payment is made two weeks after the end of a biweekly period of services, as described in §413.64(h)(5) of this chapter. The interim payments are reviewed by the intermediary at least twice during the reporting period and adjusted if necessary.

(d) Special interim payment for unusually long lengths of stay—(1) First interim payment. A hospital that is not receiving periodic interim payments under
paragraph (b) of this section may request an interim payment after a Medicare beneficiary has been in the hospital at least 60 days. Payment for the interim bill is determined as if the bill were a final discharge bill and includes any outlier payment determined as of the last day for which services have been billed.

(2) Additional interim payments. A hospital may request additional interim payments at intervals of at least 60 days after the date of the first interim bill submitted under paragraph (d)(1) of this section. Payment for these additional interim bills, as well as the final bill, is determined as if the bill were the final bill with appropriate adjustments made to the payment amount to reflect any previous interim payment made under the provisions of this paragraph (d).

(e) Outlier payments. Payments for outlier cases (described in subpart F of this part) are not made on an interim basis. The outlier payments are made based on submitted bills and represent final payment.

(f) Accelerated payments—(1) General rule. Upon request, an accelerated payment may be made to a hospital that is not receiving periodic interim payments under paragraph (b) of this section if the hospital is experiencing financial difficulties because of the following:

(i) There is a delay by the intermediary in making payment to the hospital.

(ii) Due to an exceptional situation, there is a temporary delay in the hospital’s preparation and submittal of bills to the intermediary beyond its normal billing cycle.

(2) Approval of payment. A hospital’s request for an accelerated payment must be approved by the intermediary and HCFA.

(3) Amount of payment. The amount of the accelerated payment is computed as a percentage of the net payment for unbilled or unpaid covered services.

(4) Recovery of payment. Recovery of the accelerated payment is made by recoupment as hospital bills are processed or by direct payment by the hospital.


§ 412.120 Reductions to total payments.

(a) Deductible and coinsurance. Subject to paragraph (a)(2) of this section, the total Medicare payments otherwise payable to a hospital are reduced by the applicable deductible and coinsurance amounts related to inpatient hospital services as determined in accordance with §§ 409.82, 409.83, and 409.87 of this chapter.

(b) Payment by workers’ compensation, automobile medical, no-fault or liability insurance or an employer group health plan primary to Medicare. If workers’ compensation, automobile medical, no-fault, or liability insurance or an employer group health plan which is primary to Medicare pays in full or in part, the Medicare payment is determined in accordance with the following guidelines:

(1) If workers’ compensation pays, in accordance with the applicable provisions of §§ 405.316 through 405.321 of this chapter.

(2) If automobile medical, no-fault, or liability insurance pays, in accordance with the applicable provisions of §§ 405.322 through 405.325 of this chapter.

(3) If an employer group health plan which is primary to Medicare pays for services to ESRD beneficiaries, in accordance with the applicable provisions of §§ 405.326 through 405.329 of this chapter.

(4) If an employer group health plan which is primary to Medicare pays for services to employees age 65-69 and their spouses age 65-69, in accordance with the applicable provisions of §§ 405.340 through 405.344 of this chapter.

§ 412.125 Effect of change of ownership on payments under the prospective payment systems.

When a hospital’s ownership changes, as described in §489.18 of this chapter, the following rules apply:

(a) Payment for the operating and capital-related costs of inpatient hospital services for each patient, including outlier payments, as provided in §412.112, and payments for hemophilia clotting factor costs under §412.115(b), are made to the entity that is the legal owner on the date of discharge. Payments are not prorated between the buyer and seller.

(1) The owner on the date of discharge is entitled to submit a bill for all inpatient hospital services furnished to a beneficiary regardless of when the beneficiary’s coverage began or ended during a stay, or of how long the stay lasted.

(2) Each bill submitted must include all information necessary for the intermediary to compute the payment amount, whether or not some of that information is attributable to a period during which a different party legally owned the hospital.

(b) Other payments under §412.113 and payments for bad debts as described in §412.115(a), are made to each owner or operator of the hospital (buyer and seller) in accordance with the principles of reasonable cost reimbursement.


§ 412.130 Retroactive adjustments for incorrectly excluded hospitals and units.

(a) Hospitals for which adjustment is made. The intermediary makes the payment adjustment described in paragraph (b) of this section for the following hospitals:

(1) A hospital that was excluded from the prospective payment system as a new hospital for a cost reporting period beginning on or after October 1, 1991 based on a certification under §412.23(b)(2) regarding the inpatient population the hospital planned to treat during that period, if the inpatient population actually treated in the hospital during that cost reporting period did not meet the requirements of §412.23(b)(2).

(2) A hospital that had a unit excluded from the prospective payment system as a new rehabilitation unit for a cost reporting period beginning on or after October 1, 1991 based on a certification under §412.30(a) regarding the inpatient population the hospital planned to treat in that unit during that period, if the inpatient population actually treated in the unit during that cost reporting period did not meet the requirements of §412.23(b)(2).

(3) A hospital that added new beds to its existing rehabilitation unit for a cost reporting period beginning on or after October 1, 1991 based on a certification under §412.30(c) regarding the inpatient population the hospital planned to treat in those new beds during that cost reporting period, if the inpatient population actually treated in those new beds during that cost reporting period did not meet the requirements of §412.23(b)(2).

(b) Adjustment of payment. The intermediary adjusts the payment to the hospitals described in paragraph (a) of this section as follows:

(1) The intermediary calculates the difference between the amounts actually paid during the cost reporting period for which the hospital, unit, or beds were first excluded as a new hospital, new unit, or newly added beds, and the amount that would have been paid under the prospective payment systems for services furnished during that period.

(2) The intermediary makes a retroactive adjustment for the difference between the amount paid to the hospital based on the exclusion and the amount that would have been paid under the prospective payment systems.


Subparts I–J—[Reserved]
§ 412.200 General provisions.

Beginning with discharges occurring on or after October 1, 1987, hospitals located in Puerto Rico are subject to the rules governing the prospective payment system for inpatient operating costs. Except as provided in this subpart, the provisions of subparts A, B, C, F, G, and H of this part apply to hospitals located in Puerto Rico. Except for § 412.60, which deals with DRG classification and weighting factors, the provisions of subparts D and E, which describe the methodology used to determine prospective payment rates for inpatient operating costs for hospitals, do not apply to hospitals located in Puerto Rico. Instead, the methodology for determining prospective payment rates for inpatient operating costs for these hospitals is set forth in §§ 412.204 through 412.212.

[57 FR 39825, Sept. 1, 1992]

§ 412.204 Payment to hospitals located in Puerto Rico.

(a) FY 1988 through FY 1997. For discharges occurring on or after October 1, 1997, payments for inpatient operating costs to hospitals located in Puerto Rico that are paid under the prospective payment system are equal to the sum of—

(1) 75 percent of the Puerto Rico prospective payment rate for inpatient operating costs, as determined under § 412.208 or § 412.210; and

(2) 25 percent of a national prospective payment rate for inpatient operating costs, as determined under § 412.212.


(a) General rule. HCFA determines the Puerto Rico adjusted DRG prospective payment rate for inpatient operating costs for each inpatient hospital discharge occurring in fiscal year 1988 for a prospective payment hospital. These rates are determined as described in paragraphs (b) through (i) of this section.

(b) Determining target amounts. For each hospital subject to the prospective payment system for inpatient operating costs, HCFA determines the Medicare target amount, as described in § 413.40(c) of this chapter, for the hospital's cost reporting period beginning in fiscal year 1987. Revisions in the target amounts made subsequent to establishment of the standardized amounts under paragraph (d) of this section do not affect the standardized amounts.

(c) Updating the target amounts for fiscal year 1988. HCFA updates each target amount determined under paragraph (b) of this section for fiscal year 1988 by prorating the applicable percentage increase (as defined in § 412.63(f) of this chapter) for fiscal year 1988 to the midpoint of fiscal year 1988 (April 1, 1988).

(d) Standardizing amounts. HCFA standardizes the amount updated under paragraph (c) of this section for each hospital by—

(1) Adjusting for variations in case mix among hospitals;

(2) Excluding an estimate of indirect medical education costs;

(3) Adjusting for area variations in hospital wage levels; and

(4) Excluding an estimate of the payments for hospitals that serve a disproportionate share of low-income patients.

(e) Computing urban and rural averages. HCFA computes separate discharge-weighted averages of the standardized amounts determined under
Health Care Financing Administration, HHS § 412.210

paragraph (d) of this section for urban and rural hospitals in Puerto Rico.

(f) Geographic classification. (1) For purposes of this paragraph (e) of this section, the following definitions apply:

(i) The term urban area means a Metropolitan Statistical Area (MSA), as defined by the Executive Office of Management and Budget.

(ii) The term large urban area means an MSA with a population of more than 1,000,000.

(iii) The term rural area means any area outside an urban area.

(2) A hospital classified as rural is deemed to be urban and receives the urban Puerto Rico payment amount if the county in which it is located meets the following criteria:

(i) At least 95 percent of the perimeter of the rural county is contiguous with urban counties.

(ii) The county was reclassified from an urban area to a rural area after April 20, 1983, as described in § 412.62(f)(1)(iv).

(iii) At least 15 percent of employed workers in the county commute to the central county of one of the adjacent MSAs.

(g) Reducing for value of outlier payments. HCFA reduces each of the average standardized amounts determined under paragraphs (c) through (g) of this section by a proportion equal to the proportion (estimated by HCFA) of the total amount of payments based on DRG prospective payment rates that are additional payments to hospitals located in Puerto Rico for outlier cases under subpart F of this part.

(h) Computing Puerto Rico rates established under the prospective payment system for inpatient operating costs for urban and rural hospitals. For each discharge classified within a DRG, HCFA establishes a Puerto Rico prospective payment rate, as follows:

(1) For hospitals located in an urban area, the rate equals the product of—

(i) The average standardized amount (computed under paragraphs (c) through (g) of this section) for hospitals located in a rural area; and

(ii) The weighting factor determined under § 412.60(b) for that DRG.

(2) For hospitals located in a rural area, the rate equals the product of—

(i) The average standardized amount (computed under paragraphs (c) through (g) of this section) for hospitals located in a rural area; and

(ii) The weighting factor determined under § 412.60(b) for that DRG.

(l) Adjusting for different area wage levels. HCFA adjusts the proportion (as estimated by HCFA from time to time) of Puerto Rico rates computed under paragraph (h) of this section that are attributable to wages and labor-related costs, for area differences in hospital wage levels, by a factor (established by HCFA) reflecting the relative hospital wage level in the geographic area (that is, urban or rural area as determined under the provisions of paragraph (f) of this section) of the hospital compared to the national average hospital wage level.


(a) General rule. (1) HCFA determines the Puerto Rico adjusted prospective payment rate for inpatient operating costs for each inpatient hospital discharge occurring in a Federal fiscal year after fiscal year 1988 that involves inpatient hospital services of a hospital in Puerto Rico subject to the prospective payment system for which payment may be made under Medicare Part A.

(2) The rate is determined for hospitals located in large urban, other urban, or rural areas within Puerto Rico, as described in paragraphs (b) through (e) of this section.

(b) Geographic classifications. (1) For purposes of this section, the definitions set forth in § 412.208(f)(1) apply.

(2) For discharges occurring on or after October 1, 1988, a hospital located in a rural county adjacent to one or more urban areas is deemed to be located in an urban area and receives the Federal payment amount for the urban area to which the greatest number of workers in the county commute if the rural county would otherwise be considered part of an urban area, under the standards for designating MSAs if...
§ 412.212 National rate.

(a) General rule. For purposes of payment to hospitals located in Puerto Rico, the national prospective payment rate for inpatient operating costs is determined as described in paragraphs (b) through (d) of this section.

(b) Computing a national average standardized amount. HCFA computes a discharge-weighted average of the—

(1) National urban adjusted standardized amount determined under §412.63(j)(1)(i); and

(2) National rural adjusted average standardized amount determined under §412.63(j)(2)(i).

(c) Computing a national rate. For each discharge classified within a DRG, the national rate equals the product of—

(1) The national average standardized amount computed under paragraph (b) of this section; and

(2) The weighting factor (determined under §412.60(b)) for that DRG.

(d) Adjusting for different area wage levels. HCFA adjusts the proportion (as estimated by HCFA from time to time) of the national rate computed under paragraph (c) of this section that is attributable to wages and labor-related costs for area differences in hospital wage levels by a factor (established by HCFA) reflecting the relative hospital wage level in the geographic area (that is, urban or rural area as determined under the provisions of paragraph (b) of this section) of the hospital compared to the Puerto Rico average hospital wage level.


§ 412.212 National rate.

(a) General rule. For purposes of payment to hospitals located in Puerto Rico, the national prospective payment rate for inpatient operating costs is determined as described in paragraphs (b) through (d) of this section.

(b) Computing a national average standardized amount. HCFA computes a discharge-weighted average of the—

(1) National urban adjusted standardized amount determined under §412.63(j)(1)(i); and

(2) National rural adjusted average standardized amount determined under §412.63(j)(2)(i).

(c) Computing a national rate. For each discharge classified within a DRG, the national rate equals the product of—

(1) The national average standardized amount computed under paragraph (b) of this section; and

(2) The weighting factor (determined under §412.60(b)) for that DRG.

(d) Adjusting for different area wage levels. HCFA adjusts the proportion (as estimated by HCFA from time to time) of the national rate computed under paragraph (c) of this section that is attributable to wages and labor-related costs for area differences in hospital wage levels by a factor (established by HCFA) reflecting the relative hospital wage level in the geographic area (that is, urban or rural area as determined under the provisions of paragraph (b) of this section) of the hospital compared to the Puerto Rico average hospital wage level.


§ 412.212 National rate.

(a) General rule. For purposes of payment to hospitals located in Puerto Rico, the national prospective payment rate for inpatient operating costs is determined as described in paragraphs (b) through (d) of this section.

(b) Computing a national average standardized amount. HCFA computes a discharge-weighted average of the—

(1) National urban adjusted standardized amount determined under §412.63(j)(1)(i); and

(2) National rural adjusted average standardized amount determined under §412.63(j)(2)(i).

(c) Computing a national rate. For each discharge classified within a DRG, the national rate equals the product of—

(1) The national average standardized amount computed under paragraph (b) of this section; and

(2) The weighting factor (determined under §412.60(b)) for that DRG.

(d) Adjusting for different area wage levels. HCFA adjusts the proportion (as estimated by HCFA from time to time) of the national rate computed under paragraph (c) of this section that is attributable to wages and labor-related costs for area differences in hospital wage levels by a factor (established by HCFA) reflecting the relative hospital wage level in the geographic area (that is, urban or rural area as determined under the provisions of paragraph (b) of this section) of the hospital compared to the Puerto Rico average hospital wage level.

§ 412.230 Criteria for an individual hospital seeking redesignation to another rural area or an urban area.

(a) General. (1) Purpose. Except as provided in paragraph (a)(5) of this section, an individual hospital may be redesignated from a rural area to an urban area, from a rural area to another rural area, or from an urban area to another urban area for the purposes of using the other area’s standardized amount for inpatient operating costs, wage index value, or both.

(2) Proximity. Except as provided in paragraph (a)(3) of this section, to be redesignated to another rural area or an urban area, a hospital must demonstrate a close proximity to the area to which it seeks redesignation by meeting the criteria in paragraph (b) of this section, and submitting data requested under paragraph (c) of this section.

(3) Special rules for sole community hospitals and rural referral centers. To be redesignated under the special rules in this paragraph, a hospital must be a sole community hospital or a rural referral center as of the date of the MGCRB’s review.

(i) A hospital that is a rural referral center, a sole community hospital, or both does not have to demonstrate a close proximity to the area to which it seeks redesignation.

(ii) If a hospital that is a rural referral center, a sole community hospital, or both qualifies for urban redesignation, it is redesignated to the urban area that is closest to the hospital. If the hospital is closer to another rural area than to any urban area, it may seek redesignation to either the closest rural or the closest urban area.

(iii) If a sole community hospital or rural referral center loses its special status as a result of redesignation, the hospital is considered to retain its special status for the purpose of applicability of the special rules in paragraph (a)(3) of this section.

(iv) A hospital that is redesignated under paragraph (a)(3) of this section may not be redesignated in the same fiscal year under paragraph (a)(2) of this section.

(4) Application of criteria. In applying the numeric criteria contained in §§412.230(b)(1) and (2), (d)(2), (e)(1)(iii), and (e)(1)(iv) (A) and (B), rounding of numbers to meet the mileage or qualifying percentage standard is not permitted.

(5) Limitations on redesignation. The following limitations apply to redesignation:

(i) An individual hospital may not be redesignated to another area for purposes of the wage index if the pre-reclassified average hourly wage for that area is lower than the pre-reclassified average hourly wage for the area in which the hospital is located.

(ii) For redesignations effective in fiscal years 1997 and 1998 and 2002 and thereafter, a hospital may not be redesignated for purposes of the standardized amount if the area to which the hospital seeks redesignation does not have a higher standardized amount than the standardized amount the hospital currently receives.
(iii) A hospital may not be redesignated to more than one area.

(iv) An urban hospital that has been granted redesignation as rural under §412.103 cannot receive an additional reclassification by the MGCRB based on this acquired rural status as long as such redesignation is in effect.

(b) Proximity criteria. A hospital demonstrates a close proximity with the area to which it seeks redesignation if one of the following conditions applies:

(1) The distance from the hospital to the area is no more than 15 miles for an urban hospital and no more than 35 miles for a rural hospital.

(2) At least 50 percent of the hospital’s employees reside in the area.

(c) Appropriate proximity data. For redesignation to an area, the hospital must submit appropriate data relating to its proximity to that area.

(1) To demonstrate proximity to the area, the hospital must submit evidence of the shortest route over improved roads to the area and the distance of that route.

(2) For employee address data, the hospital must submit current payroll records that include information that establishes the home addresses by zip code of its employees.

(d) Use of an area’s standardized amount for inpatient operating costs. (i) Criteria. To receive an area’s standardized amount for inpatient operating costs, a hospital must demonstrate that its incurred costs are more comparable to the amount it would be paid if it were reclassified than the amount it would be paid under its current classification, and that it has the necessary geographic relationship (as specified in paragraphs (a) and (b) of this section) with the area to which it seeks redesignation.

(2) Demonstrating comparable costs. A hospital demonstrates that its costs are more comparable to the amount it would be paid if it were reclassified if the hospital’s case mix adjusted cost per discharge is at least equal to its current rate plus 75 percent of the difference between that rate and the rate it would receive if it were reclassified.

(3) Appropriate cost data. For a standardized amount for inpatient operating costs change, the hospital must submit appropriate data as follows:

(i) For hospital-specific data, the hospital must provide data from its most recently settled and most recently filed cost report.

(ii) For data on other hospitals, the hospital must base its application on the most recent revisions to the prospective payment rates for inpatient operating costs, as published in the FEDERAL REGISTER.

(e) Use of urban or other rural area’s wage index—(1) Criteria for use of area’s wage index. Except as provided in paragraphs (e)(3) and (e)(4) of this section, to use an area’s wage index, a hospital must demonstrate the following:

(i) The hospital’s incurred wage costs are comparable to hospital wage costs in an urban or other rural area;

(ii) The hospital has the necessary geographic relationship as specified in paragraphs (a) and (b) of this section;

(iii) One of the following conditions apply:

(A) With respect to redesignations for Federal fiscal year 1994 through 2001, the hospital’s average hourly wage is at least 108 percent of the average hourly wage of hospitals in the area in which the hospital is located; or

(B) With respect to redesignations for Federal fiscal year 2002 and later years, the hospital’s average hourly wage is, in the case of a hospital located in a rural area, at least 106 percent, and, in the case of a hospital located in an urban area, at least 108 percent of the average hourly wage of hospitals in the area in which the hospital is located; and

(iv) One of the following conditions apply:

(A) For redesignations effective before fiscal year 1999, the hospital’s average hourly wage weighted for occupational categories is at least 90 percent of the average hourly wages of hospitals in the area to which it seeks redesignation.

(B) With respect to redesignations for fiscal year 1994 through 2001, the hospital’s average hourly wage is equal to at least 84 percent of the average hourly wage of hospitals in the area to which it seeks redesignation.

(C) With respect to redesignations for fiscal year 2002 and later years, the hospital’s average hourly wage is equal to, in the case of a hospital located in
a rural area, at least 82 percent, and in the case of a hospital located in an urban area, at least 84 percent of the average hourly wage of hospitals in the area to which it seeks redesignation.

(2) Appropriate wage data. For a wage index change, the hospital must submit appropriate data as follows:

(i) For hospital-specific data, the hospital must provide data from the HCFA hospital wage survey used to construct the wage index in effect for prospective payment purposes during the fiscal year prior to the fiscal year for which the hospital requests reclas-sification.

(ii) For data of other hospitals, the hospital must provide data concerning the following:

(A) The average hourly wage in the area in which the hospital is located and the average hourly wage in the area to which the hospital seeks reclas-sification. The wage data are taken from the HCFA hospital wage survey used to construct the wage index in ef-fect for prospective payment purposes during the fiscal year prior to the fiscal year for which the hospital requests reclassification and;

(B) If the hospital is requesting reclassification under §412.230(e)(1)(iv)(B), occupational-mix data to demonstrate the average occupa-tional mix for each employment cat-egory in the area to which the hospital seeks reclas-sification. Occupational-mix data can be obtained from surveys conducted by the American Hospital Association.

(3) Rural referral center exception. If a hospital was ever a rural referral cen-ter, it does not have to demonstrate that it meets the criterion set forth in paragraph (e)(1)(iii) of this section concern-ing its average hourly wage.

(4) Special dominating hospital excep-tion. The requirements of paragraph (e)(1)(i) and (e)(1)(iii) of this section do not apply if a hospital meets the fol-low criteria:

(i) Its average hourly wage is at least 108 percent of the average hourly wage of all other hospitals in the area in which the hospital is located.

(ii) It pays at least 40 percent of the adjusted uninflated wages in the MSA.

(iii) It was approved for redesignation under this paragraph (e) for each year from fiscal year 1992 through fiscal year 1997.


§412.232 Criteria for all hospitals in a rural county seeking urban redesig-nation.

(a) Criteria. For all hospitals in a rural county to be redesignated to an urban area, the following conditions must be met:

(1) The county in which the hospitals are located must be adjacent to the MSA or NECMA to which they seek re-designation.

(2) All hospitals in a rural county must apply for redesignation as a group.

(3) The hospitals must demonstrate that the rural county in which they are located currently meets the criteria for metropolitan character under para-graph (b) of this section and the wage criteria under paragraph (c) of this sec-tion.

(4) The hospitals may be redesignated only if one of the following conditions is met:

(i) The pre-reclassified average hour-ly wage for the area to which they seek redesignation is higher than the pre-re-classified average hourly wage for the area in which they are currently lo-cated.

(ii) The standardized amount for the area to which they seek redesignation is higher than the standardized amount for the area in which they are located.

(b) Metropolitan character. The group of hospitals must demonstrate that the county in which the hospitals are lo-cated meets the standards for redesig-nation to an MSA or an NECMA as an outlying county that were published in the Federal Register on March 30, 1990 (55 FR 12154) using Bureau of the Census data or Bureau of Census esti-mates made after 1990.

(c) Wage criteria. In applying the fol-low numeric criteria, rounding of numbers to meet the qualifying per-centages is not permitted.

(1) Aggregate hourly wage. The aggregate average hourly wage for all hos-pitals in the rural county must be
equal to at least 85 percent of the average hourly wage in the adjacent urban area; or
(2) Aggregate hourly wage weighted for occupational mix. For redesignations effective before fiscal year 1999, the aggregate hourly wage for all hospitals in the rural county, weighted for occupational categories, is at least 90 percent of the average hourly wage in the adjacent urban area.

(d) Appropriate data. (1) Metropolitan character. (i) To meet the criteria in paragraph (b) of this section, the hospitals may submit data, estimates, or projections, made by the Bureau of the Census concerning population density or growth, or changes in designation of urban areas.
(ii) The MGCRB only considers data developed by the Bureau of the Census.
(2) Appropriate wage data. The hospitals must submit appropriate data as follows:
(i) For hospital-specific data, the hospitals must provide the following:
(A) The average hourly wage in the adjacent area, which is taken from the HCFA hospital wage survey used to construct the wage index in effect for prospective payment purposes during the fiscal year prior to the fiscal year for which the hospitals request reclassification.
(B) Occupational-mix data to demonstrate the average occupational mix for each employment category in the adjacent area. Occupational-mix data can be obtained from surveys conducted by the American Hospital Association.

§ 412.234 Criteria for all hospitals in an urban county seeking redesignation to another urban area.

(a) General criteria. For all prospective payment hospitals in an urban county to be redesignated to another urban area, the following conditions must be met:
(1) All hospitals in an urban county must apply for redesignation as a group.
(2) The county in which the hospitals are located must be adjacent to the urban area to which they seek redesignation.
(3) The county in which the hospitals are located must be part of the Consolidated Metropolitan Statistical Area (CMSA) that includes the urban area to which they seek redesignation.
(4) The hospitals may be redesignated only if one of the following conditions is met.
(i) The pre-reclassified average hourly wage for the area to which they seek redesignation is higher than the pre-reclassified average hourly wage for the area in which they are currently located.
(ii) The standardized amount for the area to which they seek redesignation is higher than the standardized amount for the area in which they are currently located.
(b) Wage criteria. In applying the following numeric criteria, rounding of numbers to meet the qualifying percentages is not permitted.
(1) Aggregate hourly wage. The aggregate average hourly wage of all hospitals in the urban county must be at least 85 percent of the average hospital hourly wage in the MSA or NECMA to which the hospitals in the county seek reclassification; or
(2) Aggregate hourly wage weighted for occupational mix. For redesignations effective before fiscal year 1999, the aggregate average hourly wage for all hospitals in the county must be at least 90 percent of the average hourly wage in the adjacent urban area.

(c) Standardized amount inpatient operating costs—(1) Criteria. The urban hospitals must demonstrate that their average incurred costs are more comparable to the amount the hospitals would be paid if they were reclassified than the amount they would be paid under their current classification.
(2) Demonstrating comparable costs. The urban hospitals demonstrate that their costs are more comparable to the average amount they would be paid if
they were reclassified if, on average, each hospital's case-mix adjusted cost per case is at least equal to the amount it would be paid under its current classification plus 75 percent of the difference between that amount and the amount the hospital would receive if it was reclassified.

(d) Appropriate data. (1) Wage data. The hospitals must submit appropriate wage data as provided for in §412.230(e)(2).

(2) Cost data. The hospitals must submit appropriate data as provided for in §412.230(d)(3).

§412.236 Alternative criteria for hospitals located in an NECMA.

(a) General. (1) An urban hospital whose designation is affected by the implementation of NECMAs may qualify for redesignation by meeting either the criteria in §412.230 or the criterion in paragraph (b) of this section.

(2) All the hospitals in a NECMA may qualify for redesignation by meeting the criteria in either §412.234 or in paragraph (c) of this section.

(b) Criterion applicable to an individual urban hospital in a NECMA. The hospital demonstrates that it would have been designated in a different urban area under the criteria for designating MSAs in New England.

(c) Criteria applicable to a group of hospitals in a NECMA. (1) All prospective payment hospitals in a NECMA must apply for redesignation.

(2) The hospitals must demonstrate that the NECMA to which they are designated would be combined as part of the NECMA to which they seek redesignation if the criteria for combining NECMAs were the same as the criteria used for combining MSAs.

(d) Appropriate data. (1) The MGCRB only considers population and commuting data developed by the Bureau of the Census.

(2) To meet the criterion in paragraph (b) of this section or the criteria in paragraph (c) of this section, hospitals must submit data from the Bureau of the Census.

§412.246 MGCRB members.

(a) Composition. The Medicare Geographical Classification Review Board (MGCRB) consists of five members, including a Chairman, all of whom are appointed by the Secretary. The members include two members who are representative of prospective payment system hospitals located in rural areas, and at least one individual who is knowledgeable in analyzing the costs of inpatient hospital services.

(b) Term of office. The term of office for an MGCRB member may not exceed 3 years. A member may serve more than one term. The Secretary may terminate a member’s tenure prior to its full term.

§412.248 Number of members needed for a decision or a hearing.

(a) A quorum. A quorum, consisting of at least a majority of the MGCRB members, one of whom is representative of rural hospitals if possible, is required for making MGCRB decisions.

(b) Number of members for a hearing. If less than a quorum is present for an oral hearing, the chairman with the consent of the hospital may allow those members present to conduct the hearing and to prepare a recommended decision, which is then submitted to a quorum.

§412.250 Sources of MGCRB's authority.

(a) Compliance. The MGCRB, in issuing decisions under section 1886(d)(10)(C) of the Act, complies with all the provisions of title XVIII and related provisions of the Act and implementing regulations, including the criteria and conditions located at §412.230 through §412.236, issued by the Secretary under the authority of section 1886(d)(10)(D) of the Act; and HCFA Rulings issued under the authority of the Administrator.
§ 412.252 Applications.

(a) By one hospital. An individual prospective payment system hospital seeking redesignation to a different rural or urban area has the right to submit an application to the MGCRB.

(b) By a group of hospitals. A group of hospitals has the right to submit an application to the MGCRB requesting redesignation of all prospective payment hospitals in a county if all prospective payment hospitals located in a county or in a NECMA agree to the request.

§ 412.254 Proceedings before MGCRB.

(a) On-the-record decision. The MGCRB will ordinarily issue an on-the-record decision without conducting an oral hearing. The MGCRB will issue a decision based upon all documents, data, and other written evidence and comments submitted timely to the MGCRB by the parties.

(b) Oral hearing. The MGCRB may hold an oral hearing on its own motion or if a party demonstrates to the MGCRB's satisfaction that an oral hearing is necessary.

§ 412.256 Application requirements.

(a) Written application. A request for redesignation must be in writing and must constitute a complete application in accordance with paragraph (b) of this section.

(1) An application must be mailed or delivered to the MGCRB, with a copy to HCFA, and may not be submitted through the facsimile (FAX) process or by other electronic means.

(2) A complete application must be received not later than the first day of the 13-month period preceding the Federal fiscal year for which redesignation is requested.

(3) The filing date of an application is the date the application is received by the MGCRB.

(b) Criteria for a complete application. An application is complete if the application from an individual hospital or from all hospitals in a county includes the following information:

(1) The Federal fiscal year for which the hospital is applying for redesignation.

(2) Which criteria constitute the basis of the request for redesignation.

(3) An explanation of how the hospital or hospitals meet the relevant criteria in §§ 412.230 through 412.236, including any necessary data to support the application.

(c) Opportunity to complete a submitted application. (1) The MGCRB will review an application within 15 days of receipt to determine if the application is complete. If the MGCRB determines that an application is incomplete and may dismiss the application if a complete application is not filed by September 1.

(2) At the request of the hospital, the MGCRB may, for good cause, grant a hospital that has submitted an application by September 1, an extension beyond September 1 to complete its application.

(d) Appeal of MGCRB dismissal. (1) The hospital may appeal the MGCRB dismissal to the Administrator within 15 days of the date of the notice of dismissal. (2) Within 20 days of receipt of the hospital's request for appeal, the Administrator will affirm the dismissal or reverse the dismissal and remand the case to the MGCRB to determine whether redesignation is appropriate.

(e) Notification of complete application. When the MGCRB determines that the hospital's application contains all the necessary elements for a complete application, it notifies the hospital in writing, with a copy to HCFA, that the application is complete and that the case may proceed to an MGCRB decision.

§ 412.258 Parties to MGCRB proceeding.
(a) The party or parties to an MGCRB proceeding are the hospital or group of hospitals requesting a change in geographic designation.
(b) HCFA has 30 days from the date of receipt of notice of a complete application to submit written comments and recommendations (with a copy to the hospital) for consideration by the MGCRB.
(c) The hospital has 15 days from the date of receipt of HCFA’s comments to submit written comments to the MGCRB, with a copy to HCFA, for the purpose of responding to HCFA’s comments.

§ 412.260 Time and place of the oral hearing.
If the MGCRB decides that an oral hearing is necessary, it sets the time and place for the hearing and notifies the parties in writing, with a copy to HCFA, not less than 10 days before the time scheduled for the hearing. The MGCRB may reschedule, adjourn, postpone, or reconvene the hearing provided that reasonable written notice is given to the parties, with a copy to HCFA.

§ 412.262 Disqualification of an MGCRB member.
(a) Grounds for disqualification. An MGCRB member may not participate in any decision in a case in which he or she may be prejudiced or partial with respect to a party or has any other interest in the case.
(b) Request for disqualification. If a party believes that an MGCRB member should not participate in a decision, the party submits the objection in writing to the MGCRB at its earliest opportunity, explaining the grounds for the request. HCFA may also submit such a suggestion to the MGCRB.
(c) Consideration by the MGCRB member. The MGCRB member will consider the objection and, at his or her discretion, either will proceed or withdraw.
(d) Consideration by the MGCRB. If the member does not withdraw, a party may petition the MGCRB for withdrawal and the MGCRB will consider the objection and rule on whether the member may participate in the decision before it decides the case.

§ 412.264 Evidence and comments in MGCRB proceeding.
(a) Submission by the parties. Before a decision is issued and during an oral hearing, the parties may present evidence or comments to the MGCRB regarding the matters at issue in the case.
(b) Content of evidence and comments. The MGCRB may receive evidence and comments without regard for the rules of evidence applicable to court procedures.
(c) Ex parte communications. (1) The members of the MGCRB and its staff may not consult or be consulted by an individual representing the interests of an applicant hospital or by any other individual on any matter in issue before the MGCRB without notice to the hospital or HCFA. If such communication occurs, the MGCRB will disclose it to the hospital or HCFA, as appropriate, and make it part of the record after the hospital or HCFA has had an opportunity to comment. MGCRB members and staff may not consider any information outside the record about matters concerning a hospital’s application for reclassification.
(2) The provisions in paragraph (c)(1) of this section do not apply to the following:
(i) Communications among MGCRB members and staff.
(ii) Communications concerning the MGCRB’s administrative functions or procedures.
(iii) Requests from the MGCRB to a party or HCFA for a document.
(iv) Material that the MGCRB includes in the record after notice and an opportunity to comment.
(d) MGCRB rulings on evidence and comments. The MGCRB rules upon the admissibility of evidence and comments and excludes irrelevant, immaterial, or unduly repetitious evidence and comments.

§ 412.266 Availability of wage data.
A hospital may obtain the average hourly wage data necessary to prepare its application to the MGCRB from
§ 412.268 Subpoenas.

(a) In general. When reasonably necessary for the full presentation of a case, and only after a pre-decision request for information or data has failed to produce the necessary evidence, either upon its own motion or upon the request of a party, the MGCRB may issue subpoenas for the attendance and testimony of witnesses, for an oral hearing or the production of books, records, correspondence, papers, or other documents that are relevant and material to any matter at issue.

(b) Content of request. The request must designate which witnesses or documents are to be produced, and describe addresses or locations with sufficient particularity to permit these witnesses or documents to be found. The request for a subpoena must state the pertinent facts that the party expects to establish by the requested witnesses or documents and whether these facts could be established by other evidence without the use of a subpoena.

(c) Issuance. Subpoenas are issued as provided in section 205(d) of the Act.

(d) Payment for subpoena cost. HCFA pays for the cost of issuing subpoenas and the fees and mileage of any witness who is subpoenaed, as provided in section 205(d) of the Act.

§ 412.270 Witnesses.

Witnesses at an oral hearing testify under oath or affirmation, unless excused by the MGCRB for cause. The MGCRB may examine the witnesses and may allow the parties or their representatives to also examine any witnesses called.

§ 412.272 Record of proceedings before the MGCRB.

A complete record of the proceedings before the MGCRB is made in all cases. The record will not be closed until a decision has been issued by the MGCRB. A transcription of an oral hearing will be made at a party's request, at the expense of the requesting party.

§ 412.273 Withdrawing an application.

(a) Timing of a withdrawal. The MGCRB allows a hospital, or group of hospitals, to withdraw its application if the request for withdrawal is submitted to the MGCRB during the following time periods:

(1) At any time before the MGCRB issues a decision on the application; or

(2) After the MGCRB issues a decision, provided that the request for withdrawal is received by the MGCRB within 45 days of publication of HCFA’s annual notice of proposed rulemaking concerning changes to the inpatient hospital prospective payment system and proposed payment rates for the fiscal year for which the application has been filed.

(b) Written request only. A request to withdraw an application must be made in writing to the MGCRB by all hospitals that are party to the application.

(c) Appeal of the MGCRB’s denial of a hospital’s request for withdrawal. (1) A hospital may file an appeal of the MGCRB’s denial of its request for withdrawal of an application to the Administrator. The appeal must be received within 15 days of the date of the notice of the denial.

(2) Within 20 days of receipt of the hospital’s request for appeal, the Administrator affirms or reverses the denial.


§ 412.274 Scope and effect of an MGCRB decision.

(a) Scope of decision. The MGCRB may affirm or change a hospital’s geographic designation. The MGCRB’s decision is based upon the evidence of record, including the hospital’s application and other evidence obtained or received by the MGCRB.

(b) Effective date and term of the decision. Any classification change is effective for one year beginning with discharges occurring on the first day (October 1) of the second Federal fiscal year following the Federal fiscal year in which the complete application is filed and ending effective at the end of that Federal fiscal year (the end of the next September 30).
§ 412.278 Administrator's review.

(a) Hospitals requests for review. A hospital or group of hospitals dissatisfied with the MGCRB's decision regarding its geographic designation may request the Administrator to review the MGCRB decision. (A hospital or group of hospitals may also request that the Administrator review the MGCRB's dismissal of an application as untimely filed or incomplete, as provided in §412.256(d).)

(b) Procedures for hospital's request for review. (1) The hospital's request for review must be in writing and sent to the Administrator, in care of the Office of the Attorney Advisor. The request must be received by the Administrator within 15 days after the date the MGCRB issues its decision. A request for Administrator review filed by facsimile (FAX) or other electronic means will not be accepted. The hospital must also mail a copy of its request for review to HCFA's Office of Payment Policy.

(2) The request for review may contain proposed findings of fact and conclusions of law, exceptions to the MGCRB's decision, and supporting reasons therefor.

(3) Within 15 days of receipt of the hospital's request for review, HCFA may submit to the Administrator, in writing, with a copy to the party, comments and recommendations concerning the hospital's submission.

(4) Within 10 days of receipt of HCFA's submission, the hospital may submit in writing, with a copy to HCFA, a response to the Administrator.

(c) Discretionary review by the Administrator. (1) The Administrator may, at his or her discretion, review any final decision of the MGCRB.

(2) The Administrator promptly notifies the hospital that he or she has decided to review a decision of the MGCRB. The notice of review indicates the particular issues to be considered and includes copies of any comments submitted to the Administrator by HCFA staff concerning the MGCRB decision.

(3) Within 15 days of the receipt of the Administrator's notice of review, the hospital may submit a response in writing to the Administrator, with a copy of HCFA.

(d) Criteria for discretionary review. In deciding whether to review an MGCRB decision, the Administrator normally considers whether it appears that any of the following situations apply:

(1) The MGCRB made an erroneous interpretation of law, regulation, or HCFA Ruling.

(2) The MGCRB's decision is not supported by substantial evidence.

(3) The case presents a significant policy issue having a basis in law and regulations, and review is likely to lead to issuance of a HCFA Ruling or other directive needed to clarify a provision in the law or regulations.

(4) The decision of the MGCRB requires clarification, amplification, or an alternative legal basis.

(5) The MGCRB has incorrectly extended its authority to a degree not provided for by law, regulation, or HCFA Ruling.
(e) Communication procedures. All communications between HCFA staff and the Administrator concerning the Administrator's review of an MGCRB decision must be in writing. As specified in paragraphs (b) and (c) of this section, copies of comments by HCFA staff are sent to applicant hospitals within 15 days of receipt of a hospital's request for review, or, in cases in which the Administrator decides to review a case at his or her discretion, are included with the Administrator's notice of review. In the event there are additional communications between HCFA staff and the Administrator concerning MGCRB decisions reviewed by the Administrator under paragraphs (b) or (c) of this section, HCFA furnishes copies of the communications to the hospital or group of hospitals.

(f) Administrator's decision. (1) The Administrator may not receive or consider any new evidence and must issue a decision based only upon the record as it appeared before the MGCRB and comments submitted under paragraphs (b)(2), (b)(3), (b)(4), (c)(2), and (c)(3) of this section.

(2) The Administrator issues a decision in writing to the party with a copy to HCFA—

(i) Not later than 90 days following receipt of the party's request for review; or

(ii) Not later than 105 days following issuance of the MGCRB decision in the case of review at the discretion of the Administrator.

(3) The Administrator's decision issued under §412.278 (a) or (c) is the final Departmental decision, unless it is amended under §412.278(g). The final Departmental decision is not subject to judicial review.

(4) The Administrator's decision is not subject to judicial review.

(g) Amendment of Administrator decision—(1) Hospital's request for amendment. The hospital may request the Administrator to amend the decision for the limited purpose of correcting mathematical or computational errors, or to correct the decision if the evidence that was considered in making the decision clearly shows on its face that an error was made. The following procedure is followed:

(i) The hospital's request for amendment must be received by the Administrator within 10 days after the date the Administrator issues a decision. The request for amendment must be in writing, with a copy to HCFA.

(ii) The Administrator promptly reviews the hospital's request and amends the decision, if necessary, within 5 days following receipt of the hospital's request for amendment.

(2) Discretionary review by the Administrator. Within 15 days following the issuance of the Administrator's decision, the Administrator, at his or her discretion, may amend the decision to correct mathematical or computational errors, or to correct the decision if the evidence that was considered in making the decision clearly shows on its face that an error was made. The Administrator's amended decision is final and is not subject to judicial review.


§ 412.280 Representation.

(a) General. A party may be represented by legal counsel or by any other person appointed to act as its representative at any proceeding before the MGCRB or the Administrator.

(b) Rights of a representative. A representative appointed by a party may accept or give on behalf of the party any request or notice connected with any proceeding before the MGCRB or the Administrator. A representative is entitled to present evidence and argument as to facts and law in any MGCRB proceeding affecting the party represented and to obtain information to the same extent as the party represented. Notice of any action or decision sent to the representative of a party has the same effect as if it had been sent to the party itself.

Subpart M—Prospective Payment System for Inpatient Hospital Capital Costs

SOURCE: 56 FR 34449, Aug. 30, 1991, unless otherwise noted.
§ 412.300 Scope of subpart and definition.

(a) Purpose. This subpart implements section 1886(g)(1)(A) of the Act by establishing a prospective payment system for inpatient hospital capital-related costs. Under this system, payment is made on the basis described in § 412.304 through § 412.374 for inpatient hospital capital-related costs furnished by hospitals subject to the prospective payment system under subpart B of this part.

(b) Definition. For purposes of this subpart, a new hospital means a hospital that has operated (under previous or present ownership) for less than 2 years. The following hospitals are not new hospitals:

1. A hospital that builds new or replacement facilities at the same or another location even if coincidental with a change of ownership, a change in management, or a lease arrangement.

2. A hospital that closes and subsequently reopens.

3. A hospital that has been in operation for more than 2 years but has participated in the Medicare program for less than 2 years.

4. A hospital that changes its status from a hospital that is excluded from the prospective payment systems to a hospital that is subject to the capital prospective payment systems.

§ 412.302 Introduction to capital costs.

(a) New capital costs. New capital costs are allowable Medicare inpatient hospital capital-related costs under subpart G of part 413 of this chapter that are related to assets that were first put in use for patient care after December 31, 1990 (except for such costs deemed to be old capital costs based on prior obligations as described in paragraph (c) of this section) and those allowable capital-related costs related to assets in use prior to December 31, 1990 that are excluded from the definition of old capital costs described in paragraphs (b) (2) through (5) of this section, or are betterment or improvement costs related to those old capital assets.

(b) Old capital costs. Except as provided in paragraph (c) of this section with respect to capital obligations that qualify for recognition as old capital, old capital costs are allowable capital-related costs for land and depreciable assets that were put in use for patient care on or before December 31, 1990. However, for a new hospital as defined in § 412.300(b), old capital costs are defined as those allowable capital-related costs for land and depreciable assets that were put in use for patient care on or before the later of December 31, 1990 or the last day of the hospital’s base year cost reporting period under § 412.328(a)(2). Old capital costs include the following:

1. Allowable depreciation on assets based on the useful life guidelines used to determine depreciation expense in the hospital’s base period.

2. Allowable capital-related interest expense. Except as provided below, the amount of allowable capital-related interest expense that will be recognized as old capital is limited to the amount the hospital was legally obligated to pay as of December 31, 1990. Any allowable interest expense in excess of this limitation will be recognized as new capital.

   (i) An increase in interest expense is recognized if the increase is due to periodic fluctuations of rates in variable interest rate loans or at the time of conversion from a variable rate loan to a fixed rate loan when no other changes in the terms of the loan are made.

   (ii) If the terms of a debt instrument are revised after December 31, 1990, the amount of interest that will be recognized as old capital during the transition cannot exceed the amount that would have been recognized during the same period prior to the revision of the debt instrument.

   (iii) If short-term financing was used to acquire old capital assets and the debt is extended or “rolled-over”, a portion of the extended debt will be recognized as old capital. The portion will equal the ratio of the net book value as of the beginning of the applicable cost reporting period for depreciable assets that were in use in the
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base year, to the net book value as of the beginning of the base year cost reporting period for those assets. The net book value for the base year will not be adjusted to exclude assets that have been fully depreciated or removed from service since the base year. If the debt is related to specific assets, the ratio will be determined based on the values for those assets. The ratio will exclude assets that were acquired with other identifiable debt instruments. For purposes of this paragraph, short term financing is a debt that becomes due in no later than the earlier of 5 years or half of the average useful life of the assets to which the debt is related.

(iv) If old capital indebtedness is commingled with new capital debt, the allowable interest expense will be apportioned to old capital costs based on the ratio of the portion of the loan principal related to old capital indebtedness to the total loan principal.

(v) Investment income, excluding income from funded depreciation accounts, is used to reduce old capital interest expense based on the ratio of total old capital interest expense to total allowable interest expense in each cost reporting period.

(3) Allowable capital-related lease and rental costs for land and depreciable assets that were obligated as of December 31, 1990.

(i) Lease renewals up to the annual lease payment level obligated as of December 31, 1990 are recognized provided the same asset remains in use, the asset has a useful life of at least 3 years, and the annual lease payment is $1,000 or more for each item or service.

(ii) If a hospital-owned asset is sold or given to another party and that same asset is then leased back by the hospital, the amount of allowable capital-related costs recognized as old capital costs is limited to the amount allowed for that asset in the last cost reporting period that it was owned by the hospital.

(iii) If an entire hospital is leased without assumption of the hospital’s asset costs after December 31, 1990, the amount of allowable capital-related costs recognized as old capital costs is limited to the amount allowed for old capital costs in the base year or the last cost reporting period these costs were recognized under this subpart, whichever is later.

(4) The portion of allowable costs for other capital-related expenses (including but not limited to, taxes, insurance, license and royalty fees on depreciable assets) resulting from applying the ratio of the hospital’s gross old asset value to total asset value in each cost reporting period.

(5) The appropriate portion of the capital-related costs of related organizations under §413.17 that would be recognized as old capital costs if these costs had been incurred directly by the hospital.

(6) Obligated capital costs that are recognized as old capital costs in accordance with paragraph (c) of this section.

(7) If a hospital had nonreimbursable costs applicable to an old capital asset as of December 31, 1990 that subsequently become allowable inpatient capital-related costs, the allowable costs for such an asset that are attributable to inpatient hospital services are recognized as old capital costs if a portion of the asset was in use for inpatient hospital care on December 31, 1990 and the costs meet all other provisions for recognition of old capital costs contained in this section.

(c) Obligated capital costs—(1) General rule. Under the conditions described below, capital-related costs attributable to assets that are put in use after December 31, 1990 may be recognized as old capital costs. Any allowable capital-related costs for these assets that are not recognized as old capital costs are recognized as new capital costs.

(i) Fixed assets. The costs of capital-related items and services defined in subpart G of part 413 for which there was a contractual obligation entered into by a hospital or related party with an outside, unrelated party for the construction, reconstruction, lease, rental, or financing of a fixed asset may be recognized as old capital costs if all the following conditions are met:

(A) The obligation must arise from a binding written agreement that was executed on or before December 31, 1990 and that obligates the hospital on or before December 31, 1990.
(B) The capital asset must be put in use for patient care before October 1, 1994 except as provided in paragraph (c)(1)(iv) of this section.

(C) The hospital notifies the intermediary of the existence of obligated capital costs as provided in paragraph (c)(1)(v) of this section.

(D) The amount that is recognized as old capital cost is limited to the lesser of the actual allowable costs when the asset is put in use or the estimated costs of the capital expenditure at the time it was obligated as provided in paragraph (c)(1)(vi) of this section.

(ii) Moveable equipment. Moveable equipment is recognized as old capital cost only if all of the conditions specified in paragraphs (c)(1)(i) (B) through (D) of this section are met and one of the following conditions is met:

(A) There was a binding contractual agreement that was executed on or before December 31, 1990 and obligates the hospital on or before December 31, 1990 for the lease or purchase of the item of equipment on or before December 31, 1990.

(B) There was a binding contractual agreement that was executed on or before December 31, 1990 and obligates the hospital on or before December 31, 1990 for financing the acquisition of the equipment; the item of equipment costs at least $100,000; and the item was specifically listed in an equipment purchase plan approved by the Board of Directors on or before December 31, 1990.

(iii) Agreements not recognized. Agreements for planning, design or feasibility that do not commit the hospital to undertake a project are not recognized as obligating capital expenditures for purposes of this subsection.

(iv) Extension of deadline. HCFA may extend the deadline in paragraph (c)(1)(i)(B) of this section, under which an asset must be put in use for patient care before October 1, 1994, to no later than September 30, 1996 for extraordinary circumstances beyond the hospital’s control. Extraordinary circumstances include, but are not limited to, a construction strike or atypically severe weather that significantly delayed completion of a construction project. Normal construction delays do not constitute extraordinary circumstances.

(A) The hospital must submit its request for an extended deadline with documentation of the extraordinary circumstances by the later of January 1, 1993 or 180 days after the extraordinary circumstance.

(B) The intermediary reviews the request and verifies the hospital’s documentation, and forwards the request to HCFA within 60 days. Within 90 days, HCFA notifies the intermediary of its decision and, if an extension is granted, of the revised deadline for putting the asset in use for patient care service.

(v) The hospital must submit to its intermediary the binding agreement and supporting documents that relate to the obligated capital expenditure by the later of October 1, 1992, or within 90 days after the start of the hospital’s first cost reporting period beginning on or after October 1, 1991. This documentation must include a project description (including details of any phased construction or financing) and an estimate of costs that were prepared no later than December 31, 1990.

(vi) Cost limitation—(A) Leases, Rentals or Purchases. The amount of obligated capital costs recognized as old capital costs cannot exceed the amount specified in the lease, rental, or purchase agreement. If moveable equipment is recognized as old capital under paragraph (c)(1)(ii)(B) of this section, the amount recognized as old capital costs cannot exceed the estimated cost identified in the equipment purchase plan approved by the hospital’s Board of Directors.

(B) Construction contracts. The amount of obligated capital costs recognized as old capital costs cannot exceed the estimated construction costs for the project as of December 31, 1990. Additional costs will be recognized as old capital costs only if the additional costs are directly attributable to changes in life safety codes or other building requirements established by government ordinance that occurred after the project was obligated.

(C) Financing costs. The amount of obligated interest expense that will be recognized as old capital costs cannot
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exceed the amount for which the hospital was legally obligated as of December 31, 1990 or, in the case of financing that is arranged after December 31, 1990 for a capital acquisition that was legally obligated as of December 31, 1990, the amount specified in a detailed financing plan approved by the hospital’s Board of Directors prior to January 1, 1991.

(vii) Determining old capital costs. (A) The intermediary determines whether the applicable criteria are met for recognition of obligated capital costs as old capital costs and the maximum allowable cost that will be recognized as old capital costs.

(B) The intermediary advises the hospital of its determination by the later of the end of the hospital’s first cost reporting period subject to the capital prospective payment system or 9 months after the receipt of the hospital’s notification under paragraph (c)(1)(v) of this section.

(C) The actual amount that will be recognized as old capital costs is based on the lesser of the allowable costs for the asset when it is put into patient use or the amounts determined under paragraph (c)(1)(vi) of this section.

(viii) Multi-phase project. If the hospital has a multi-phase capital project, the provisions of paragraphs (c)(1)(i) through (vii) of this section apply independently to each phase of the project.

(2) Lengthy certificate-of-need process. (i) If a hospital does not meet the criteria under paragraph (c)(1)(i) or paragraph (c)(1)(ii) of this section, but meets all of the following criteria, the estimated cost for the project as of December 31, 1990 may be recognized as old capital costs:

(A) The hospital received any required certificate of need approval on or before December 31, 1990.

(B) The hospital’s Board of Directors formally authorized the project with a detailed description of its scope and costs on or before December 31, 1990.

(C) The estimated cost of the project as of December 31, 1990 exceeds 5 percent of the hospital’s total patient revenues during its base year.

(D) The capitalized cost that had been incurred for the project as of December 31, 1990 exceeded the lesser of $750,000 or 10 percent of the estimated project cost.

(E) The hospital began actual construction or renovation (“groundbreaking”) on or before March 31, 1991.

(F) The project is completed before October 1, 1994.

(ii) The provisions of paragraphs (c)(1)(iv) through (viii) of this section apply to projects that meet the criteria in paragraph (c)(3)(i) of this section.

(3) Construction in process. (i) If a hospital that initiates construction on a capital project does not meet the requirements of paragraphs (c)(1)(i) or (ii) or (c)(2)(i) of this section, the project costs may be recognized as old capital costs if all the following conditions are met:

(A) The hospital received any required certificate of need approval on or before December 31, 1990.

(B) The hospital’s Board of Directors formally authorized the project with a detailed description of its scope and costs on or before December 31, 1990.

(C) The estimated cost of the project as of December 31, 1990 exceeds 5 percent of the hospital’s total patient revenues during its base year.

(D) The capitalized cost that had been incurred for the project as of December 31, 1990 exceeded the lesser of $750,000 or 10 percent of the estimated project cost.

(E) The hospital began actual construction or renovation (“groundbreaking”) on or before March 31, 1991.

(F) The project is completed before October 1, 1994.

(ii) The provisions of paragraphs (c)(1)(iv) through (viii) of this section apply to projects that meet the criteria in paragraph (c)(3)(i) of this section.

(d) Consistency in cost reporting—(1) General rule. For cost reporting periods beginning on or after October 1, 1991, and before October 1, 2001, the hospital must follow consistent cost finding
methods for classifying and allocating capital-related costs, except as otherwise provided in paragraph (d)(4) of this section.

(2) Old capital costs. Unless there is a change of ownership, the hospital must continue the same cost finding methods for old capital costs, including its practices for the direct assignment of capital-related costs and its cost allocation bases, that were in effect in the hospital’s last cost reporting period ending on or before October 1, 1991. If there is a change of ownership, the new owners may request that the intermediary approve a change in order to be consistent with their established cost finding practices.

(3) New capital costs. If a hospital desires to change its cost finding methods for new capital costs, the request for change must be made in writing to the intermediary prior to the beginning of the cost reporting period for which the change is to apply. The request must include justification as to why the change will result in more accurate and more appropriate cost finding. The intermediary will not approve the change unless it determines that there is reasonable justification for the change.

(4) Hospitals may elect the simplified cost allocation methodology under the terms and conditions provided in the instructions for HCFA Form 2552.

§ 412.304 Implementation of the capital prospective payment system.

(a) General rule. As described in §§412.312 through 412.370, effective with cost reporting periods beginning on or after October 1, 1991, HCFA pays an amount determined under the capital prospective payment system for each inpatient hospital discharge as defined in §412.4. This amount is in addition to the amount payable under the prospective payment system for inpatient hospital operating costs as determined under §412.63.

(b) Cost reporting periods beginning on or after October 1, 1991 and before October 1, 2001. For cost reporting periods beginning on or after October 1, 1991 and before October 1, 2001, the capital payment amount is based on either a combination of payments for old capital costs and new capital costs or a fully prospective rate, as determined under §§412.324 through 412.348.

(c) Cost reporting periods beginning on or after October 1, 2001. For cost reporting periods beginning on or after October 1, 2001, the capital payment amount is based solely on the Federal rate determined under paragraphs (a) and (b) of §412.308 and updated under paragraph (c) of §412.308.

(d) Interim payments. Interim payments are made to the hospital as provided in §412.116.

§ 412.308 Determining and updating the Federal rate.

(a) FY 1992 national average cost per discharge. HCFA determines the FY 1992 estimated national average cost per discharge by updating the discharge weighted national average Medicare inpatient hospital capital-related cost per discharge for FY 1989 by the estimated increase in Medicare inpatient hospital capital costs per discharge.

(b) Standard Federal rate. The standard Federal rate is used to determine the Federal rate each year under paragraph (c) of this section.

(1) HCFA determines the standard Federal rate by adjusting the FY 1992 estimated national average cost per discharge by a factor so that estimated aggregate payments based on the standard Federal rate adjusted by the payment adjustments described in §412.312(b) equal estimated aggregate payments based solely on the national average cost per discharge.

(2) Effective FY 1994, the standard Federal rate used to determine the Federal rate each year under paragraph (c) of this section is reduced by 7.4 percent.

(3) Effective FY 1996, the standard Federal rate used to determine the Federal rate each year under paragraph (c) of this section is reduced by 0.28
percent to account for the effect of the revised policy for payment of transfers under §412.4(d).

(4) Effective FY 1998, the unadjusted standard Federal capital payment rate in effect on September 30, 1997, used to determine the Federal rate each year under paragraph (c) of this section is reduced by 15.68 percent.

(5) For discharges occurring on or after October 1, 1997 through September 30, 2002, the unadjusted standard Federal capital payment rate as in effect on September 30, 1997, used to determine the Federal rate each year under paragraph (c) of this section is further reduced by 2.1 percent.

(c) The Federal rate. HCFA determines the Federal rate each year by adjusting the standard Federal rate by the following factors.

(1) Update factor. After FY 1992, HCFA updates the standard Federal rate as follows:

(i) FY 1993 through FY 1995. For FY 1993 through FY 1995, the standard Federal rate is updated based on a moving two-year average of actual increases in capital-related costs per discharge for the period three and four years before the fiscal year in question, excluding the portion of the increase attributable to changes in case mix.

(ii) Effective FY 1996. Effective FY 1996, the standard Federal rate is updated based on an analytical framework. The framework includes a capital input price index, which measures the annual change in the prices associated with capital-related costs during the year. HCFA adjusts the capital input price index rate of change to take into account forecast errors, changes in the case mix index, the effect of changes to DRG classification and relative weights, and allowable changes in the intensity of hospital services.

(2) Outlier payment adjustment factor. HCFA reduces the updated standard Federal rate by an adjustment factor equal to the estimated additional payments for exceptions under §412.348 determined as a proportion of total payments under the hospital-specific rate and Federal rate.

(4) Budget neutrality adjustment factor.

(i) For FY 1992 through FY 1995, HCFA adjusts the updated standard Federal rate by a budget neutrality factor determined under §412.352.

(ii) HCFA makes an adjustment to the Federal rate so that estimated aggregate payments for the fiscal year based on the Federal rate after any changes resulting from the annual reclassification and recalibration of the DRG weight in accordance with §412.60(e) and in the geographic adjustment factors described in §412.312(b)(2) equal estimated aggregate payments based on the Federal rate that would have been made without such changes.


§ 412.312 Payment based on the Federal rate.

(a) General. The payment amount for each discharge based on the Federal rate determined under §412.308(c) is determined under the following formula:

\[
\text{Payment} = \text{Federal rate} \times \text{DRG weight} \times \text{Geographic adjustment factor} \times \text{Large urban add-on} \times (1 + \text{Capital disproportionate share adjustment factor + capital indirect medical education adjustment factor}) \times \text{(for hospitals located in Alaska and Hawaii, a cost-of-living adjustment factor)} + \text{(Any applicable outlier payment)}.
\]

(b) Payment adjustments—(1) DRG weights. The relative resource requirements of the discharge are taken into account by applying the DRG weighting factor that is assigned to the discharge under §412.60.

(2) Geographic adjustment factors—(i) Local cost variation. A geographic adjustment factor is applied that takes into account geographic variation in costs.

(ii) Large urban add-on. An additional adjustment is made for hospitals located in a large urban area to reflect the higher costs incurred by hospitals located in those areas.
(iii) Cost-of-living adjustment. An additional adjustment is made for hospitals located in Alaska and Hawaii to account for the higher cost-of-living in those States.

(3) Disproportionate share adjustment. For hospitals with at least 100 beds located in an urban area and serving low-income patients, a disproportionate share adjustment factor is applied that reflects the higher costs attributable to furnishing services to low income patients.

(4) Indirect medical education adjustment. An additional adjustment is made based on the ratio of residents to the average daily patient census of the hospital to account for the indirect costs of medical education.

§ 412.316 Geographic adjustment factors.

(a) Local cost variation. HCFA adjusts for local cost variation based on the hospital wage index value that is applicable to the hospital under §412.63(k). The adjustment factor equals the hospital wage index value applicable to the hospital raised to the .6848 power and is applied to 100 percent of the Federal rate.

(b) Large urban location. HCFA provides an additional payment to a hospital located, for purposes of receiving payment under §412.63(a), in a large urban area equal to 3.0 percent of what would otherwise be payable to the hospital based on the Federal rate.

(c) Cost-of-living adjustment. HCFA provides an additional payment to a hospital located in Alaska and Hawaii equal to \(0.3152 \times (\text{cost-of-living adjustment factor used to determine payments under } \$412.115 - 1)\) percent.

§ 412.322 Disproportionate share adjustment factor.

(a) Criteria for classification. A hospital is classified as a "disproportionate share hospital" for the purposes of capital prospective payments if either of the following conditions is met:

(1) The hospital is located, for purposes of receiving payment under §412.63(a), in an urban area, has 100 or more beds as determined in accordance with §412.105(b) and serves low-income patients, as determined under §412.106(b).

(2) The hospital meets the criteria in §412.106(c)(2).

(b) Payment adjustment factor. (1) If a hospital meets the criteria in paragraph (a)(1) of this section for a disproportionate share hospital for purposes of capital prospective payments, the disproportionate share payment adjustment factor equals \(e^{(0.2025 \times \text{disproportionate patient percentage as determined under } \$412.106(b)(5))} - 1\)

(2) If a hospital meets the criteria in §412.106(c)(2) for purposes of hospital inpatient operating prospective payments, the disproportionate share adjustment factor is the factor that results from deeming the hospital to have the same disproportionate share patient percentage that would yield its operating disproportionate share adjustment.

§412.324 General description.

(a) Hospitals under Medicare in FY 1991. During the ten-year transition period, payments to a hospital with a hospital-specific rate below the Federal rate are based on the fully prospective payment methodology under §412.340 or for a hospital with a hospital-specific rate above the Federal rate, the hold-harmless payment methodology under §412.344.

(b) New hospitals. (1) A new hospital, as defined under §412.300(b), is paid 85 percent of its allowable Medicare inpatient hospital capital-related costs through its cost reporting period ending at least 2 years after the hospital accepts its first patient.

(2) For the third year through the remainder of the transition period, the hospital is paid based on the fully prospective payment methodology or the hold-harmless payment methodology using the base period determined under §412.328(a)(2).

(c) Hospitals with 52-53 week fiscal years ending September 25 through September 29. For purposes of this subpart, a hospital with a 52-53 week fiscal year period beginning September 26 through September 30, 1992 is deemed to have the same beginning date for all cost reporting periods beginning before October 1, 2000 (unless the hospital later changes its cost reporting period).

§412.328 Determining and updating the hospital-specific rate.

(a) Base-year cost reporting period. (1) Last 12 month cost reporting period ending on or before December 31, 1990. For each hospital, the intermediary uses the hospital’s latest 12-month or longer cost reporting period ending on or before December 31, 1990 as the base period to determine a hospital’s hospital-specific rate.

(2) New hospitals. The base-year cost reporting period for a new hospital is its 12-month cost reporting period (or a combination of cost reporting periods covering at least 12 months) that begins at least 1 year after the hospital accepts its first patient.

(3) Other hospitals. For other than a new hospital as defined in §412.300(b), if a hospital does not have a 12-month cost reporting period or does not have adequate Medicare utilization to file a cost report in a period ending on or before December 31, 1990, the hospital-specific rate is based on the hospital’s old capital costs (per discharge) in its first 12-month cost reporting period (or combination of cost reporting periods covering at least 12 months) ending after December 31, 1990.

(b) Base-year costs per discharge. (1) Base period allowable inpatient capital costs per discharge. (i) Determination. The intermediary determines the base period allowable inpatient capital costs per discharge for the hospital by dividing the hospital’s total allowable Medicare inpatient hospital capital-related cost in the base period by the number of Medicare discharges in the base period.

(ii) Disposal of assets in the base year. When a depreciable asset has been disposed of in the base year, only that portion of the gain or loss that is allocated to the base-year cost reporting period is reflected in the hospital-specific rate.

(iii) Disposal of assets subsequent to the base year. If an asset for which the Medicare program had recognized depreciation during the base year is disposed of subsequent to the base year, the hospital-specific rate will not be revised to recognize the portion of the gain or loss allocated to the base year.

(2) Discharges. For the purpose of determining a hospital’s base period capital costs per discharge, a discharge includes discharges as defined in §412.4(a) and transfers as defined in §412.4(b)(2), adjusted by the transfer adjustment factor that is determined under paragraph (b)(3) of this section.

(3) Transfer adjustment factor. (i) For base year cost reporting periods ending on or before December 31, 1990, HCFA uses the base year MEDPAR data received as of June 30, 1991 to develop an adjustment to discharges to account for transfers. HCFA divides the length of stay for each transfer case by the geometric mean length of stay for the DRG (but in no case using a number greater than 1.0) and assigns each non-transfer case a value of 1.0. To determine the transfer adjustment factor, HCFA adds together the adjusted discharges and divides the result by total discharges including transfers.

(ii) For base year cost reporting periods beginning before October 1, 1991, HCFA determines a transfer adjustment factor as described in paragraph (b)(3)(i) of this section for a hospital using the applicable base year MEDPAR data on file as of the December 31 or June 30 occurring at least 6 months after the close of the approved base year.

(iii) For base year cost reporting periods beginning on or after October 1, 1991, the intermediary determines the transfer adjustment factor in place of HCFA as described in paragraph (b)(3)(i) of this section based on the most recent billing data available as of the date of the final determination of the hospital-specific rate.

(c) Case-mix adjustment. (1) Determining transfer-adjusted case mix value. Step 1: For base year cost reporting periods ending on or before December 31, 1990, HCFA uses the base year MEDPAR data received as of June 30, 1991 to determine the hospital's transfer-adjusted case mix value. For base year cost reporting periods ending after December 31, 1990 and beginning before October 1, 1991, HCFA determines a transfer-adjusted case mix value for a hospital using the applicable base year MEDPAR data on file as of the December 31 or June 30 occurring at least 6 months after the close of the base year. For base year cost reporting periods beginning on or after October 1, 1991, the intermediary determines the transfer-adjusted case mix value based on the most recent billing data available as of the date of the final determination of the hospital-specific rate. HCFA, or the intermediary, as appropriate, multiplies the DRG weight for each case by one of the following factors:

(i) If the case is not a transfer, the factor equals 1.0.

(ii) If the case is a transfer, the factor equals the lesser of 1.0 or the ratio of the length of stay for the case divided by the geometric mean length of stay for the DRG.

Step 2: The products derived for all cases under Step 1 are added together and the result is divided by the adjusted discharges used to calculate the transfer adjustment factor determined under paragraph (b)(3) of this section.

(2) Adjusting base period capital costs per discharge by the hospital's transfer-adjusted case mix value. The intermediary divides the base period capital costs per discharge for each hospital as determined in paragraph (b) of this section by the hospital's transfer-adjusted case mix value for the cost reporting period determined under paragraph (c)(3) of this section.

(d) Updating to FY 1992. The intermediary updates the case mix-adjusted base period costs per discharge to FY 1992 based on the national average increase in Medicare inpatient capital costs per discharge as estimated by HCFA, excluding the portion of the increase in capital costs per discharge attributable to changes in case mix.

(e) Hospital-specific rate. The intermediary determines the hospital-specific rate each year by adjusting the amount determined under paragraph (d) of this section by the following factors:

(1) Update factor. After FY 1992, the intermediary updates the hospital-specific rate each year by adjusting the amount determined under paragraph (d) of this section by an adjustment factor.
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equal to the estimated additional payments for capital-related costs for exceptions under § 412.348, determined as a proportion of the total amount of payments under the hospital-specific rate and Federal rate.

(3) Budget neutrality adjustment factor. For FY 1992 through FY 1995, the intermediary adjusts the updated amount determined in paragraph (d) of this section by a budget neutrality adjustment factor determined under § 412.352.

(4) Payment for transfer cases. Effective FY 1996, the intermediary reduces the updated amount determined in paragraph (d) of this section by 0.28 percent to account for the effect of the revised policy for payment of transfers under § 412.4(d).

(5) Reduction of rate: FY 1998. Effective FY 1998, the unadjusted hospital-specific rate as in effect on September 30, 1997 described in paragraph (e)(1) of this section is reduced by 15.68 percent.

(6) Reduction of rate: FY 1998 through FY 2002. For discharges occurring on or after October 1, 1997 through September 30, 2002, the unadjusted hospital-specific rate in effect on September 30, 1997, described in paragraph (e)(1) of this section is further reduced by 2.1 percent.

(f) Redetermination of hospital-specific rate. (1) General. (i) Upon request by a hospital, the intermediary redetermines the hospital-specific rate to reflect an increase in old capital costs as determined in the cost reporting period subsequent to the base year. An increase in Medicare old capital cost per discharge that is related solely to a decline in utilization is not recognized as an increase in old capital costs for purposes of this section. New capital costs are excluded from the redetermination of the hospital-specific rate.

(ii) The hospital may request redetermination for any cost reporting period beginning subsequent to the base period but no later than the later of the hospital’s cost reporting period beginning in FY 1994 or the cost reporting period beginning after obligated capital that is recognized as old capital under § 412.302(b) is put in use.

(iii) The hospital must request a redetermination in writing no later than the date the cost report must be filed with the hospital’s intermediary for the first cost reporting period beginning on or after October 1, 1993 or the cost reporting period that will serve as the new base period, whichever is later. The hospital’s redetermination request must include the cost report for the new base period and an estimate of the revised hospital-specific rate indicating that the new rate exceeds the hospital’s current hospital-specific rate.

(2) Determination of old capital costs. The intermediary determines the hospital’s old capital costs for the subsequent cost reporting period that will serve as the new base period. The intermediary includes the costs of obligated capital that are recognized as old capital costs under § 412.302(b), excludes the costs of assets disposed of subsequent to the initial base year, and reflects changes in allowable old capital costs occurring subsequent to the initial base period.

(3) Redetermined hospital-specific rate. The intermediary redetermines the hospital-specific rate based on the old capital costs that are determined under paragraph (f)(2) of this section for the new base period. The intermediary—

(i) Divides the hospital’s old capital costs for the new base period by the number of Medicare discharges in that cost reporting period (consistent with paragraph (b) of this section);

(ii) Divides the old capital costs per discharge by the hospital’s transfer adjusted case-mix value for the new base period (consistent with paragraph (c) of this section);

(iii) Applies an update factor, if appropriate, to account for inflation occurring subsequent to the new base year, an exceptions payment adjustment factor, and a budget neutrality adjustment factor (consistent with paragraphs (d) and (e) of this section).

(4) Denial by intermediary. If the intermediary determines, after audit, that the revised hospital-specific rate is lower than the current hospital-specific rate, it advises the hospital that its request is denied and explains the basis for the denial.

(5) Implementation date. The redetermined hospital-specific rate applies to discharges occurring on or after the beginning date of the new base period.
(g) Review and revision of the hospital-specific rate. (1) Interim determination. The intermediary makes an interim determination of the hospital-specific rate based on the best data available and notifies the hospital at least 30 days before the beginning of the hospital's first cost reporting period beginning on or after October 1, 1991.

(2) Final determination. (i) The intermediary makes a final determination of the hospital-specific rate based on the final settlement of the base period cost report.

(ii) The final determination of the hospital-specific rate is effective retroactively to the beginning of the hospital's first cost reporting period beginning on or after October 1, 1991 or, in the case of a redetermination of the hospital-specific rate under §412.328, to the beginning of the new base period.

(iii) The final determination of the hospital-specific rate is subject to administrative and judicial review in accordance with subpart R of part 405 of this chapter, governing provider reimbursement determinations and appeals.

(iv) The intermediary adjusts the hospital-specific rate to reflect any revisions that result from administrative or judicial review of the final determination of hospital-specific rate. The revised determination is effective retroactively to the same extent as in paragraph (g)(2)(ii) of this section.

§ 412.331 Determining hospital-specific rates in cases of hospital merger, consolidation, or dissolution.

(a) New hospital merger or consolidation. If, after a new hospital accepts its first patient but before the end of its base year, it merges with one or more existing hospitals, and two or more separately located hospital campuses are maintained, the hospital-specific rate and payment determination for the merged entity are determined as follows—

(1) Post-merger base year payment methodology. The new campus is paid based on reasonable costs until the end of the new campus' base year. Effective with the first cost reporting period beginning after the the end of the new campus' base year, the intermediary determines a hospital-specific rate applicable to the new campus in accordance with §412.328, and then determines a revised hospital-specific rate for the merged entity in accordance with paragraph (a)(2) of this section.

(2) Revised hospital-specific rate. Using each hospital's base period data, the intermediary determines a combined average discharge-weighted hospital-specific rate.

(3) Post-base year payment determination. To determine the applicable payment methodology under §412.336 and for payment purposes under §412.340 or §412.344, the discharge-weighted hospital-specific rate determined by the intermediary is compared to the Federal rate. The revised payment methodology is effective on the first day of the cost reporting period beginning after the end of the new campus' base year.

(b) Hospital merger or consolidation. If, after the base year, two or more hospitals merge or consolidate into one hospital as provided for under §413.134(k) of this chapter and the provisions of paragraph (a) of this section do not apply, the intermediary determines a revised hospital-specific rate applicable to the combined facility under §412.328, which is effective beginning with the date of merger or consolidation. The following rules apply to the revised hospital-specific rate and payment determination:

(1) Revised hospital-specific rate. Using each hospital's base period data, the intermediary determines a combined average discharge weighted hospital-specific rate.

(2) Payment determination. The discharge-weighted hospital-specific rate determined by the intermediary is compared to the Federal rate to establish the appropriate payment methodology under §412.336 and for payment purposes under §§412.340 or 412.344. The revised payment methodology is effective as of the date of merger or consolidation.

(3) Old capital cost determination. The capital-related costs related to the assets of each merged or consolidated
§ 412.332 Payment based on the hospital-specific rate.

The payment amount for each discharge (as defined in § 412.4(a)) based on the hospital-specific rate determined under § 412.328 (e) or (f) is determined by multiplying the applicable hospital-specific rate by the DRG weighting factor applicable to the discharge under § 412.60 and the applicable hospital-specific rate percentage for the pertinent cost reporting period under § 412.340.

§ 412.336 Transition period payment methodologies.

(a) General. For discharges occurring in cost reporting periods beginning on or after October 1, 1991 and before October 1, 2001, a hospital is paid under one of two payment methodologies described in § 412.340 and § 412.344. Except as provided under paragraph (b) of this section, a hospital is paid under the same methodology throughout the transition period.

(1) Hospital-specific rate below the Federal rate. A hospital with a hospital-specific rate below the Federal rate (after taking into account the estimated effect of the payment adjustments and outlier payments) is paid under the fully prospective payment methodology as described in § 412.340.

(2) Hospital-specific rate above the Federal rate. A hospital with a hospital-specific rate that is above the Federal rate (after taking into account the estimated effect of the payment adjustments and outlier payments) is paid under the hold-harmless payment methodology as described in § 412.344.

(b) Special rule for revised hospital-specific rate. If a hospital with a hospital-specific rate below the Federal rate requests that its hospital-specific rate be redetermined, the redetermined hospital-specific rate is compared to the Federal rate that is applicable to the new base period (after taking into account the estimated effect of the payment adjustments and outlier payments). If the redetermined hospital-specific rate is higher than the Federal rate, the hospital is paid under the
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Hold-harmless methodology effective with the beginning of the new base period and continuing throughout the remainder of the transition.

(c) Interim and final determinations of applicable payment methodology—(1) Interim determination. The intermediary makes an interim determination of the applicable payment methodology based on the best data available and notifies the hospital of its determination at least 30 days before the beginning of the hospital’s first cost reporting period beginning on or after October 1, 1991.

(2) Final determination. (i) The intermediary makes a final determination of the applicable payment methodology based on its final determination of the hospital’s hospital-specific rate. The final determination of the applicable payment methodology is effective retroactively to the beginning of the hospital’s first cost reporting period beginning on or after October 1, 1991.

(ii) If the hospital-specific rate is reetermined in accordance with §412.328(f), the intermediary makes a new determination of the applicable payment methodology. The new determination is effective retroactively to the beginning of the new base period.

(iii) If the hospital-specific rate is revised under §412.328(g) as a result of administrative or judicial review, the intermediary makes a new determination of the applicable payment methodology. The new determination is effective retroactively to the beginning of the hospital’s first cost reporting period beginning on or after October 1, 1991 or to the beginning of the new base period.

(d) Special Rule for Redetermination of Hospital Payment Methodology. For cost reporting periods beginning on or after October 1, 1993, the intermediary redetermines the hospital payment methodologies to take into account the reduction to the standard Federal rate provided in §412.308(b)(2).

(1) For a hospital paid under the fully prospective payment methodology in the last cost reporting period beginning before October 1, 1993, the intermediary compares the hospital’s FY 1994 hospital-specific rate with the hospital’s FY 1994 Federal rate (after taking into account the estimated effect of the payment adjustments and outlier payments).

(i) A hospital with a FY 1994 hospital-specific rate that is above the FY 1994 adjusted Federal rate is paid under the hold-harmless payment methodology described in §412.344.

(ii) Subject to the provisions of §412.328(f), a hospital with a FY 1994 hospital-specific rate that is below the FY 1994 adjusted Federal rate continues to be paid under the fully prospective payment methodology as described in §412.340.

(iii) The intermediary notifies the hospital of the new determination of the hospital’s payment methodology within 90 days of the hospital’s first cost reporting period beginning on or after October 1, 1993.

(2) A hospital paid under the hold-harmless payment methodology in the last cost reporting period beginning before October 1, 1993, will continue to be paid in accordance with the provisions of §412.344.

(1) 85 percent of reasonable costs for old capital costs (100 percent for sole community hospitals) plus an amount for new capital costs based on a proportion of the Federal rate. The proportion is equal to the ratio of the hospital's Medicare inpatient costs for new capital to total Medicare inpatient capital costs; or
(2) 100 percent of the Federal rate.

(3) Exceptions. (i) A hospital that would receive higher payment under paragraph (a)(1) of this section may elect payment based on 100 percent of the Federal rate under paragraph (a)(2) of this section.
(ii) A hospital that does not maintain records that are adequate to identify its old capital costs is deemed to have elected payment per discharge based on 100 percent of the Federal rate.

(b) Continued basis of payment. A hospital paid based on 100 percent of the Federal rate during the later of its cost reporting period beginning in FY 1994 or its first cost reporting period beginning after obligated capital that is recognized as old capital under §412.302(b) is put in use continues to be paid on that basis in subsequent cost reporting periods during the transition period and does not receive a reasonable cost payment for old capital costs under paragraph (a)(1) of this section.

(c) Basis of determination. The determination under paragraph (a) of this section regarding which payment alternative is applicable is made without regard to additional payments under the exceptions process under §412.348.

(d) Interim and final payment determinations. (1) Using the best data available, the intermediary makes an interim payment determination under paragraph (a) of this section concerning the applicable payment alternative, and, in the case of payment under paragraph (a)(1) of this section, the payment amounts for old and new capital. The intermediary notifies the hospital of its determination at least 30 days before the beginning of the hospital's first cost reporting period beginning on or after October 1, 1991. The intermediary may revise its determination based on additional information submitted by the hospital and make appropriate adjustments retroactively.

(2) The final determination of the amount payable under paragraph (a) of this section is based on final settlement of the Medicare cost report for the applicable cost reporting period and is effective retroactively to the beginning of that cost reporting period. This final determination is subject to administrative and judicial review in accordance with subpart R of part 405 of this chapter, governing provider reimbursement determinations and appeals.

§412.348 Exception payments.
(a) Definitions. As used in this section—

Annual operating expenses. Annual operating expenses means the sum of net expenses for all reimbursable cost centers for a 12 month cost reporting period. Annual operating expenses are obtained from the Medicare cost report.

Average age of fixed assets. The average age of fixed assets is the ratio of accumulated depreciation for buildings and fixed equipment to current depreciation expense for buildings and fixed equipment. The average age of fixed assets is determined from information on the Medicare cost report.

Fixed assets. Fixed assets mean buildings and fixed equipment.

(b) Criterion for additional payment during the transition period. An additional payment is made to a hospital paid under either the fully prospective payment methodology or the hold-harmless payment methodology as determined under paragraph (c) of this section for cost reporting periods beginning on or after October 1, 1991, and before October 1, 2001.

(c) Minimum payment level by class of hospital. (1) HCFA establishes a minimum payment level by class of hospital. The minimum payment level for a hospital will equal a fixed percentage of the hospital's capital-related costs. The minimum payment levels may be no greater than the percentages of allowable capital-related costs that follow:

(i) 90 percent for sole community hospitals.
(ii) 80 percent for hospitals located in an urban area for purposes of §412.63(a)
with at least 100 beds, as determined under §412.105(b), that have a disproportionate share patient percentage of at least 20.2 percent as determined under §412.63(a) with at least 100 beds that qualify for disproportionate share payments under §412.106(c)(2).

(iii) 70 percent for all other hospitals.

(2) When it is necessary to adjust the minimum payment levels set by class of hospitals specified in paragraphs (c)(1)(i) and (g)(6) of this section, HCFA will adjust those levels for each class of hospitals in one percentage point increments as necessary to satisfy the requirement specified in paragraph (h) of this section that total estimated payments under the exception process not exceed 10 percent of the total estimated capital prospective payments (exclusive of hold-harmless payments for old capital) for the same fiscal year.

(d) Additional payments. A hospital is entitled to an additional payment if its capital payments for the cost reporting period would otherwise be less than the applicable minimum payment level. The additional payment equals the difference between the applicable minimum payment level and the capital payments that the hospital would otherwise receive minus any offset amount determined under paragraph (e)(2) of this section.

(e) Determining a hospital’s exception payment amount—(1) Cumulative comparison. For each cost reporting period beginning before October 1, 2001, the hospital’s exception payment is determined by comparing the cumulative payments made to the hospital under the capital prospective payment system to the cumulative minimum payment levels applicable to the hospital for each cost reporting period subject to the prospective payment system.

(2) Offsetting amounts. Any amount by which the hospital’s cumulative payments exceed its cumulative minimum payment levels is deducted from the additional payment that would otherwise be payable for a cost reporting period.

(f) Additional payment exception for extraordinary circumstances. (1) A hospital may request an additional payment if the hospital incurs unanticipated capital expenditures in excess of $5 million (net of proceeds from other payment sources such as insurance, litigation decisions and other State, local or Federal government funding programs) due to extraordinary circumstances beyond the hospital’s control. Extraordinary circumstances include, but are not limited to, a flood, fire, or earthquake.

(2) A hospital must apply to its HCFA Regional Office by the later of October 1, 1992 or 180 days after the extraordinary circumstance causing the unanticipated expenditures for a determination by HCFA of whether the hospital is eligible for an additional payment based on the nature of the circumstances and the amount of financial loss documented by the hospital.

(3) Except for sole community hospitals, the additional payment is based on a minimum payment amount of 85 percent for Medicare’s share of allowable capital-related costs attributable to the extraordinary circumstances. For sole community hospitals, the minimum payment amount is 100 percent.

(4) The minimum payment level applicable under paragraph (c)(1) of this section is adjusted to take into account the 85 percent minimum payment level (100 percent for sole community hospitals) under paragraph (f)(3) of this section for the unanticipated capital-related costs. The additional payment for the cost reporting period equals the difference between the adjusted minimum payment level and the capital payments the hospital would otherwise receive less any offset amount determined under paragraph (e)(2) of this section.

(g) Special exceptions process. For eligible hospitals that meet a project need requirement, a project size requirement, and, in the case of certain urban hospitals, meet an excess capacity test, an additional payment may be made for up to 10 years beyond the end of the capital prospective payment system transition period.

(1) Eligible hospitals. The following classes of hospitals are eligible to receive exceptions payments under this special exceptions provision:

(i) Sole community hospitals.
(ii) Hospitals located in an urban area under §412.63(a) with at least 100 beds, as determined under §412.105(b), that either have a disproportionate share of at least 20.2 percent as determined under §412.106(b) or qualify for disproportionate share payments under §412.106(c)(2).

(iii) Hospitals with a combined inpatient Medicare and Medicaid utilization of at least 70 percent.

(2) Project need requirement. A hospital must show that it has obtained any required approval from a State or local planning authority. If a hospital is not required to obtain approval from a planning authority, it must satisfy the age of asset test specified in paragraph (g)(3) of this section and, in the case of an urban hospital, the excess capacity test under paragraph (g)(4) of this section.

(3) Age of assets test. A hospital must show that its average age of fixed assets is at or above the 75th percentile for the hospital’s first cost reporting period beginning on or after October 1, 1991.

(4) Excess capacity test for urban hospitals. Urban hospitals that are not required to receive approval from a State or local planning authority must demonstrate that either—

(i) The overall average occupancy rate in its metropolitan statistical area is at least 80 percent; or

(ii) After completion of the project, its capacity is no more than 80 percent of its prior capacity (in terms of bed size).

(5) Project size requirement. A hospital must complete, during the period from the beginning of its first cost reporting period beginning on or after October 1, 1991 to the end of its last cost reporting period beginning on or after October 1, 2001, a project whose costs for replacement and/or renovation of fixed assets related to patient care are at least:

(i) $200 million; or

(ii) 100 percent of its operating cost during the first 12 month cost reporting period beginning on or after October 1, 1991.

(6) Minimum payment level. The minimum payment level for qualifying hospitals will be 70 percent.

(7) Limitation on the period for exception payments. A qualifying hospital may receive an exceptions payment for up to 10 years from the year in which it completes a project for replacement or renovation of capital assets that meets project need and project size requirements (and, if applicable, excess capacity test), provided that it completes the project no later than the end of the hospital’s last cost reporting period beginning before October 1, 2001. A project is considered to be completed when the assets are put into use for patient care.

(B) Determining a hospital’s exception payment amount—

(i) Cumulative comparison. For each cost reporting period, the hospital’s exception payment amount is determined by comparing the cumulative payments made to the hospital under the capital prospective payment system to the cumulative minimum payment levels applicable to the hospital for each cost reporting period subject to the prospective payment system.

(ii) Offsetting amounts. Offsetting amounts are applied in the following order—

(A) Any amount by which the hospital’s cumulative payments exceed its cumulative minimum payment levels is deducted from the additional payment that would otherwise be payable for the cost reporting period.

(B) Any amount by which the hospital’s current year Medicare inpatient operating and capital prospective payment system payments (excluding, if applicable, 75 percent of the hospital’s operating prospective payment system disproportionate share payments) exceed its Medicare inpatient operating and capital costs is deducted from the additional payment that would otherwise be payable for the cost reporting period.

For purposes of calculating the offset, the costs and payments for services that are not subject to the hospital inpatient prospective payment system are excluded.

(h) Limit on exception payments. Total estimated payments under the exception process may not exceed 10 percent of the total estimated capital prospective payments (exclusive of hold-harmless payments for old capital) for the same fiscal year.

For FY 1992 through FY 1995, HCFA will determine an adjustment to the hospital-specific rate and the Federal rate proportionately so that the estimated aggregate payments under this subpart for inpatient hospital capital costs each fiscal year will equal 90 percent of what HCFA estimates would have been paid for capital-related costs on a reasonable cost basis under §413.130 of this chapter.

Special Rules for Puerto Rico Hospitals

§412.370 General provisions for hospitals located in Puerto Rico.

Except as provided in §412.374, hospitals located in Puerto Rico are subject to the rules in this subpart governing the prospective payment system for inpatient hospital capital-related costs.

§412.374 Payments to hospitals located in Puerto Rico.

(a) Payments for capital-related costs to hospitals located in Puerto Rico that are paid under the prospective payment system are equal to the sum of the following:

(1) 50 percent of a Puerto Rico capital rate based on data from Puerto Rico hospitals only, which is determined in accordance with procedures for developing the Federal rate; and

(2) 50 percent of the Federal rate, as determined under §412.308.

(b) Effective for fiscal year 1998, the Puerto Rico capital rate described in paragraph (a) of this section in effect on September 30, 1997, is reduced by 15.68 percent.

(c) For discharges occurring on or after October 1, 1997 through September 30, 2002, the Puerto Rico capital rate described in paragraph (a) of this section in effect on September 30, 1997 is further reduced by 2.1 percent.

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§ 413.1 Introduction.

(a) Basis, scope, and applicability—(1) Statutory basis—(i) Basic provisions. (A) Section 1815 of the Act requires that the Secretary make interim payments to providers and periodically determine the amount that should be paid under Part A of Medicare to each provider for the services it furnishes. (B) Section 1814(b) of the Act (for Part A) and section 1833(a) (for Part B) provide for payment on the basis of the lesser of a provider's reasonable costs or customary charges. (C) Section 1861(v) of the Act defines "reasonable cost". (ii) Additional provisions. (A) Section 1138(b) of the Act specifies the conditions for Medicare payment for organ procurement costs. (B) Section 1814(j) of the Act provides for exceptions to the "lower of costs or charges" provisions. (C) Sections 1815(a) and 1833(e) of the Act provide the Secretary with authority to request information from providers to determine the amount of Medicare payment due providers. (D) Section 1833(a)(4) and (i)(3) of the Act provide for payment of a blended amount for certain surgical services furnished in a hospital's outpatient department. (E) Section 1833(n) of the Act provides for payment of a blended amount for outpatient hospital diagnostic procedures such as radiology. (F) Section 1834(c)(1)(C) of the Act establishes the method for determining Medicare payment for screening mammograms performed by hospitals. (G) Section 1834(g) of the Act provides that payment for critical access hospital (CAH) outpatient services is the reasonable costs of the CAH in providing these services, as determined in accordance with section 1861(v)(1)(A) of the Act and the applicable principles of cost reimbursement in this part and in part 415 of this chapter. (H) Section 1881 of the Act authorizes payment for services furnished to ESRD patients. (I) Section 1883 of the Act provides for payment for post-hospital SNF care furnished by a rural hospital that has swing-bed approval. (J) Sections 1886(a) and (b) of the Act impose a ceiling on the rate of increase in hospital inpatient costs. (K) Section 1886(h) of the Act provides for payment to a hospital for the services of interns and residents in approved teaching programs on the basis of a "per resident" amount. (2) Scope. This part sets forth regulations governing Medicare payment for services furnished to beneficiaries by—(i) Hospitals and critical access hospitals (CAHs); (ii) Skilled nursing facilities (SNFs); (iii) Home health agencies (HHAs); (iv) Comprehensive outpatient rehabilitation facilities (CORFs); (v) End-stage renal disease (ESRD) facilities; (vi) Providers of outpatient physical therapy and speech pathology services (OPTs); and (vii) Organ procurement agencies (OPAs) and histocompatibility laboratories. (3) Applicability. The payment principles and related policies set forth in this part are binding on HCFA and its fiscal intermediaries, on the Provider Reimbursement Review Board, and on the entities listed in paragraph (a)(2) of this section. (b) Reasonable cost reimbursement. Except as provided under paragraphs (c) through (h) of this section, Medicare is generally required, under section 1814(b) of the Act (for services covered under Part A) and under section 1833(a)(2) of the Act (for services covered under Part B) to pay for services furnished by providers on the basis of reasonable costs as defined in section 1861(v) of the Act, or the provider's customary charges for those services, if lower. Regulations implementing section 1861(v) are found generally in this part beginning at §413.5. (c) Outpatient maintenance dialysis and related services. Section 1881 of the Act authorizes special rules for the
computed.

Paragraph (d) Payment for inpatient hospital services.

(1) For cost reporting periods beginning before October 1, 1983, the amount paid for inpatient hospital services is determined on a reasonable cost basis.

(2) Payment to short-term general hospitals located in the 50 States and the District of Columbia for the operating costs of hospital inpatient services for cost reporting periods beginning on or after October 1, 1983, and for the capital-related costs of inpatient services for cost reporting periods beginning on or after October 1, 1991, are determined prospectively on a per discharge basis under part 412 of this chapter except as follows:

(i) Payment for capital-related costs for cost reporting periods beginning before October 1, 1991, medical education costs, kidney acquisition costs, and the costs of certain anesthesia services, is described in §412.113 of this chapter.

(ii) Payment to children's, psychiatric, rehabilitation and long-term hospitals (as well as separate psychiatric and rehabilitation units (distinct parts) of short-term general hospitals), which are excluded from the prospective payment system under subpart B of part 412 of this chapter, and to hospitals outside the 50 States and the District of Columbia is on a reasonable cost basis, subject to the provisions of §413.40.

(iii) Payment to hospitals subject to a State reimbursement control system is described in paragraph (e) of this section.

(e) State reimbursement control systems. Beginning October 1, 1983, Medicare reimbursement for inpatient hospital services may be made in accordance with a State reimbursement control system rather than under the Medicare reimbursement principles set forth in this part, if the State system is approved by HCFA. Regulations implementing this alternative reimbursement authority are set forth in subpart C of part 403 of this chapter.

(f) Payment for services furnished by HHAs. The amount paid for home health services as defined in section 1861(m) of the Act (except durable medical equipment and the covered osteoporosis drug as provided for in that section) that are furnished beginning on or after October 1, 2000 to an eligible beneficiary under a home health plan of care is determined according to the prospectively determined payment
§ 413.5 Cost reimbursement: General.

(a) In formulating methods for making fair and equitable reimbursement for services rendered beneficiaries of the program, payment is to be made on the basis of current costs of the individual provider, rather than costs of a past period or a fixed negotiated rate. All necessary and proper expenses of an institution in the production of services, including normal standby costs, are recognized. Furthermore, the share of the total institutional cost that is borne by the program is related to the care furnished beneficiaries so that no part of their cost would need to be borne by other patients. Conversely, costs attributable to other patients of the institution are not to be borne by the program. Thus, the application of this approach, with appropriate accounting support, will result in meeting actual costs of services to beneficiaries as such costs vary from institution to institution. However, payments to providers of services for services furnished Medicare beneficiaries are subject to the provisions of §§ 413.13 and 413.30.

(b) Putting these several points together, certain tests have been evolved for the principles of reimbursement and certain goals have been established that they should be designed to accomplish. In general terms, these are the tests or objectives:

1. That the methods of reimbursement should result in current payment so that institutions will not be disadvantaged, as they sometimes are under other arrangements, by having to put up money for the purchase of goods and services well before they receive reimbursement.

2. That, in addition to current payment, there should be retroactive adjustment so that increases in costs are taken fully into account as they actually occurred, not just prospectively.

3. That there be a division of the allowable costs between the beneficiaries of this program and the other patients of the provider that takes account of the actual use of services by the beneficiaries of this program and that is fair to each provider individually.

4. That there be sufficient flexibility in the methods of reimbursement to be used, particularly at the beginning of the program, to take account of the great differences in the present state of development of recordkeeping.

5. That the principles should result in the equitable treatment of both non-profit organizations and profit-making organizations.

6. That there should be a recognition of the need of hospitals and other providers to keep pace with growing needs and to make improvements.

(c) As formulated herein, the principles given recognition to such factors as depreciation, interest, bad debts, educational costs, compensation of owners, and an allowance for a reasonable return on equity capital (in the case of certain proprietary providers). With respect to allowable costs some items of inclusion and exclusion are:

1. An appropriate part of the net cost of approved educational activities will be included.

2. Costs incurred for research purposes, over and above usual patient care, will not be included.

3. [Reserved]

4. The value of services provided by nonpaid workers, as members of an organization (including services of members of religious orders) having an agreement with the provider to furnish such services, is includable in the amount that would be paid others for similar work.

5. Discounts and allowances received on the purchase of goods or services are reductions of the cost to which they relate.

6. Bad debts growing out of the failure of a beneficiary to pay the deductible, or the coinsurance, will be reimbursed (after bona fide efforts at collection).

7. Charity and courtesy allowances are not includable, although “fringe benefit” allowances for employees
§ 413.9 Cost related to patient care.

(a) Principle. All payments to providers of services must be based on the reasonable cost of services covered under Medicare and related to the care of beneficiaries. Reasonable cost includes all necessary and proper costs incurred in furnishing the services, subject to principles relating to specific items of revenue and cost. However, for cost reporting periods beginning after December 31, 1973, payments to providers of services are based on the lesser of the reasonable cost of services covered under Medicare and furnished to program beneficiaries or the customary charges to the general public for such services, as provided for in § 413.13.

(b) Definitions—(1) Reasonable cost. Reasonable cost of any services must be determined in accordance with regulations establishing the method or methods to be used, and the items to be included. The regulations in this part take into account both direct and indirect costs of providers of services. The objective is that under the methods of determining costs, the costs with respect to individuals covered by the program will not be borne by individuals not so covered, and the costs with respect to individuals not so covered will not be borne by the program. These regulations also provide for the making of suitable retroactive adjustments after the provider has submitted fiscal and statistical reports. The retroactive adjustment will represent the difference between the amount received by the provider during the year for covered services from both Medicare and the beneficiaries and the amount determined in accordance with an accepted method of cost apportionment to be the actual cost of services furnished to beneficiaries during the year.

(2) Necessary and proper costs. Necessary and proper costs are costs that are appropriate and helpful in developing and maintaining the operation of
§ 413.13 Amount of payment if customary charges for services furnished are less than reasonable costs.

(a) Definitions. As used in this section—

Customary charges means the regular rates that providers charge both beneficiaries and other paying patients for the services furnished to them.

Fair compensation means the reasonable cost of covered services.

Nominal charge means a charge equal to 60 percent or less of the reasonable cost of a service.

Public provider means a provider operated by a Federal, State, county, city, or other local government agency or instrumentality.

Reasonable cost means cost actually incurred, to the extent that cost is necessary for the efficient delivery of the service, and subject to the exclusions specified in paragraph (d) of this section.

(b) Application of the lesser of costs or charges (LCC) principle.—(1) General rule. Except as provided in paragraph (c) of this section, HCFA pays providers the lesser of the reasonable cost or the customary charges for services furnished to Medicare beneficiaries. Reasonable cost and customary charges are compared separately for Part A services and Part B services.

(2) Example. (i) A provider’s reasonable cost for covered services furnished to Medicare beneficiaries during a cost reporting period is $125,000.

(ii) The provider’s customary charges for those services is $110,000.

(iii) HCFA pays the provider $110,000 less the deductible and coinsurance amounts for which the beneficiaries are responsible.

(c) Exceptions to the LCC principle. (1) Providers not subject to the LCC principle.

HCFA pays the following providers the fair compensation for the services they furnish:

(i) CORFs.

(ii) Public providers that furnish services free of charge or at a nominal charge.

(iii) Any provider that requests payment of fair compensation and can demonstrate to its intermediary that a significant portion of its patients are

§ 413.13 Amount of payment if customary charges for services furnished are less than reasonable costs.

(a) Definitions. As used in this section—

Customary charges means the regular rates that providers charge both beneficiaries and other paying patients for the services furnished to them.

Fair compensation means the reasonable cost of covered services.

Nominal charge means a charge equal to 60 percent or less of the reasonable cost of a service.

Public provider means a provider operated by a Federal, State, county, city, or other local government agency or instrumentality.

Reasonable cost means cost actually incurred, to the extent that cost is necessary for the efficient delivery of the service, and subject to the exclusions specified in paragraph (d) of this section.

(b) Application of the lesser of costs or charges (LCC) principle.—(1) General rule. Except as provided in paragraph (c) of this section, HCFA pays providers the lesser of the reasonable cost or the customary charges for services furnished to Medicare beneficiaries. Reasonable cost and customary charges are compared separately for Part A services and Part B services.

(2) Example. (i) A provider’s reasonable cost for covered services furnished to Medicare beneficiaries during a cost reporting period is $125,000.

(ii) The provider’s customary charges for those services is $110,000.

(iii) HCFA pays the provider $110,000 less the deductible and coinsurance amounts for which the beneficiaries are responsible.

(c) Exceptions to the LCC principle. (1) Providers not subject to the LCC principle.

HCFA pays the following providers the fair compensation for the services they furnish:

(i) CORFs.

(ii) Public providers that furnish services free of charge or at a nominal charge.

(iii) Any provider that requests payment of fair compensation and can demonstrate to its intermediary that a significant portion of its patients are
low income and that its charges are less than costs because its customary practice is to charge patients on the basis of their ability to pay.

(2) Services not subject to the LCC principle. The following services are not subject to the LCC principle:

(i) Part A inpatient hospital services. Inpatient hospital services are not subject to the LCC principle if they are subject to either of the following:

(A) The prospective payment system under part 422 of this chapter.

(ii) Facility services related to ambulatory surgical procedures performed in outpatient hospital departments. Facility services related to ambulatory surgical procedures performed in hospital outpatient departments are subject to the payment methodology set forth in §413.101.

(iii) Skilled nursing facility services. Skilled nursing facility services subject to the payment methodology set forth in §413.106.

(iv) Hospital outpatient radiology services. Hospital outpatient radiology services are subject to the payment methodology set forth in §413.102.

(vi) Skilled nursing facility services. Skilled nursing facility services subject to the payment methodology set forth in §413.106.

(d) Exclusions from reasonable cost. For purposes of comparison with customary charges under this section, reasonable cost does not include the following:

(1) Payments made to a provider as reimbursement for bad debts arising from noncollection of Medicare deductible and coinsurance amounts, as provided in §413.80.

(2) Amounts that represent the recovery of excess depreciation resulting from termination from the Medicare program or a decrease in Medicare utilization applicable to prior cost reporting periods, as provided in §413.134.

(4) Payments to funds for the donated services of teaching physicians, as provided in §413.85.

(5) Except as provided in paragraph (f)(2)(iii) of this section for making nominal charge determinations in special situations, graduate medical education costs.

(e) Reductions in customary charges. Customary charges are reduced in proportion to the ratio of the aggregate amount actually collected from charge-paying non-Medicare patients to the amount that would have been realized had customary charges been paid, if the provider—

(1) Did not actually impose charges on most of the patients liable for payment for its services on a charge basis; or

(2) Failed to make a reasonable effort to collect those charges.

(f) Nominal charge determinations. In determining whether a provider's customary charges equal 60 percent or less of its reasonable costs, the following rules apply:

(1) General rule. The determination is based on charges actually billed to charge-paying, non-Medicare patients, and (except for clinical diagnostic laboratory tests that are paid under section 1833(h) of the Act) is made separately for Part A services and Part B services.

(2) Determination in special situations.

(i) Charges based on ability to pay. For providers that have a sliding scale or discounted charges based on patients' ability to pay, the determination—

(A) Is based on charges billed to all charge-paying patients;

(B) Uses the ratio of the sliding scale charges to the provider's full customary charges; and

(C) Applies the ratio to the discounted charges to equate those charges to customary charges.

(ii) HHA services. In determining nominal charges for HHAs, all Part A and Part B services, with the exception of DME, are considered together.

(iii) Graduate medical education. When making the nominal charge determination, graduate medical education payments (or the provider's reasonable costs for that education, if supported
§ 413.17 Cost to related organizations.

(a) Principle. Except as provided in paragraph (d) of this section, costs applicable to services, facilities, and supplies furnished to the provider by organizations related to the provider by common ownership or control are includable in the allowable cost of the provider at the cost to the related organization. However, such cost must not exceed the price of comparable services, facilities, or supplies that could be purchased elsewhere.

(b) Definitions.

(1) Related to the provider. Related to the provider means that the provider to a significant extent is associated or affiliated with or has control of or is controlled by the organization furnishing the services, facilities, or supplies.

(2) Common ownership. Common ownership exists if an individual or individuals possess significant ownership or equity in the provider and the institution or organization serving the provider.

(3) Control. Control exists if an individual or an organization has the power, directly or indirectly, significantly to influence or direct the actions or policies of an organization or institution.

(c) Application. (1) Individuals and organizations associate with others for various reasons and by various means. Some deem it appropriate to do so to assure a steady flow of supplies or services, to reduce competition, to gain a tax advantage, to extend influence, and for other reasons. These goals may be accomplished by means of ownership or control, by financial assistance, by management assistance, and other ways.

(2) If the provider obtains items of services, facilities, or supplies from an organization, even though it is a separate legal entity, and the organization is owned or controlled by the owner(s) of the provider, in effect the items are obtained from itself. An example would be a corporation building a hospital or a nursing home and then leasing it to another corporation controlled by the owner. Therefore, reimbursable cost should include the costs for these items at the cost to the supplying organization. However, if the price in the open market for comparable services, facilities, or supplies is lower than the cost to the supplier, the allowable cost to the provider may not exceed the market price.

(d) Exception. (1) An exception is provided to this general principle if the provider demonstrates by convincing evidence to the satisfaction of the fiscal intermediary (or, if the provider has not nominated a fiscal intermediary, HCFA), that—

(i) The supplying organization is a bona fide separate organization;

(ii) A substantial part of its business activity of the type carried on with the provider is transacted with others than the provider and organizations related to the supplier by common ownership or control and there is an open, competitive market for the type of services, facilities, or supplies furnished by the organization;

(iii) The services, facilities, or supplies are those that commonly are obtained by institutions such as the provider from other organizations and are not a basic element of patient care ordinarily furnished directly to patients by such institutions; and

(iv) The charge to the provider is in line with the charge for such services, facilities, or supplies in the open market and no more than the charge made under comparable circumstances to others by the organization for such services, facilities, or supplies.

(2) In such cases, the charge by the supplier to the provider for such services, facilities, or supplies is allowable as cost.

Subpart B—Accounting Records and Reports

§ 413.20 Financial data and reports.

(a) General. The principles of cost reimbursement require that providers maintain sufficient financial records and statistical data for proper determination of costs payable under the program. Standardized definitions, accounting, statistics, and reporting practices that are widely accepted in the hospital and related fields are followed. Changes in these practices and
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systems will not be required in order to determine costs payable under the principles of reimbursement. Essentially the methods of determining costs payable under Medicare involve making use of data available from the institution's basis accounts, as usually maintained, to arrive at equitable and proper payment for services to beneficiaries.

(b) Frequency of cost reports. Cost reports are required from providers on an annual basis with reporting periods based on the provider's accounting year. In the interpretation and application of the principles of reimbursement, the fiscal intermediaries will be an important source of consultative assistance to providers and will be available to deal with questions and problems on a day-to-day basis.

(c) Recordkeeping requirements for new providers. A newly participating provider of services (as defined in §400.202 of this chapter) must make available to its selected intermediary for examination its fiscal and other records for the purpose of determining such provider's ongoing recordkeeping capability and inform the intermediary of the date its initial Medicare cost reporting period ends. This examination is intended to assure that—

(1) The provider has an adequate ongoing system for furnishing the records needed to provide accurate cost data and other information capable of verification by qualified auditors and adequate for cost reporting purposes under section 1815 of the Act; and

(2) No financial arrangements exist that will thwart the commitment of the Medicare program to reimburse providers the reasonable cost of services furnished beneficiaries. The data and information to be examined include cost, revenue, statistical, and other information pertinent to reimbursement including, but not limited to, that described in paragraph (d) of this section and in §413.24.

(d) Continuing provider recordkeeping requirements. (1) The provider must furnish such information to the intermediary as may be necessary to—

(i) Assure proper payment by the program, including the extent to which there is any common ownership or control (as described in §413.17(b)(2) and (3)) between providers or other organizations, and as may be needed to identify the parties responsible for submitting program cost reports;

(ii) Receive program payments; and

(iii) Satisfy program overpayment determinations.

(2) The provider must permit the intermediary to examine such records and documents as are necessary to ascertain information pertinent to the determination of the proper amount of program payments due. These records include, but are not limited to, matters pertaining to—

(i) Provider ownership, organization, and operation;

(ii) Fiscal, medical, and other recordkeeping systems;

(iii) Federal income tax status;

(iv) Asset acquisition, lease, sale, or other action;

(v) Franchise or management arrangements;

(vi) Patient service charge schedules;

(vii) Costs of operation;

(viii) Amounts of income received by source and purpose; and

(ix) Flow of funds and working capital.

(3) The provider, upon request, must furnish the intermediary copies of patient service charge schedules and changes thereto as they are put into effect. The intermediary will evaluate such charge schedules to determine the extent to which they may be used for determining program payment.

(e) Suspension of program payments to a provider. If an intermediary determines that a provider does not maintain or no longer maintains adequate records for the determination of reasonable cost under the Medicare program, payments to such provider will be suspended until the intermediary is assured that adequate records are maintained. Before suspending payments to a provider, the intermediary will, in accordance with the provisions in §405.372(a) of this chapter, send written notice to such provider of its intent to suspend payments. The notice will explain the basis for the intermediary's determination with respect to the provider's records and will identify the provider's recordkeeping deficiencies. The provider must be given the opportunity, in accordance with §405.372(b)
of this chapter, to submit a statement (including any pertinent evidence) as to why the suspension must not be put into effect.

§ 413.24 Adequate cost data and cost finding.

(a) Principle. Providers receiving payment on the basis of reimbursable cost must provide adequate cost data. This must be based on their financial and statistical records which must be capable of verification by qualified auditors. The cost data must be based on an approved method of cost finding and on the accrual basis of accounting. However, if governmental institutions operate on a cash basis of accounting, cost data based on such basis of accounting will be acceptable, subject to appropriate treatment of capital expenditures.

(b) Definitions. (1) Cost finding. Cost finding is the process of recasting the data derived from the accounts ordinarily kept by a provider to ascertain costs of the various types of services furnished. It is the determination of these costs by the allocation of direct costs and proration of indirect costs.

(2) Accrual basis of accounting. As used in this part, the term accrual basis of accounting means that revenue is reported in the period in which it is earned, regardless of when it is collected; and an expense is reported in the period in which it is incurred, regardless of when it is paid. (See §413.100 regarding limitations on allowable accrued costs in situations in which the related liabilities are not liquidated timely.)

(c) Adequacy of cost information. Adequate cost information must be obtained from the provider’s records to support payments made for services furnished to beneficiaries. The requirement of adequacy of data implies that the data be accurate and in sufficient detail to accomplish the purposes for which it is intended. Adequate data capable of being audited is consistent with good business concepts and effective and efficient management of any organization, whether it is operated for profit or on a nonprofit basis. It is a reasonable expectation on the part of any agency paying for services on a cost-reimbursement basis. In order to provide the required cost data and not impair comparability, financial and statistical records should be maintained in a manner consistent from one period to another. However, a proper regard for consistency need not preclude a desirable change in accounting procedures if there is reason to effect such change.

(d) Cost finding methods. After the close of the accounting period, providers must use one of the following methods of cost finding to determine the actual costs of services furnished during that period. (These provisions do not apply to SNFs that elect and qualify for prospectively determined payment rates under subpart I of this part for cost reporting periods beginning on or after October 1, 1986. For the special rules that are applicable to those SNFs, see §413.321.) For cost reporting periods beginning after December 31, 1971, providers using the departmental method of cost apportionment must use the step-down method described in paragraph (d)(1) of this section or an “other method” described in paragraph (d)(2) of this section. For cost reporting periods beginning after December 31, 1971, providers using the combination method of cost apportionment must use the modified cost finding method described in paragraph (d)(3) of this section. Effective for cost reporting periods beginning on or after October 1, 1980, HHAs not based in hospitals or SNFs must use the step-down method described in paragraph (d)(1) of this section. (HHAs based in hospitals or SNFs must use the method applicable to the parent institution.) However, an HHA not based in a hospital or SNF that received less than $35,000 in Medicare payment for the immediately preceding cost reporting period, and for whom this payment represented less than 50 percent of the total operating cost of the agency, may use a simplified version of the step-down method, as specified in instructions for the cost report issued by HCFA.

(1) Step-down Method. This method recognizes that services furnished by certain nonrevenue-producing departments or centers are utilized by certain other nonrevenue-producing centers as
well as by the revenue-producing centers. All costs of nonrevenue-producing centers are allocated to all centers that they serve, regardless of whether or not these centers produce revenue. The cost of the nonrevenue-producing center serving the greatest number of other centers, while receiving benefits from the least number of centers, is apportioned first. Following the apportionment of the cost of the nonrevenue-producing center, that center will be considered “closed” and no further costs are apportioned to that center. This applies even though it may have received some service from a center whose cost is apportioned later. Generally, if two centers furnish services to an equal number of centers while receiving benefits from an equal number, that center which has the greatest amount of expense should be allocated first.

(2) Other methods. (i) The double-apportionment method. The double-apportionment method may be used by a provider upon approval of the intermediary. This method also recognizes that the nonrevenue-producing departments or centers furnish services to other nonrevenue-producing centers as well as to revenue-producing centers. A preliminary allocation of the costs of non-revenue-producing centers is made. These centers or departments are not “closed” after this preliminary allocation. Instead, they remain “open,” accumulating a portion of the costs of all other centers from which services are received. Thus, after the first or preliminary allocation, some costs will remain in each center representing services received from other centers. The first or preliminary allocation is followed by a second or final apportionment of expenses involving the allocation of all costs remaining in the nonrevenue-producing functions directly to revenue-producing centers.

(ii) More sophisticated methods. A more sophisticated method designed to allocate costs more accurately may be used by the provider upon approval of the intermediary. However, having elected to use the double-apportionment method, the provider may not thereafter use the step-down method without approval of the intermediary. Written request for the approval must be made on a prospective basis and must be submitted before the end of the fourth month of the prospective reporting period. Likewise, once having elected to use a more sophisticated method, the provider may not thereafter use either the double-apportionment or step-down methods without similar request and approval.

(3) Modified cost finding for providers using the Combination Method for reporting periods beginning after December 31, 1971. This method differs from the step-down method in that services furnished by nonrevenue-producing departments or centers are allocated directly to revenue-producing departments or centers even though these services may be utilized by other nonrevenue-producing departments or centers. In the application of this method the cost of nonrevenue-producing centers having a common basis of allocation are combined and the total distributed to revenue-producing centers. All nonrevenue-producing centers having significant percentages of cost in relation to total costs will be allocated this way. The combined total costs of remaining non-revenue-producing costs centers will be allocated to revenue-producing cost centers in the proportion that each bears to total costs, direct and indirect, already allocated. The bases which are to be used and the centers which are to be combined for allocation are not optional but are identified and incorporated in the cost report forms developed for this method. Providers using this method must use the program cost report forms devised for it. Alternative forms may not be used without prior approval by HCFA based upon a written request by the provider submitted through the intermediary.

(4) Temporary method for initial period. If the provider is unable to use either cost-finding method when it first participates in the program, it may apply to the intermediary for permission to use some other acceptable method that would accurately identify costs by department or center, and appropriately segregate inpatient and outpatient costs. Such other method may be used for cost reports covering periods ending before January 1, 1968.
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(5) Simplified optional reimbursement method for small, rural hospitals with distinct parts for cost reporting periods beginning on or after July 20, 1982. (i) A rural hospital with a Medicare-certified distinct part SNF may elect to be reimbursed for services furnished in its hospital general routine service area and distinct part SNF using the reimbursement method specified in § 413.53 for swing-bed hospitals, if it meets the following conditions:

(A) The institution is located in a rural area as defined in § 482.66 of this chapter.

(B) On the first day of the cost reporting period, the hospital and distinct part SNF have fewer than 50 beds in total (with the exception of beds for newborns and beds in intensive care type inpatient units).

(ii) In applying the optional reimbursement method, only those beds located in the hospital general routine service area and in the distinct part SNF certified by Medicare are combined into a single cost center for purposes of cost finding.

(iii) The reasonable cost of the routine extended care services is determined in accordance with § 413.114(c). The reasonable cost of the hospital general routine services is determined in accordance with § 413.53(a)(2).

(iv) The hospital must make its election to use the optional swing-bed reimbursement method in writing to the intermediary before the beginning of the hospital’s cost reporting year. The hospital must make any request to revoke the election in writing before the beginning of the affected cost reporting period.

(v) The intermediary must approve requests to terminate use of the optional swing-bed reimbursement method. If a hospital terminates use of this optional method, no further elections may be made by the facility to use the optional method.

(6) Management contracts. (i) If the main provider purchases services for a department of the provider or a provider-based entity through a management contract, the like costs of the main provider purchased through a management contract must be included in the main provider’s administrative and general costs and allocated among the provider’s overall statistics.

(ii) Costs of free-standing entities may not be shown in the provider’s trial balance for purposes of stepping down overhead costs to these entities. However, governmental institutions that operate on a cash basis of accounting may submit cost data on the cash basis subject to appropriate treatment of capital expenditures.

(e) Accounting basis. The cost data submitted must be based on the accrual basis of accounting which is recognized as the most accurate basis for determining costs. However, governmental institutions that operate on a cash basis of accounting may submit cost data on the cash basis subject to appropriate treatment of capital expenditures.

(f) Cost reports. For cost reporting purposes, the Medicare program requires each provider of services to submit periodic reports of its operations that generally cover a consecutive 12-month period of the provider’s operations. Amended cost reports to revise cost report information that has been previously submitted by a provider may be permitted or required as determined by HCFA.

(1) Cost reports—Terminated providers and changes of ownership. A provider that voluntarily or involuntarily ceases to participate in the Medicare program or experiences a change of ownership must file a cost report for that period under the program beginning with the first day not included in a previous cost reporting period and ending with the effective date of termination of its provider agreement or change of ownership.

(2) Due dates for cost reports. (i) Cost reports are due on or before the last day of the fifth month following the close of the period covered by the report. For cost reports ending on a day other than the last day of the month,
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Cost reports are due 150 days after the last day of the cost reporting period.

(ii) Extensions of the due date for filing a cost report may be granted by the intermediary only when a provider's operations are significantly adversely affected due to extraordinary circumstances over which the provider has no control, such as flood or fire.

(3) Changes in cost reporting periods. A provider may change its cost reporting period if a change in ownership is experienced or if the—

(i) Provider requests the change in writing from its intermediary;

(ii) Intermediary receives the request at least 120 days before the close of the new reporting period requested by the provider; and

(iii) Intermediary determines that good cause for the change exists. Good cause would not be found to exist if the effect is to change the initial date that a hospital would be affected by the rate increase ceiling (see § 413.40), or be paid under the prospective payment systems (see part 412 of this chapter).

(4) Electronic submission of cost reports.

(i) As used in this paragraph, "provider" means a hospital, skilled nursing facility, or home health agency.

(ii) Effective for cost reporting periods beginning on or after October 1, 1989, for hospitals, and cost reporting periods ending on or after February 1, 1994, for skilled nursing facilities and home health agencies, a provider must submit a hard copy of a settlement summary, a statement of certain worksheet totals found within the electronic file, and a statement signed by its administrator or chief financial officer certifying the accuracy of the electronic file or the manually prepared cost report. During a transition period, skilled nursing facilities and home health agencies must submit a hard copy of the completed cost report forms in addition to the electronic file. The following statement must immediately precede the dated signature of the provider’s administrator or chief financial officer:

I hereby certify that I have read the above certification statement and that I have examined the accompanying electronically filed or manually submitted cost report and the Balance Sheet Statement of Revenue and Expenses prepared by (Provider Name(s) and Number(s)) for the cost reporting period beginning ______ and ending ______ and that to the best of my knowledge and belief, this report and statement are true, correct, complete and prepared from the books and records of the provider in accordance with applicable instructions, except as noted. I further certify that I am familiar with the laws and regulations regarding the provision of health care services, and that the services identified in this cost report were provided in compliance with such laws and regulations.

(v) A provider may request a delay or waiver of the electronic submission requirement in paragraph (f)(4)(ii) of this section if this requirement would cause a financial hardship or if the provider...
§ 413.30 Limitations on payable costs.

(a) Introduction—(1) Scope. This section implements section 1861(v)(1)(A) of the Act by setting forth the general rules under which HCFA may establish limits on SNF and HHA costs recognized as reasonable in determining Medicare program payments. It also sets forth rules governing exemptions and exceptions to limits established under this section that HCFA may make as appropriate in considering special needs or situations of particular providers.

(2) General principle. Reimbursable provider costs may not exceed HCFA estimates to be necessary for the efficient delivery of needed health care services. HCFA may establish estimated cost limits for direct or indirect overall costs or for costs of specific services or groups of services. HCFA imposes these limits prospectively and may calculate them on a per

(h) Waiver of full or simplified cost reporting for low program utilization. (1) If the provider has had low utilization of covered services by Medicare beneficiaries (as determined by the intermediary) and has received correspondingly low interim payments for the cost reporting period, the intermediary may waive a full cost report or the simplified cost report described in §413.321 if it decides that it can determine, without a full or simplified report, the reasonable cost of covered services provided during that period.

(2) If a full or simplified cost report is waived, the provider must submit within the same time period required for full or simplified cost reports:

(i) The cost reporting forms prescribed by HCFA for this situation; and

(ii) Any other financial and statistical data the intermediary requires.


EFFECTIVE DATE NOTE: At 65 FR 18537, Apr. 7, 2000, §413.34 was amended by adding paragraph (d)(6), effective October 10, 2000.

Subpart C—Limits on Cost Reimbursement

§ 413.30 Limitations on payable costs.

(a) Introduction—(1) Scope. This section implements section 1861(v)(1)(A) of the Act by setting forth the general rules under which HCFA may establish limits on SNF and HHA costs recognized as reasonable in determining Medicare program payments. It also sets forth rules governing exemptions and exceptions to limits established under this section that HCFA may make as appropriate in considering special needs or situations of particular providers.

(2) General principle. Reimbursable provider costs may not exceed HCFA estimates to be necessary for the efficient delivery of needed health care services. HCFA may establish estimated cost limits for direct or indirect overall costs or for costs of specific services or groups of services. HCFA imposes these limits prospectively and may calculate them on a per
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admission, per discharge, per diem, per visit, or other basis.

(b) Procedure for establishing limits. (1) In establishing limits under this section, HCFA may classify SNFs and HHAs by factors that HCFA finds appropriate and practical, including the following:

(i) Type of services furnished.

(ii) Geographical area where services are furnished, allowing for grouping of noncontiguous areas having similar demographic and economic characteristics.

(iii) Size of institution.

(iv) Nature and mix of services furnished.

(v) Type and mix of patients treated.

(2) HCFA bases its estimates of the costs necessary for efficient delivery of health services on cost reports or other data providing indicators of current costs. HCFA adjusts current and past period data to arrive at estimated costs for the prospective periods to which limits are applied.

(3) Before the beginning of a cost period to which revised limits will be applied, HCFA publishes a notice in the Federal Register, establishing cost limits and explaining the basis on which they are calculated.

(4) In establishing limits under paragraph (b)(1) of this section, HCFA may find it inappropriate to apply particular limits to a class of SNFs or HHAs due to the characteristics of the SNF or HHA class, the data on which HCFA bases those limits, or the method by which HCFA determines the limits. In these cases, HCFA may exclude that class of SNFs or HHAs from the limits, explaining the basis of the exclusion in the notice setting forth the limits for the appropriate cost reporting periods.

(c) Requests regarding applicability of cost limits. For cost reporting periods beginning before July 1, 1998, a SNF may request an exception or exemption to the cost limits imposed under this section. An HHA may request only an exception to the cost limits. The SNF or HHA must make its request to its fiscal intermediary within 180 days of the date on the intermediary’s notice of program reimbursement.

(d) Home health agencies. The intermediary makes a recommendation on the HHA’s request to HCFA, which makes the decision. HCFA responds to the request within 180 days from the date HCFA receives the request from the intermediary. The intermediary notifies the HHA of HCFA’s decision. The time required by HCFA to review the request is considered good cause for the granting of an extension of the time limit for the HHA to apply for a PRRB review, as specified in §405.1841 of this chapter. HCFA’s decision is subject to review under subpart R of part 405 of this chapter.

(2) Skilled nursing facility exception. The intermediary makes the final determination on the SNF’s exception request and notifies the SNF of its determination within 90 days from the date that the intermediary receives the request from the SNF. If the intermediary determines that the SNF did not provide adequate documentation from which a proper determination can be made, the intermediary notifies the SNF that the request is denied. The intermediary also notifies the SNF that it has 45 days from the date on the intermediary’s denial letter to submit a new exception request with the complete documentation and that otherwise, the denial is the final determination. The time required by the intermediary to review the request is considered good cause for the granting of an extension of the time limit for the SNF to apply for a PRRB review, as specified in §405.1841 of this chapter. The intermediary’s determination is subject to review under subpart R of part 405 of this chapter.

(d) Exemptions. Exemptions from the limits imposed under this section may be granted to a new SNF with cost reporting periods beginning before July 1, 1998 as stated in §413.1(g)(1). The intermediary makes a recommendation on the provider’s request to HCFA, which makes the decision. A new SNF is a provider of inpatient services that has operated a SNF (or the equivalent) for which it is certified for Medicare, under present and previous ownership, for less than 3 full years. An exemption granted under this paragraph expires at the end of the SNF’s first cost reporting period beginning at least 2 years after the provider accepts its first inpatient.
(e) Exceptions. Limits established under this section may be adjusted upward for a SNF or HHA under the circumstances specified in paragraphs (e)(1) through (e)(5) of this section. An adjustment is made only to the extent that the costs are reasonable, attributable to the circumstances specified, separately identified by the SNF or HHA, and verified by the intermediary.

(1) Atypical services. The SNF or HHA can show that the—

(i) Actual cost of services furnished by a SNF or HHA exceeds the applicable limit because the services are atypical in nature and scope, compared to the services generally furnished by SNFs or HHAs similarly classified; and

(ii) Atypical services are furnished because of the special needs of the patients treated and are necessary in the efficient delivery of needed health care.

(2) Extraordinary circumstances. The SNF or HHA can show that it incurred higher costs due to extraordinary circumstances beyond its control. These circumstances include, but are not limited to, strikes, fire, earthquake, flood, or other unusual occurrences with substantial cost effects.

(3) Areas with fluctuating populations. The SNF meets the following conditions:

(i) Is located in an area (for example, a resort area) that has a population that varies significantly during the year.

(ii) Is furnishing services in an area for which the appropriate health planning agency has determined does not have a surplus of beds or similar services and has certified that the beds or similar services furnished by the SNF are necessary.

(iii) Meets occupancy or capacity standards established by the Secretary.

(4) Medical and paramedical education. The SNF or HHA can demonstrate that, if compared to other SNFs or HHAs in its group, it incurs increased costs for services covered by limits under this section because of its operation of an approved education program specified in §413.85.

(5) Unusual labor costs. The SNF or HHA has a percentage of labor costs that varies more than 10 percent from that included in the promulgation of the limits.

(f) Operational review. Any SNF or HHA that applies for an exception to the limits established under paragraph (e) of this section must agree to an operational review at the discretion of HCFA. The findings from this review may be the basis for recommendations for improvements in the efficiency and economy of the SNF's or the HHA's operations. If recommendations are made, any future exceptions are contingent on the SNF's or HHA's implementation of these recommendations.

[64 FR 42612, Aug. 5, 1999; 65 FR 60104, Oct. 10, 2000]
in excess of the costs determined to be necessary in the efficient delivery of needed health services under Medicare; and

(5) The provider has, in the manner described in paragraph (e) of this section, identified such charges to such individual or person acting on his behalf as charges to meet the costs in excess of the costs determined to be necessary in the efficient delivery of needed health services under Medicare.

(b) Provider request to charge beneficiaries for costs in excess of limits.

(1) If a provider’s actual costs (or, if less, the customary charges) in the second preceding cost period exceed the prospective limits established for such costs, the intermediary will, at the provider’s request, validate in advance the charges that may be made to the beneficiaries for the excess.

(2) If a provider does not have a second preceding cost period and is a new provider as defined in §413.30(e), the provider, subject to validation by the intermediary, will estimate the current cost of the service to which a limit is being applied. Such amount will be adjusted to an amount equivalent to costs in the second preceding year by use of a factor to be developed based on estimates of cost increases during the preceding two years and published by SSA or HCFA. The amount thus derived will be used in lieu of the second preceding cost period amount in determining the charge to the beneficiary.

(3) To obtain consideration of such a request, the provider must submit to the intermediary a statement indicating the charge for which it is seeking validation and providing the data and method used to determine the amount. Such statement should include the—

(i) Provider’s name and number;

(ii) Identity of class and prospective cost limit for the class in which the provider has been included;

(iii) Amount of charge and cost period in which the charge is to be imposed;

(iv) Cost and customary charge for items and services furnished to beneficiaries; and

(v) Cost period ending date of the second reporting period immediately preceding the cost period in which the charge is to be imposed. The intermediary may request such additional information as it finds necessary with respect to the request.

(c) Provider charges.—(1) Establishing the charges. If the actual cost incurred (or, if less, the customary charges) in the prior period determined under paragraph (a) of this section exceeds the limits applicable to the pertinent period, the provider may charge the beneficiary to the extent costs in the second preceding cost reporting period (or the equivalent when there is no second preceding period) exceed the current cost limits. (Data from the most recently submitted appropriate cost report will be used in determining the actual cost.) For example, if a limit of $58 per day is applied to the cost of general routine services for the provider’s cost reporting period starting in calendar year 1975 and if the provider’s actual general routine cost in the second preceding reporting period, that is, the reporting period starting in calendar year 1973, was $60 per day, the provider (after first having obtained intermediary validation and subject to the considerations and requirements specified in paragraph (a) of this section) may charge Medicare Part A beneficiaries up to $2 per day for general routine services.

(2) Adjusting cost. Program reimbursement for the costs to which limits imposed under §413.30 are applied in any cost reporting period will not exceed the lesser of the provider’s actual cost or the limits imposed under §413.30. If program reimbursement for items or services to which such limits are applied plus the charges to beneficiaries for such items or services imposed under this section exceed the provider’s actual cost for such items or services, program payment to the provider will be reduced to the extent program payment plus charges to the beneficiaries exceed actual cost. If the provider’s actual cost for general routine services in 1975 was $57,000, the cost limit was $58,000, and billed charges to Medicare Part A beneficiaries were $2,000, the provider would receive $55,000 from the program ($57,000 actual cost minus the $2,000 in charges to the beneficiaries).

(d) Definition of emergency services. For purposes of paragraph (a)(2) of this
section, emergency services are those hospital services that are necessary to prevent the death or serious impairment of the health of the individual, and which, because of the threat to the life or health of the individual, necessitate the use of the most accessible hospital (as determined under § 424.106 of this chapter) available and equipped to furnish such services. If an individual has been admitted to such hospital as an inpatient because of an emergency, the emergency will be deemed to continue until it is safe from a medical standpoint to move the individual to another hospital or other institution or to discharge him.

(e) Identification of charges to individual. For purposes of paragraph (a)(5) of this section, a provider must give or send to the individual or his representative, a schedule of all items and services that the individual might need and for which the provider imposes charges under this section, and the charge for each. Such schedule must specify that the charges are necessary to meet the costs in excess of the costs determined to be necessary in the efficient delivery of needed health services under Medicare and include such other information as HCFA considers necessary to protect the individual’s rights under this section. The provider, in arranging for the individual’s admission, first service, or start of care, must give or send this schedule to the individual or his representative when arrangements are being made for such services or if this is not feasible, as soon thereafter as is practicable but no later than at the initiation of services.


§ 413.40 Ceiling on the rate of increase in hospital inpatient costs.

(a) Introduction—(1) Scope. This section implements section 1886(b) of the Act, establishing a ceiling on the rate of increase in operating costs per case for hospital inpatient services furnished to Medicare beneficiaries that will be recognized as reasonable for purposes of determining the amount of Medicare payment. This rate-of-increase ceiling applies to hospital cost reporting periods beginning on or after October 1, 1982. This section also sets forth rules governing exemptions from and adjustments to the ceiling.

(2) Applicability. (i) This section is not applicable to—

(A) Hospitals reimbursed in accordance with section 1814(b)(3) of the Act or under State reimbursement control systems that have been approved under section 1886(c) of the Act and subpart C of part 403 of this chapter; or

(B) Hospitals that are paid under the prospective payment systems for inpatient hospital services in accordance with section 1886(d) and (g) of the Act and part 412 of this chapter.

(ii) For cost reporting periods beginning on or after October 1, 1983, this section applies to hospitals excluded from the prospective payment system in accordance with §412.23 of this chapter, and psychiatric and rehabilitation units excluded from the prospective payment system in accordance with §§412.25 through 412.30 of this chapter.

(3) Definitions. As used in this section—

Ceiling is the aggregate upper limit on the amount of a hospital’s net Medicare inpatient operating costs that the program will recognize for payment purposes. For each cost reporting period, the ceiling is determined by multiplying the updated target amount, as defined in this paragraph, for that period by the number of Medicare discharges during that period. For a hospital-within-a-hospital, as described in §412.22(e) of this chapter, the number of Medicare discharges in a cost reporting period does not include discharges of a patient to another hospital in the same building on or on the same campus, if—

(A) The patient is subsequently readmitted to the hospital-within-a-hospital directly from the other hospital; and

(B) The hospital-within-a-hospital has discharged to the other hospital and subsequently readmitted more than 5 percent (that is, in excess of 5.0 percent) of the total number of Medicare inpatients discharged from the hospital-within-a-hospital in that cost reporting period.

Date of discharge is the earliest of the following dates:
(A) The date the patient has exhausted Medicare Part A hospital inpatient benefits (including the election to use lifetime reserve days) during his or her spell of illness.

(B) The date the patient is formally released as specified in §412.4(a)(1) of this chapter.

(C) The date the patient is transferred to another facility.

(D) The date the patient dies.

Market basket index is HCFA’s projection of the annual percentage increase in hospital inpatient operating costs. The market basket index is a wage and price index that incorporates weighted indicators of changes in wages and prices that are representative of the mix of goods and services included in the most common categories of hospital inpatient operating costs subject to the ceiling, as described in paragraph (c)(1) of this section.

Net inpatient operating costs include the costs of certain preadmission services as specified in §413.40(c)(2), the costs of routine services, ancillary services, and intensive care services (as defined in §413.53(b)) incurred by a hospital in furnishing covered inpatient services to Medicare beneficiaries. Net inpatient operating costs exclude capital-related costs as described in §413.130, the costs of approved medical education programs as described in §§413.85 and 413.86, heart, kidney, and liver acquisition costs incurred by approved transplantation centers. These costs are identified and excluded from inpatient operating costs before the application of the ceiling.

Rate-of-increase percentage is the percentage by which each hospital’s target amount from the preceding Federal fiscal year is increased.

Target amount is the per discharge (case) limitation, derived from the hospital’s allowable net Medicare inpatient operating costs in the hospital’s base year, and updated for each subsequent hospital cost reporting period by the appropriate annual rate-of-increase percentage.

Update adjustment percentage is the percentage by which a hospital’s allowable inpatient operating service costs for the 12-month cost reporting period beginning in Federal fiscal year 1990 exceeds the hospital’s ceiling for that period.

Update factor is the decimal equivalent of the rate-of-increase percentage. The update factor is the value by which a hospital’s target amount for the preceding year is multiplied in order to determine the target amount for the following year. For example, if the rate-of-increase percentage for a year is 2.7 percent, the update factor for that year is 1.027.

(b) Cost reporting periods subject to the rate-of-increase ceiling.

(i) The target amount established under this provision remains applicable to a hospital or excluded hospital unit, as described in §§412.25 through 412.30 of this chapter, despite intervening cost reporting periods during which the hospital or excluded hospital unit is not subject to the ceiling as a result of other provisions of the law or regulations, or nonparticipation in the Medicare program, unless the hospital or excluded hospital unit qualifies as a new hospital or excluded part hospital unit under the provisions of paragraph (f) of this section.

(ii) The base period for a newly established excluded unit is the first cost reporting period of at least 12 months following the unit’s certification to participate in the Medicare program.

(iii) When the operational structure of a hospital or unit changes (that is, a freestanding hospital becomes an excluded unit or an excluded unit becomes a freestanding hospital, or an entity of a multicampus hospital becomes a newly created hospital or unit or a hospital or unit becomes a part of a multicampus hospital), the base period for the hospital or unit that changed its operational structure is the first cost reporting period of at
least 12 months effective with the revised Medicare certification classification.

(iv) Request for rebased target amount for the cost reporting period beginning on or after October 1, 1997 and on or before September 30, 1998. Except for qualified long-term care hospitals as defined in paragraph (b)(1)(v) of this section, each hospital or unit under present or previous ownership that received payment under section 1886(b) of the Act during cost reporting periods beginning before October 1, 1990, may submit a request to its fiscal intermediary to rebase its target amount. The request must be received by the fiscal intermediary by the later of November 1, 1997 or 60 days before the beginning of its cost reporting period beginning during fiscal year 1998. The rebased target amount for the cost reporting period beginning during fiscal year 1998 is determined as follows:

(A) Determine the hospital’s inpatient operating costs per case for each of the five most recent settled cost reports as of August 5, 1997.

(B) For each of the five cost reports, update the operating costs per case by the applicable update factors up to the hospital’s cost reporting period beginning during FY 1998.

(C) Exclude the highest and lowest of the five updated amounts determined under paragraph (b)(1)(iv)(B) of this section.

(D) Compute the average for the remaining three updated amounts for operating cost per case.

(v) Request by qualified long-term care hospital. A qualified long-term care hospital may file a request to its fiscal intermediary for a rebased FY 1998 target amount. The request must be received by the fiscal intermediary by the later of November 1, 1997 or 60 days before the beginning of its cost reporting period beginning during fiscal year 1998. The rebased FY 1998 target amount is the hospital’s FY 1996 inpatient operating costs updated to FY 1997. A qualified long-term care hospital means a long-term care hospital that meets the following two conditions for its two most recent settled cost reports as of August 5, 1997:

(A) Its Medicare inpatient operating costs exceed 115 percent of the ceiling.

(B) The hospital would have had a disproportionate patient percentage (as defined in §412.106) equal to or greater than 70 percent if it were a prospective payment hospital.

(2) Periods subject to the ceiling. The ceiling established under this section applies to all cost reporting periods that—

(i) Begin on or after October 1, 1982; and

(ii) Immediately follow the base period established under paragraph (b)(1) of this section unless the exception in paragraph (b)(3) of this section is applicable.

(3) Periods of other than 12 months. The ceiling established under this section does not apply to cost reporting periods of fewer than 12 months that occur in conjunction with a change in operation of the facility, as defined in paragraph (b)(1)(iii) of this section, as a result of changes in ownership, merger, or consolidation. However, the ceiling applies to cost reporting periods of fewer than 12 months that result solely from the approval of a hospital’s request for a change in accounting cycle, as specified in §413.24(f)(3).

(c) Costs subject to the ceiling—(1) Applicability. The ceiling established under this section applies to net operating costs incurred by a hospital in furnishing inpatient hospital services to Medicare beneficiaries.

(2) Preadmission services otherwise payable under Medicare Part B furnished to a beneficiary during the calendar day immediately preceding the date of the beneficiary’s admission to the hospital that meet the following conditions:

(i) The services are furnished by the hospital or any entity wholly owned or operated by the hospital. An entity is wholly owned by the hospital if the hospital is the sole owner of the entity. An entity is wholly operated by a hospital if the hospital has exclusive responsibility for conducting and overseeing the entity’s routine operations, regardless of whether the hospital also has policymaking authority over the entity.

(ii) For services furnished after January 1, 1991, the services are diagnostic (including clinical diagnostic laboratory tests).
(iii) For services furnished on or after October 1, 1991, the services are furnished in connection with the principal diagnosis that requires the beneficiary to be admitted as an inpatient and are not the following:

(A) Ambulance services.

(B) Maintenance renal dialysis.

(3) Rate-of-increase percentages and update factors. The applicable rate-of-increase percentages and update factors are determined as follows:

(i) Federal fiscal year 1986. The applicable rate-of-increase percentage for cost reporting periods beginning on or after October 1, 1985 and before September 30, 1986 is five twenty-fourths of one percent, and the update factor is 1.00208333. For purposes of determining the target amount for cost reporting periods beginning on or after October 1, 1986, the applicable percentage increase for cost reporting periods beginning during Federal fiscal year 1986 is deemed to have been one-half percent, and the update factor is 1.005.

(ii) Federal fiscal year 1987. The applicable rate-of-increase percentage for cost reporting periods beginning on or after October 1, 1986 and before September 30, 1987 is 1.15 percent; the update factor is 1.0115.

(iii) Federal fiscal year 1988. The applicable rate-of-increase percentage for cost reporting periods beginning on or after October 1, 1987 and before October 1, 1988 is 2.3238 percent; the update factor is 1.023238. For purposes of updating the target amount for cost reporting periods beginning during FY 1988, the rate-of-increase percentage for cost reporting periods beginning during FY 1988 is deemed to have been 2.7 percent; the update factor is deemed to have been 1.027.

(iv) Federal fiscal year 1989 through Federal fiscal year 1993. The applicable rate-of-increase percentage for cost reporting periods beginning on or after October 1, 1988, and before October 1, 1993, is the percentage increase projected by the hospital market basket index (as defined in paragraph (a)(3) of this section).

(v) Federal fiscal year 1994 through Federal fiscal year 1997. The applicable rate-of-increase percentage for cost reporting periods beginning on or after October 1, 1993, and before October 1, 1998, is the market basket percentage increase minus the lesser of, 1 percentage point, or the percentage point difference between 10 percent and the hospital’s “update adjustment percentage” (as defined in paragraph (a)(3) of this section); for hospitals with an “update adjustment percentage” of at least 10 percent, the applicable rate-of-increase percentage is the market basket percentage increase. The “update adjustment percentage” is increased in each Federal fiscal year by the sum of the hospital’s applicable reductions applied to the market basket percentage increase for previous Federal fiscal years.

(vi) Federal fiscal year 1998. The applicable rate-of-increase percentage for cost reporting periods beginning on or after October 1, 1997 is 0 percent.

(vii) Federal fiscal year 1999 through Federal fiscal year 2002. The applicable rate-of-increase percentage for cost reporting periods beginning on or after October 1, 1998, and before October 1, 2002, based on data from the most recent available cost report, is:

(A) The percentage increase in the market basket, if inpatient operating costs are equal to or exceed the ceiling amount by 10 percent or more of the ceiling.

(B) The percentage increase in the market basket minus .25 percentage points for each percentage point by which inpatient operating costs are less than 10 percent over the ceiling (but not less than 0), if inpatient operating costs exceed the ceiling by less than 10 percent of the ceiling.

(C) The greater of the percentage increase in the market basket minus 2.5 percentage points or 0 percent, if inpatient operating costs are equal to or less than the ceiling but greater than 66.7 percent of the ceiling.

(D) 0 percent, if inpatient operating costs do not exceed 66.7 percent of the ceiling.

(viii) Federal fiscal year 2003 and following. The applicable rate-of-increase percentage for cost reporting periods beginning on or after October 1, 2002, is the percentage increase projected by the hospital market basket index.

(4) Target amounts. The intermediary will establish a target amount for each hospital. The target amount for a cost

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reporting period is determined as follows:

(i) Except as provided in paragraph (c)(4)(iv) of this section, and subject to the provisions of paragraph (c)(4)(iii) of this section, for the first cost reporting period to which this ceiling applies, the target amount equals the hospital’s allowable net inpatient operating costs per case for the hospital’s base period increased by the update factor for the subject period.

(ii) Subject to the provisions of paragraph (c)(4)(iii) of this section, for subsequent cost reporting periods, the target amount equals the hospital’s target amount for the previous cost reporting period increased by the update factor for the subject cost reporting period, unless the provisions of paragraph (c)(5)(ii) of this section apply.

(iii) In the case of a psychiatric hospital or unit, rehabilitation hospital or unit, or long-term care hospital, the target amount is the lower of—

(A) The hospital-specific target amount (the net allowable costs in a base period increased by the applicable update factors); or

(B) One of the following for the applicable cost reporting period—

(1) For cost reporting periods beginning during fiscal year 1996, the 75th percentile of target amounts for hospitals in the same class (psychiatric hospital or unit, rehabilitation hospital or unit, or long-term care hospital) for cost reporting periods ending during FY 1996, increased by the applicable market basket percentage up to the first cost reporting period beginning on or after October 1, 1996.

(2) For cost reporting periods beginning during fiscal year 1996, the amount determined under paragraph (c)(4)(iii)(B)(1) of this section, increased by the market basket percentage up through the subject period, subject to the provisions of paragraph (c)(4)(iv) of this section.

(3) For cost reporting periods beginning during fiscal year 2000—

(i) The labor-related portion and the nonlabor-related portion of the wage-neutralized 75th percentile target amounts for hospitals in the same class (psychiatric hospital or unit, rehabilitation hospital or unit, or long-term care hospital) for cost reporting periods ending during FY 1999, are increased by the applicable market basket percentage up through the subject period.

(ii) The labor-related portion of the wage-neutralized 75th percentile target amounts for hospitals in the same class is determined by adding the nonlabor-related portion of the wage-neutralized 75th percentile target amounts determined under paragraph (c)(4)(iii)(B)(3)(i) of this section and the hospital’s wage-adjusted labor-related portion of the wage-neutralized 75th percentile target amounts determined under paragraph (c)(4)(iii)(B)(3)(ii) of this section, subject to the provisions of paragraph (c)(4)(iv) of this section.

(4) For cost reporting periods beginning during fiscal years 2001 and 2002—

(i) The amounts determined under paragraph (c)(4)(iii)(B)(3)(i) of this section are increased by the applicable market basket percentage up through the subject period.

(ii) The labor-related portion of the wage-neutralized 75th percentile target amounts, for cost reporting periods beginning during fiscal year 2001 and the hospital’s hospital inpatient prospective payment system wage index, for cost reporting periods beginning during fiscal year 2002.

(iii) The wage-adjusted 75th percentile target amounts for hospitals in the same class are determined by adding the nonlabor-related portion of the wage-neutralized 75th percentile target amounts determined under paragraph (c)(4)(iii)(B)(4)(i) of this section, subject to the provisions of paragraph (c)(4)(iv) of this section.

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(iv) For purposes of the limits on target amounts established under paragraph (c)(4)(iii) of this section, each hospital or unit that qualifies for exclusion as a member of only one class of excluded facility (psychiatric hospital or unit, rehabilitation hospital or unit, or long-term care hospital) will be subject to the limit applicable to that class. If a hospital or unit qualifies to be classified in more than one way under the exclusion criteria in subpart B of part 412 of this chapter, the hospital’s or unit’s target amount may not exceed the lowest applicable limit.

(v) In the case of a hospital that received payments under paragraph (f)(2)(ii) of this section as a newly created hospital or unit, to determine the hospital’s target amount for the hospital’s third 12-month cost reporting period, the payment amount determined under paragraph (f)(2)(ii)(A) of this section for the preceding cost reporting period is updated to the third cost reporting period.

(5) Applicable update factor. (i) The applicable update factor is derived from the prospectively determined rate-of-increase percentage published by HCFA. The update factor for each Federal fiscal year is applied prospectively to the target amount for each cost reporting period beginning during the Federal fiscal year.

(ii) In the case of cost reporting periods of less than 12 months, the target amount determined for a hospital’s first cost reporting period beginning in a Federal fiscal year applies to subsequent periods beginning in the same Federal fiscal year.

(d) Application of the target amount in determining the amount of payment. (1) General process. (i) At the end of each cost reporting period subject to this section, the hospital’s intermediary will compare a hospital’s allowable net inpatient operating costs with that hospital’s ceiling (as defined in paragraph (a)(3) of this section) for that period.

(ii) The hospital’s actual allowable costs will be determined without regard to the lesser of cost or charges provisions of §413.13, and in accordance with the provisions of paragraphs (d)(2) or (d)(3) of this section, as applicable.

(2) Net inpatient operating costs are less than or equal to the ceiling. For cost reporting periods beginning on or after October 1, 1997, if a hospital’s allowable net inpatient operating costs do not exceed the hospital’s ceiling, payment to the hospital will be determined on the basis of the lower of the—

(i) Net inpatient operating costs plus 15 percent of the difference between inpatient operating costs and the ceiling; or

(ii) Net inpatient operating costs plus 2 percent of the ceiling.

(3) Net inpatient operating costs are greater than the ceiling. For cost reporting periods beginning on or after October 1, 1997—

(i) If a hospital’s allowable net inpatient operating costs do not exceed 110 percent of the ceiling (or the adjusted ceiling, if applicable), payment will be the ceiling (or the adjusted ceiling, if applicable);

(ii) If a hospital’s allowable net inpatient operating costs are greater than 110 percent of the ceiling (or the adjusted ceiling, if applicable), payment will be the ceiling (or the adjusted ceiling, if applicable) plus the lesser of:

(A) 50 percent of the allowable net inpatient operating costs in excess of 110 percent of the ceiling (or the adjusted ceiling, if applicable); or

(B) 10 percent of the ceiling (or the adjusted ceiling, if applicable).

(4) Continuous improvement bonus payments. (i) For cost reporting periods beginning on or after October 1, 1997 and ending before October 1, 2000, eligible hospitals (as defined in paragraph (d)(5) of this section) receive payments in addition to those in paragraph (d)(2) of this section, as applicable. These payments are equal to the lesser of—

(A) 50 percent of the amount by which the operating costs are less than the expected costs for the period; or

(B) 1 percent of the ceiling.

(ii) For cost reporting periods beginning on or after October 1, 2000, and ending before September 30, 2001, eligible psychiatric hospitals and units and long-term care hospitals (as defined in paragraph (d)(5) of this section) receive payments in addition to those in paragraph (d)(2) of this section, as applicable. These payments are equal to the lesser of—

(A) 50 percent of the amount by which the operating costs are less than the expected costs for the period; or

(B) 1 percent of the ceiling.
(A) 50 percent of the amount by which the operating costs are less than the expected costs for the period; or
(B) 1.5 percent of the ceiling.

(iii) For cost reporting periods beginning on or after October 1, 2001, and before September 30, 2002, eligible psychiatric hospitals and units and long-term care hospitals receive payments in addition to those in paragraph (d)(5) of this section, as applicable. These payments are equal to the lesser of—
(A) 50 percent of the amount by which the operating costs are less than the expected costs for the periods; or
(B) 2 percent of the ceiling.

(5) Eligibility requirements for continuous improvement bonus payments. To qualify, a hospital must have been paid as a prospective payment excluded hospital for at least three full cost reporting periods prior to the applicable period, and the hospital's operating costs per discharge for the period must be less than the least of the following:
(i) The hospital's target amount.
(ii) The hospital's trended costs.
(A) For a hospital for which its cost reporting period ending during fiscal year 1996 was its third or subsequent full cost reporting period, trended costs are the lesser of the allowable inpatient operating costs per discharge or the target amount for the cost reporting period ending in fiscal year 1996, increased in a compounded manner for each succeeding fiscal year by the market basket percentage increase;
(B) For all other hospitals, trended costs are the allowable inpatient operating costs per discharge for its third full cost reporting period increased in a compounded manner for each succeeding fiscal year by the market basket increase.
(iii) The hospital's expected costs. The hospital's expected costs are the lesser of its allowable inpatient operating costs per discharge or the target amount for the previous cost reporting period, updated by the market basket percentage increase for the fiscal year.

(6) Hospital requests regarding adjustments to the payment allowed under the rate-of-increase ceiling. (1) Timing of application. A hospital may request an adjustment to the rate-of-increase ceiling imposed under this section. The hospital's request must be received by the hospital's fiscal intermediary no later than 180 days after the date on the intermediary's initial notice of amount of program reimbursement (NPR) for the cost reporting period for which the hospital requests an adjustment.

(2) Intermediary recommendation. Unless HCFA has authorized the intermediary to make the decision, the intermediary makes a recommendation on the hospital's request to HCFA, which makes the decision. HCFA issues a decision to the intermediary no later than 180 days after receipt of the completed application and the intermediary's recommendation.

(3) Intermediary decision. If HCFA has authorized the intermediary to make the decision, the intermediary issues a decision no later than 180 days after receipt of the completed application.

(4) Notification and review. (i) The intermediary notifies the hospital of the decision, including a full explanation of the grounds for the decision. A decision issued under paragraph (e)(2) or (e)(3) of this section is considered final unless the hospital submits additional information and requests a review of the decision no later than 180 days after the date on the intermediary's notice of the decision.

(ii) The final decision is subject to review under the provider reimbursement determination and appeal procedures in subpart R of part 405 of this chapter, provided the hospital has received an NPR for the cost reporting period in question, and the NPR disallows costs for which the hospital had requested an adjustment (see the definitions in §405.1801(a) of this chapter and the provisions regarding a provider's right to a Board hearing in §405.1835 of this chapter).

(5) Extending time limit for PRRB review of NPR. The time required to review the request is considered good cause for the granting of an extension of the time limit to apply for review of the notice of amount of program reimbursement by the Provider Reimbursement Review Board, as specified in §405.1841(b) of this chapter.

(6) Applicability. The provisions in paragraphs (e)(1) through (e)(5) of this
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section apply to a hospital’s initial request for an adjustment and to a request for a review of the original decision based on additional data.

(f) Comparison to the target amount for new hospitals and units—(1) New hospitals and units—(i) New hospitals. For purposes of this section, a new hospital is a provider of hospital inpatient services that—

(A) Has operated as the type of hospital for which HCFA granted it approval to participate in the Medicare program, under present or previous ownership (or both), for less than 2 full years; and

(B) Has provided the type of hospital inpatient services for which HCFA granted it approval to participate in the Medicare program, for less than 2 years.

(ii) New units. A newly established unit that is excluded from the prospective payments system under the provisions of §§412.25 through 412.30 of this chapter does not qualify for the exemption afforded to a new hospital under paragraph (f)(1)(i) of this section unless the unit is located in an acute care hospital that, if it were subject to the provisions of this section, would qualify as a new hospital under paragraph (f)(1)(i) of this section.

(2) Comparison—(i) Exemptions. (A) A new children’s hospital is exempt from the rate-of-increase ceiling imposed under this section. The exemption begins when the hospital accepts its first patient and ends at the end of the first cost reporting period ending at least 2 years after the hospital accepts its first patient. The first cost reporting period of at least 12 months beginning at least 1 year after the hospital accepts its first patient is the base year, in accordance with paragraph (b) of this section.

(B) Within 180 days of the date a hospital is excluded from the prospective payment system, the intermediary determines whether the hospital is exempt from the rate-of-increase ceiling. The intermediary notifies the hospital of its determination and the hospital’s base period.

(C) A decision issued under paragraph (f)(2)(ii)(B) of this section is considered final unless the hospital submits additional information and requests a review of the decision no later than 180 days after the date on the intermediary’s notice of the decision. The final decision is subject to review under subpart R of part 405 of this chapter, provided the hospital has received a notice of program reimbursement (NPR) for the cost reporting period in question and the NPR does not reflect an exemption (see the definitions in §405.1801(a) of this chapter and the provisions regarding a provider’s right to a Board hearing in §405.1835 of this chapter).

(ii) Median target amount. (A) For cost reporting periods beginning on or after October 1, 1997, the amount of payment for a new psychiatric hospital or unit, a new rehabilitation hospital or unit, or a new long-term care hospital that was not paid as an excluded hospital prior to October 1, 1997, is the lower of the hospital’s net inpatient operating cost per case or 110 percent of the national median of the target amounts for the class of excluded hospitals and units (psychiatric, rehabilitation, long-term care) as adjusted for differences in wage levels and updated to the first cost reporting period in which the hospital receives payment. The second cost reporting period is subject to the same target amount as the first cost reporting period.

(B) The national median of the target amounts is the FY 1996 median target amount—

(1) Adjusted to account for differences in area wage levels;

(2) Updated by the market basket percentage increase to the fiscal year in which the hospital first received payments as an excluded provider.

(3) Risk-basis HMOs. Items or services that are furnished to beneficiaries enrolled in an HMO by a hospital that is either owned or operated by a risk-basis HMO or related to a risk-basis HMO by common ownership or control are exempt from the rate-of-increase ceiling (see the definition of an entity with a risk sharing contract in §417.401 of this chapter).

(g) Adjustments—(1) General rules. (i) HCFA adjusts the amount of the operating costs considered in establishing the rate-of-increase ceiling for one or more cost reporting periods, including both periods subject to the ceiling and
the hospital’s base period, under the circumstances specified in paragraphs (g)(2), (g)(3), and (g)(4) of this section.

(ii) When the hospital requests an adjustment, HCFA makes an adjustment only to the extent that the hospital’s operating costs are reasonable, attributable to the circumstances specified separately, identified by the hospital, and verified by the intermediary.

(iii) When the hospital requests an adjustment, HCFA makes an adjustment only if the hospital’s operating costs exceed the rate-of-increase ceiling imposed under this section.

(iv) In the case of a psychiatric hospital or unit, rehabilitation hospital or unit, or long-term care hospital, the amount of payment under paragraph (g)(3) of this section may not exceed the payment amount based on the target amount determined under paragraph (c)(4)(iii) of this section.

(v) In the case of a hospital or unit that received a revised FY 1998 target amount under the rebasing provisions of paragraph (b)(1)(iv) of this section, the amount of an adjustment payment for a cost reporting period is based on a comparison of the hospital’s operating costs for the cost reporting period to the average costs and statistics for the cost reporting periods used to determine the FY 1998 rebased target amount.

(2) Extraordinary circumstances. HCFA may make an adjustment to take into account unusual costs (in either a cost reporting period subject to the ceiling or the hospital’s base period) due to extraordinary circumstances beyond the hospital’s control. These circumstances include, but are not limited to, strikes, fire, earthquakes, floods, or similar unusual occurrences with substantial cost effects.

(3) Comparability of cost reporting periods—(i) Adjustment for distortion. HCFA may make an adjustment to take into account factors that would result in a significant distortion in the operating costs of inpatient hospital services between the base year and the cost reporting period subject to the limits.

(ii) Factors. The adjustments described in paragraph (g)(3)(i) of this section, include, but are not limited to, adjustments to take into account:

(A) FICA taxes (if the hospital did not incur costs for FICA taxes in its base period).

(B) Services billed under part B of Medicare during the base period, but paid under part A during the subject cost reporting period.

(C) Malpractice insurance costs (if malpractice costs were not included in the base year operating costs).

(D) Increases in service intensity or length of stay attributable to changes in the type of patient served.

(E) A change in the inpatient hospital services that a hospital provides, and that are customarily provided directly by similar hospitals, such as an addition or discontinuation of services or treatment programs.

(F) The manipulation of discharges to increase reimbursement.

(iii) Adjusting operating costs. Without a formal request from a hospital, HCFA may adjust the amount of operating costs determined under paragraph (c)(1) of this section to take into account certain adjustments. These adjustments include, but are not limited to, adjustments under paragraphs (g)(3)(i)(A), (B), (C), (E), and (F) of this section.

(4) Significant wage increase. (i) Criteria. HCFA may make an adjustment to take into account a significant increase in wages occurring between the base period and the cost reporting period subject to the ceiling if there is a significant increase in the average hourly wage for the geographic area in which the hospital is located (determined by reference to the wage index for prospective payment hospitals without regard to geographic reclassifications under sections 1886(d)(8) and (10) of the Act). For this purpose, there is a significant wage increase if the wage index value based on wage survey data collected for the cost reporting period subject to the ceiling is at least 8.0 percent higher than the wage index value based on survey data collected for the base year cost reporting period. If survey data are not available for the cost reporting periods used in the comparison, the wage index value based on the latest available survey data collected prior to that cost reporting period is used.
(ii) Amount of the adjustment. The adjustment for a significant wage increase equals the amount by which the lesser of the following calculations exceeds 108 percent of the increase in the national average hourly earnings for hospital workers:

(A) The rate of increase in the average hourly wage in the geographic area (determined by applying the applicable increase in the area wage index value to the rate of increase in the national average hourly earnings for hospital workers).

(B) The rate of increase in the hospital’s average hourly wage.

(5) Adjustment limitations. For cost reporting periods beginning on or after October 1, 1993, and before October 1, 2003, the payment reductions under paragraph (c)(3)(v) through (c)(3)(vii) of this section will not be considered when determining adjustments under this paragraph.

(h) [Reserved]

(i) Assignment of a new base period. (1) General rule. Effective with cost reporting periods beginning on or after April 1, 1990, HCFA may assign a new base period to establish a revised ceiling if the new base period is more representative of the reasonable and necessary cost of furnishing inpatient services and all the following conditions apply:

(A) The actual allowable inpatient costs of the hospital in the cost reporting period that would be affected by the revised ceiling exceed the target amount established under paragraph (c) of this section.

(B) The hospital documents that the higher costs are the result of substantial and permanent changes in furnishing patient care services since the base period. In making this determination, HCFA takes into consideration the following factors:

(1) Changes in the services provided by the hospital.

(2) Changes in applicable technologies and medical practices.

(3) Differences in the severity of illness among patients or types of patients served.

(C) The adjustments described in paragraph (g) of this section would not result in recognition of the reasonable and necessary costs of providing inpatient services.

(ii) The revised ceiling is based on the necessary and proper costs incurred during the new base period.

(A) Increases in overhead costs (for example, administrative and general costs and housekeeping costs) are not taken into consideration unless the hospital documents that these increases result from substantial and permanent changes in furnishing patient care services.

(B) In determining whether wage increases are necessary and proper, HCFA takes into consideration whether increases in wages and wage-related costs for hospitals in the labor market area exceed the national average increase.

(2) New base period. The new base period is the first cost reporting period that is 12 months or longer that reflects the substantial and permanent change.

(3) New applicable rate-of-increase percentages and update factors. The revised target amount resulting from the assignment of a new base period is increased by the applicable rate-of-increase percentages (update factors) described in paragraph (c)(3) of this section.

(j) Reduction to capital-related costs. For psychiatric hospital and units, rehabilitation hospitals and units, and long-term care hospitals, the amount otherwise payable for capital-related costs for hospital inpatient services is reduced by 15 percent for portions of cost reporting periods occurring on or after October 1, 1997 through September 30, 2002.


Subpart D—Apportionment

§413.50 Apportionment of allowable costs.

(a) Consistent with prevailing practice in which third-party organizations
(b) In the study and consideration devoted to the method of apportioning costs, the objective has been to adopt methods for use under Medicare that would, to the extent reasonably possible, result in the program’s share of a provider’s total allowable costs being the same as the program’s share of the provider’s total services. This result is essential for carrying out the statutory directive that the program’s payments to providers should be such that the costs of covered services for beneficiaries would not be passed on to nonbeneficiaries, nor would the cost of services for nonbeneficiaries be borne by the program.

(c) A basic factor bearing upon apportionment of costs is that Medicare beneficiaries are not a cross section of the total population. Nor will they constitute a cross section of all patients receiving services from most of the providers that participate in the program. Available evidence shows that the use of services by persons age 65 and over differs significantly from other groups. Consequently, the objective sought in the determination of the Medicare share of a provider’s total costs means that the methods used for apportionment must take into account the differences in the amount of services received by patients who are beneficiaries and other patients serviced by the provider.

(d) The method of cost reimbursement most widely used at the present time by third-party purchasers of inpatient hospital care apportions a provider’s total costs among groups served on the basis of the relative number of days of care used. This method, commonly referred to as average-per-diem cost, does not take into account, variations in the amount of service which a day of care may represent and thereby assumes that the patients for whom payment is made on this basis are average in their use of service.

(e) In considering the average-per-diem method of apportioning cost for use under the program, the difficulty encountered is that the preponderance of presently available evidence strongly indicates that the over-age 65 patient is not typical from the standpoint of average-per-diem cost. On the average this patient stays in the hospital twice as long and therefore the ancillary services that he uses are averaged over the longer period of time, resulting in an average-per-diem cost for the aged alone, significantly below the average-per-diem for all patients.

(f) Moreover, the relative use of services by aged patients as compared to other patients differs significantly among institutions. Consequently, considerations of equity among institutions are involved as well as that of effectiveness of the apportionment method under the program in accomplishing the objective of paying each provider fully, but only for services to beneficiaries.

(g) A further consideration of long-range importance is that the relative use of services by aged and other patients can be expected to change, possibly to a significant extent in future years. The ability of apportionment methods used under the program to reflect such change is an element of flexibility which has been regarded as important in the formulation of the cost reimbursement principles.

(h) An alternative to the relative number of days of care as a basis for apportioning costs is the relative amount of charges billed by the provider for services to patients. The amount of charges is the basis upon which the cost of hospital care is distributed among patients who pay directly for the services they receive. Payment for services on the basis of charges applies generally under insurance programs in which individuals are indemnified for incurred expenses, a form of health insurance widely held throughout the United States. Also, charges to patients are commonly a factor in determining the amount of...
payment to hospitals under insurance programs providing service benefits, many of which pay "costs or charges, whichever is less" and some of which pay exclusively on the basis of charges. In all of these instances, the provider’s own charge structure and method of itemizing services for the purpose of assessing charges is utilized as a measure of the amount of services received and as the basis for allocating responsibility for payment among those receiving the provider's services.

(i) An increasing number of third-party purchasers who pay for services on the basis of cost are developing methods that utilize charges to measure the amount of services for which they have responsibility for payment. In this approach, the amount of charges for such services as a proportion of the provider's total charges to all patients is used to determine the proportion of the provider's total costs for which the third-party purchaser assumes responsibility. The approach is subject to numerous variations. It can be applied to the total of charges for all services combined or it can be applied to components of the provider’s activities for which the amount of costs and charges are ascertained through a breakdown of data from the provider’s accounting records.

(j) For the application of the approach to components, which represent types of services, the breakdown of total costs is accomplished by "cost-finding" techniques under which indirect costs and nonrevenue activities are allocated to revenue producing components for which charges are made as services are furnished.

§413.53 Determination of cost of services to beneficiaries.

(a) Principle. Total allowable costs of a provider will be apportioned between program beneficiaries and other patients so that the share borne by the program is based upon actual services received by program beneficiaries. The methods of apportionment are defined as follows:

(1) Departmental method—(i) Methodology. Except as provided in paragraph (a)(1)(ii) of this section with respect to the treatment of the private room cost differential for cost reporting periods starting on or after October 1, 1982, the ratio of beneficiary charges to total patient charges for the services of each ancillary department is applied to the cost of the department; to this is added the cost of routine services for program beneficiaries, determined on the basis of a separate average cost per diem for general routine patient care areas as defined in paragraph (b) of this section, taking into account, in hospitals, a separate average cost per diem for each intensive care unit, coronary care unit, and other intensive care type inpatient hospital units.

(ii) Exception: Indirect cost of private rooms. For cost reporting periods starting on or after October 1, 1982, except with respect to a hospital receiving payment under part 412 of this chapter (relating to the prospective payment system), the additional cost of furnishing services in private room accommodations is apportioned to Medicare only if these accommodations are furnished to Medicare patients and are medically necessary. To determine routine service cost applicable to beneficiaries—

(A) Multiply the average cost per diem (as defined in paragraph (b) of this section) by the total number of Medicare patient days (including private room days whether or not medically necessary);

(B) Add the product of the average per diem private room cost differential (as defined in paragraph (b) of this section) and the number of medically necessary private room days used by beneficiaries; and

(C) Effective October 1, 1990, do not include private rooms furnished for SNF-type and NF-type services under the swing-bed provision in the number of days in paragraphs (a)(1)(ii)(A) and (B) of this section.

(2) Carve-out out method—(i) The carve-out out method is used to allocate hospital inpatient general routine service costs in a participating swing-bed hospital, as defined in §413.114(b). Under this method, effective for services furnished on or after October 1, 1990, the reasonable costs attributable to the inpatient routine SNF-type and NF-type services furnished to all classes of patients are subtracted from total inpatient routine service costs before
computing the average cost per diem for inpatient routine hospital care.

(ii) The cost per diem attributable to the routine SNF-type services covered by Medicare is based on the regional Medicare swing-bed SNF rate in effect for a given calendar year, as described in §413.114(c). The Medicare SNF rate applies only to days covered and paid as Medicare days. When Medicare coverage runs out, the Medicare rate no longer applies.

(iii) The cost per diem attributable to all non-Medicare swing-bed days is based on the average statewide Medicaid NF rate for the prior calendar year, adjusted to approximate the average NF rate for the current calendar year.

(iv) The sum of total Medicare SNF-type days multiplied by the cost per diem attributable to Medicare SNF-type services and the total NF-type days multiplied by the cost per diem attributable to all non-Medicare days is subtracted from total inpatient general routine service costs. The cost per diem for inpatient routine hospital care is computed based on the remaining inpatient routine service costs.

(3) Cost per visit by type-of-service method—HHAs.

For cost reporting periods beginning on or after October 1, 1980, all HHAs must use the cost per visit by type-of-service method of apportioning costs between Medicare and non-Medicare beneficiaries. Under this method, the total allowable cost of all visits for each type of service is divided by the total number of visits for that type of service. Next, for each type of service, the number of Medicare covered visits is multiplied by the average cost per visit just computed. This represents the cost Medicare will recognize as the cost for that service, subject to cost limits published by HCFA (see §413.30).

(b) Definitions. As used in this section—

Ancillary services means the services for which charges are customarily made in addition to routine services.

Apportionment means an allocation or distribution of allowable cost between the beneficiaries of the Medicare program and other patients.

Average cost per diem for general routine services means the following:

(1) For cost reporting periods beginning on or after October 1, 1982, subject to the provisions on swing-bed hospitals, the average cost of general routine services net of the private room cost differential. The average cost per diem is computed by the following methodology:

(i) Determine the total private room cost differential by multiplying the average per diem private room cost differential determined in paragraph (c) of this section by the total number of private room patient days.

(ii) Determine the total inpatient general routine service costs net of the total private room cost differential by subtracting the total private room cost differential from total inpatient general routine service costs.

(iii) Determine the average cost per diem by dividing the total inpatient general routine service costs net of private room cost differential by all inpatient general routine days, including total private room days.

(2) For swing-bed hospitals, the amount computed by—

(i) Subtracting the routine costs associated with Medicare SNF-type days and non-Medicare NF-type days from the total allowable inpatient cost for routine services (excluding the cost of services provided in intensive care units, coronary care units, and other intensive care type inpatient hospital units and nursery costs); and

(ii) Dividing the remainder (excluding the total private room cost differential) by the total number of inpatient hospital days of care (excluding Medicare SNF-type days and non-Medicare NF-type days of care, days of care in intensive care units, coronary care units, and other intensive care type inpatient hospital units; and newborn days; but including total private room days).

Average cost per diem for hospital intensive care type units means the amount computed by dividing the total allowable costs for routine services in each of these units by the total number of inpatient days of care furnished in each of these units.

Average per diem private room cost differential means the difference in the average per diem cost of furnishing routine services in a private room and in a
semi-private room. (This differential is not applicable to hospital intensive care type units.) (The method for computing this differential is described in paragraph (c) of this section.)

Charges means the regular rates for various services that are charged to both beneficiaries and other paying patients who receive the services. Implicit in the use of charges as the basis for apportionment is the objective that charges for services be related to the cost of the services.

Intensive care type inpatient hospital unit means a hospital unit that furnishes services to critically ill inpatients. Examples of intensive care type units include, but are not limited to, intensive care units, trauma units, coronary care units, pulmonary care units, and burn units. Excluded as intensive care type units are postoperative recovery rooms, postanesthesia recovery rooms, maternity labor rooms, and subintensive or intermediate care units. (The unit must also meet the criteria of paragraph (d) of this section.)

Nursing facility (NF)-type services, formerly known as ICF and SNF-type services, are routine services furnished by a swing-bed hospital to Medicaid and other non-Medicare patients. Under the Medicaid program, effective October 1, 1990, facilities are no longer certified as SNFs or ICFs but instead are certified only as NFs and can provide services as defined in section 1919(a)(1) of the Act.

Skilled nursing facility (SNF)-type services are routine services furnished by a swing-bed hospital that would constitute extended care services if furnished by an SNF. SNF-type services include routine SNF services furnished in the distinct part SNF of a hospital complex that is combined with the hospital general routine service area cost center under §413.24(d)(5). Effective October 1, 1990, only Medicare covered services are included in the definition of SNF-type services.

Ratio of beneficiary charges to total charges on a departmental basis means the ratio of charges to beneficiaries of the Medicare program for services of a revenue-producing department or center to the charges to all patients for that center during an accounting period. After each revenue-producing center’s ratio is determined, the cost of services furnished to beneficiaries of the Medicare program is computed by applying the individual ratio for the center to the cost of the related center for the period.

Routine services means the regular room, dietary, and nursing services, minor medical and surgical supplies, and the use of equipment and facilities for which a separate charge is not customarily made.

(c) Method for computing the average per diem private room cost differential. Compute the average per diem private room charge differential by subtracting the average per diem charge for all semi-private room accommodations from the average per diem charge for all private room accommodations. The average per diem charge for private room accommodations is determined by dividing the total charges for private room accommodations by the total number of days of care furnished in private room accommodations. The average per diem charge for semi-private accommodations is determined by dividing the total charges for semi-private room accommodations by the total number of days of care furnished in semi-private accommodations.

(1) Determine the average per diem private room charge differential by subtracting the average per diem charge for all semi-private room accommodations from the average per diem charge for all private room accommodations. The average per diem charge for private room accommodations is determined by dividing the total charges for private room accommodations by the total number of days of care furnished in private room accommodations. The average per diem charge for semi-private accommodations is determined by dividing the total charges for semi-private room accommodations by the total number of days of care furnished in semi-private accommodations.

(2) Determine the inpatient general routine service cost to charge ratio by dividing total inpatient general routine service cost by the total inpatient general routine service charges.

(3) Determine the average per diem private room cost differential by multiplying the average per diem private room charge differential determined in paragraph (c)(1) of this section by the ratio determined in paragraph (c)(2) of this section.

(d) Criteria for identifying intensive care type units. For purposes of determining costs under this section, a unit will be identified as an intensive care type inpatient hospital unit only if the unit—

(1) Is in a hospital;
(2) Is physically and identifiably separate from general routine patient care.
areas, including subintensive or intermediate care units, and ancillary service areas. There cannot be a concurrent sharing of nursing staff between an intensive care type unit and units or areas furnishing different levels or types of care. However, two or more intensive care type units that concurrently share nursing staff can be reimbursed as one combined intensive care type unit if all other criteria are met. Float nurses (nurses who work in different units on an as-needed basis) can be utilized in the intensive care type unit. If a float nurse works in two different units during the same eight hour shift, then the costs must be allocated to the appropriate units depending upon the time spent in those units. The hospital must maintain adequate records to support the allocation. If such records are not available, then the costs must be allocated to the general routine services cost areas;

(3) Has specific written policies that include criteria for admission to, and discharge from, the unit;

(4) Has registered nursing care available on a continuous 24-hour basis with at least one registered nurse present in the unit at all times;

(5) Maintains a minimum nurse-patient ratio of one nurse to two patients per patient day. Included in the calculation of this nurse-patient ratio are registered nurses, licensed vocational nurses, licensed practical nurses, and nursing assistants who provide patient care. Not included are general support personnel such as ward clerks, custodians, and housekeeping personnel; and

(6) Is equipped, or has available for immediate use, life-saving equipment necessary to treat the critically ill patients for which it is designed. This equipment may include, but is not limited to, respiratory and cardiac monitoring equipment, respirators, cardiac defibrillators, and wall or canister oxygen and compressed air.

(e) Application—(1) Departmental method; Cost reporting periods beginning on or after October 1, 1982.

(i) The following example illustrates how costs would be determined, using only inpatient data, for cost reporting periods beginning on or after October 1, 1982, based on apportionment of—

(A) The average cost per diem for general routine services (subject to the private room differential provisions of paragraph (a)(1)(iii) of this section);

(B) The average cost per diem for each intensive care type unit;

(C) The ratio of beneficiary charges to total charges applied to cost by department.

<table>
<thead>
<tr>
<th>Department</th>
<th>Charges to program beneficiaries</th>
<th>Total charges</th>
<th>Ratio of beneficiary charges to total charges</th>
<th>Total cost</th>
<th>Cost of beneficiary services</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Percent</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Operating rooms</td>
<td></td>
<td>$20,000</td>
<td>$70,000 (28%)</td>
<td>$77,000</td>
<td>$22,000</td>
</tr>
<tr>
<td>Delivery rooms</td>
<td></td>
<td>0</td>
<td>12,000 (33%)</td>
<td>30,000</td>
<td>0</td>
</tr>
<tr>
<td>Pharmacy</td>
<td></td>
<td>20,000</td>
<td>60,000 (28%)</td>
<td>45,000</td>
<td>15,000</td>
</tr>
<tr>
<td>X-ray</td>
<td></td>
<td>24,000</td>
<td>100,000 (24%)</td>
<td>75,000</td>
<td>18,000</td>
</tr>
<tr>
<td>Laboratory</td>
<td></td>
<td>40,000</td>
<td>140,000 (28%)</td>
<td>98,000</td>
<td>26,000</td>
</tr>
<tr>
<td>Others</td>
<td></td>
<td>6,000</td>
<td>30,000 (20%)</td>
<td>25,000</td>
<td>5,000</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td>110,000</td>
<td>412,000 (9,200%)</td>
<td>350,000</td>
<td>86,000</td>
</tr>
</tbody>
</table>

Total inpatient days | Total cost | Average cost per diem | Program inpatient days | Cost of beneficiary services |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>General routine</td>
<td>30,000</td>
<td>$630,000</td>
<td>$21</td>
<td>8,000</td>
</tr>
<tr>
<td>Coronary care unit</td>
<td>500</td>
<td>20,000</td>
<td>40</td>
<td>200</td>
</tr>
<tr>
<td>Intensive care unit</td>
<td>3,000</td>
<td>108,000</td>
<td>36</td>
<td>1,000</td>
</tr>
<tr>
<td>Total</td>
<td>33,500</td>
<td>758,000</td>
<td>9,200</td>
<td>212,000</td>
</tr>
</tbody>
</table>
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(ii) The following illustrates how apportionment based on an average cost per diem for general routine services is determined.

**HOSPITAL E**

<table>
<thead>
<tr>
<th></th>
<th>Facts</th>
<th>Private accommodations</th>
<th>Semi-private accommodations</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total charges</td>
<td>$20,000</td>
<td>$175,000</td>
<td>$195,000</td>
<td></td>
</tr>
<tr>
<td>Total days</td>
<td>100</td>
<td>1,000</td>
<td>1,100</td>
<td></td>
</tr>
<tr>
<td>Programs days</td>
<td>70</td>
<td>400</td>
<td>470</td>
<td></td>
</tr>
<tr>
<td>Medically necessary for program beneficiaries</td>
<td>$162,885</td>
<td>$2,115</td>
<td>$165,000</td>
<td></td>
</tr>
<tr>
<td>Average private room per diem charge ($20,000 private room charges ÷ 100 days)</td>
<td>$200</td>
<td>$175</td>
<td>$375</td>
<td></td>
</tr>
<tr>
<td>Average semi-private room per diem charge ($175 private room charge + 1,100 days)</td>
<td>$1,115</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1 Per diem.
2 Average per diem private room cost differential.
3 Inpatient general routine cost/charge ratio ($165,000 total costs ÷ $195,000 total charges), 0.8461538.
4 Average per diem private room cost differential ($25 charge differential × $175 average per diem private room charge), $21.15.
5 Total private room cost differential ($21.15 average per diem private room cost differential × 100 private room days), $2,115.
6 Average cost per diem for inpatient general routine services ($162,885 routine cost net of private room cost differential = $2,115 private room cost differential), $162.885.
7 Total routine per diem cost applicable to Medicare ($148.08 average cost per diem × 470 Medicare private and semi-private patient days), $69,598.
8 Medicare inpatient general routine service cost ($21.15 average per diem private room cost differential × 20 medically necessary private room days), $423.
9 Medicare inpatient general routine service cost ($423 Medicare private room cost differential + $69,598 Medicare cost of general routine inpatient services), $70,021.

(2) Carve out method. The following illustrates how apportionment is determined in a hospital reimbursed under the carve out method (subject to the private room differential provisions of paragraph (a)(1)(ii) of this section):

**HOSPITAL K**

[Determination of cost of routine SNF-type and ICF-type services and general routine hospital services]

<table>
<thead>
<tr>
<th></th>
<th>Facts</th>
<th>Days of care</th>
<th>General routine hospital</th>
<th>SNF-type</th>
<th>ICF-type</th>
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</thead>
<tbody>
<tr>
<td>Total days of care</td>
<td>2,000</td>
<td>400</td>
<td>100</td>
<td>600</td>
<td>300</td>
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<tr>
<td>Medicare days of care</td>
<td></td>
<td></td>
<td></td>
<td>N/A</td>
<td>$35</td>
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<tr>
<td>Average Medicaid rate</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>$20</td>
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<tr>
<td>Total inpatient general service costs</td>
<td>$250,000</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Calculation of cost of routine SNF-type services applicable to Medicare:

$35 × 300 = $10,500

Calculation of cost of general routine hospital services:

Cost of SNF-type services: $35 × 400 = $14,000
Cost of ICF-type services: $20 × 100 = $2,000
Total cost of the general routine hospital services: $16,000

Average cost per diem general routine hospital services: $250,000 ÷ $16,000 = 2,000 days = $10
Medicare general routine hospital costs: $117 × 600 = $70,200
Total Medicare reasonable cost for general routine inpatient days:

$10.50 × 70,200 = $703,000

Subpart E—Payments to Providers

§ 413.60 Payments to providers: General.

(a) The fiscal intermediaries will establish a basis for interim payments to each provider. This may be done by one of several methods. If an intermediary is already paying the provider on a cost basis, the intermediary may adjust its rate of payment to an estimate of the result under the Medicare principles of reimbursement. If no organization is paying the provider on a cost basis, the intermediary may obtain the previous year’s financial statement from the provider and, by applying the principles of reimbursement, compute or approximate an appropriate rate of payment. The interim payment may be related to the last year’s average per diem, or to charges, or to any other ready basis of approximating costs.

(b) At the end of the period, the actual apportionment, based on the cost finding and apportionment methods selected by the provider, determines the
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§ 413.64 Payments to providers: Specific rules.

(a) Reimbursement on a reasonable cost basis. Providers of services paid on the basis of the reasonable cost of services furnished to beneficiaries will receive interim payments approximating the actual costs of the provider. These payments will be made on the most expeditious schedule administratively feasible but not less often than monthly. A retroactive adjustment based on actual costs will be made at the end of a reporting period.

(b) Amount and frequency of payment. Medicare states that providers of services will be paid the reasonable cost of services furnished to beneficiaries. Since actual costs of services cannot be determined until the end of the accounting period, the providers must be paid on an estimated cost basis during the year. While Medicare provides that interim payments will be made no less often than monthly, intermediaries are expected to make payments on the most expeditious basis administratively feasible. Whatever estimated cost basis is used for determining interim payments during the year, the intent is that the interim payments shall approximate actual costs as nearly as is practicable so that the retroactive adjustment based on actual costs will be as small as possible.

(c) Interim payments during initial reporting period. At the beginning of the program or when a provider first participates in the program, it will be necessary to establish interim rates of payment to providers of services. Once a provider has filed a cost report under the Medicare program, the cost report may be used as a basis for determining the interim rate of reimbursement for the following period. However, since initially there is no previous history of cost under the program, the interim rate of payment must be determined by other methods, including the following:

(1) If the intermediary is already paying the provider on a cost or cost-related basis, the intermediary will adjust its rate of payment to the program’s principles of reimbursement. This rate may be either an amount per inpatient day, or a percent of the provider’s charges for services furnished to the program’s beneficiaries.

(2) If an organization other than the intermediary is paying the provider for services on a cost or cost-related basis, the intermediary may obtain from that organization or from the provider itself the rate of payment being used and other cost information as may be needed to adjust that rate of payment to give recognition to the program’s principles of reimbursement.

(3) If no organization is paying the provider on a cost or cost-related basis, the intermediary will obtain the previous year’s financial statement from the provider. By analysis of such statement in light of the principles of reimbursement, the intermediary will compute an appropriate rate of payment.

(4) After the initial interim rate has been set, the provider may at any time request, and be allowed, an appropriate increase in the computed rate, upon presentation of satisfactory evidence to the intermediary that costs have increased. Likewise, the intermediary may adjust the interim rate of payment if it has evidence that actual costs may fall significantly below the computed rate.

(d) Interim payments for new providers. (1) Newly-established providers will not have cost experience on which to base a determination of an interim rate of payment. In such cases, the intermediary will use the following methods to determine an appropriate rate:

(i) If there is a provider or providers comparable in substantially all relevant factors to the provider for which the rate is needed, the intermediary will base an interim rate of payment on the costs of the comparable provider.
(ii) If there are no substantially comparable providers from whom data are available, the intermediary will determine an interim rate of payment based on the budgeted or projected costs of the provider.

(2) Under either method, the intermediary will review the provider’s cost experience after a period of three months. If need for an adjustment is indicated, the interim rate of payment will be adjusted in line with the provider’s cost experience.

(2) Interim payments after initial reporting period. Interim rates of payment for services provided after the initial reporting period will be established on the basis of the cost report filed for the previous year covering Medicare services. The current rate will be determined—whether on a per diem or percentage of charges basis—using the previous year’s costs of covered services and making any appropriate adjustments required to bring, as closely as possible, the current year’s rate of interim payment into agreement with current year’s costs. This interim rate of payment may be adjusted by the intermediary during an accounting period if the provider submits appropriate evidence that its actual costs are or will be significantly higher than the computed rate. Likewise, the intermediary may adjust the interim rate of payment if it has evidence that actual costs may fall significantly below the computed rate. (f) Retroactive adjustment. (1) Medicare provides that providers of services will be paid amounts determined to be due, but not less often than monthly, with necessary adjustments due to previously made overpayments or underpayments. Interim payments are made on the basis of estimated costs. Actual costs reimbursable to a provider cannot be determined until the cost reports are filed and costs are verified. Therefore, a retroactive adjustment will be made at the end of the reporting period to bring the interim payments made to the provider during the period into agreement with the reimbursable amount payable to the provider for the services furnished to program beneficiaries during that period.

(2) In order to reimburse the provider as quickly as possible, an initial retroactive adjustment will be made as soon as the cost report is received. For this purpose, the costs will be accepted as reported, unless there are obvious errors or inconsistencies, subject to later audit. When an audit is made and the final liability of the program is determined, a final adjustment will be made.

(3) To determine the retroactive adjustment, the amount of the provider’s total allowable cost apportioned to the program for the reporting year is computed. This is the total amount of reimbursement the provider is due to receive from the program and the beneficiaries for covered services furnished during the reporting period. The total of the interim payments made by the program in the reporting year and the deductibles and coinsurance amounts receivable from beneficiaries is computed. The difference between the reimbursement due and the payments made is the amount of the retroactive adjustment.

(g) Accelerated payments to providers. Upon request, an accelerated payment may be made to a provider of services that is not receiving periodic interim payments under paragraph (h) of this section if the provider has experienced financial difficulties due to a delay by the intermediary in making payments or in exceptional situations, in which the provider has experienced a temporary delay in preparing and submitting bills to the intermediary beyond its normal billing cycle. Any such payment must be approved first by the intermediary and then by HCFA. The amount of the payment is computed as a percentage of the net reimbursement for unbilled or unpaid covered services. Recovery of the accelerated payment may be made by recoupment as provider bills are processed or by direct payment.

(h) Periodic interim payment method of reimbursement—(1) Covered services furnished before July 1, 1987. In addition to the regular methods of interim payment on individual provider billings for covered services, the periodic interim payment (PIP) method is available for Part A hospital and SNF inpatient services.

(2) Covered services furnished on or after July 1, 1987. Effective with claims
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received on or after July 1, 1987, the periodic interim payment (PIP) method is available for the following:

(i) Part A inpatient hospital services furnished in hospitals that are excluded from the prospective payment systems under subpart B of part 412 of this chapter.

(ii) Part A services furnished in hospitals receiving payment in accordance with a demonstration project authorized under section 402(a) of Public Law 90-248 (42 U.S.C. 1395b-1) or section 222(a) of Public Law 92-603 (42 U.S.C. 1395b-1 (note)), or a State reimbursement control system approved under section 1886(c) of the Act and subpart C of part 403 of this chapter, if that type of payment is specifically approved by HCFA as an integral part of the demonstration or control system. If that type of payment is not an integral part of the demonstration or control system, PIP is available for the hospital under paragraph (h)(1)(i) of this section for hospitals excluded from the prospective payment systems or under § 412.116(b) of this chapter for prospective payment hospitals.

(iii) Part A SNF services furnished in cost reporting periods beginning before July 1, 1998. (For services furnished in subsequent cost reporting periods, see § 413.350 regarding periodic interim payments for skilled nursing facilities).

(iv) Part A services furnished in hospitals paid under the prospective payment system, including distinct part psychiatric or rehabilitation units, as described in § 412.116(b) of this chapter.

(v) Services furnished in a hospice as specified in part 418 of this chapter. Payment on a PIP basis is described in § 418.307 of this chapter.

(3) Any participating provider furnishing the services described in paragraph (h)(1) of this section that establishes to the satisfaction of the intermediary that it meets the following requirements may elect to be reimbursed under the PIP method, beginning with the first month after its request that the intermediary finds administratively feasible:

(i) The provider’s estimated total Medicare reimbursement for inpatient services is at least $25,000 a year computed under the PIP formula; or

(B) Medicare reimbursement computed under the PIP formula is at least 50 percent of estimated total allowable cost.

(ii) The provider has filed at least one completed Medicare cost report accepted by the intermediary as providing an accurate basis for computation of program payment (except in the case of a provider requesting reimbursement under the PIP method upon first entering the Medicare program).

(iii) The provider has the continuing capability of maintaining in its records the cost, charge, and statistical data needed to accurately complete a Medicare cost report on a timely basis.

(iv) The provider has repaid or agrees to repay any outstanding current financing payment in full, such payment to be made before the effective date of its requested conversion from a regular interim payment method to the PIP method.

(4) No conversion to the PIP method may be made with respect to any provider until after that provider has repaid in full its outstanding current financing payment.

(5) The intermediary’s approval of a provider’s request for reimbursement under the PIP method will be conditioned upon the intermediary’s best judgment as to whether payment can be made to the provider under the PIP method without undue risk of its resulting in an overpayment because of greatly varying or substantially declining Medicare utilization, inadequate billing practices, or other circumstances. The intermediary may terminate PIP reimbursement to a provider at any time it determines that the provider no longer meets the qualifying requirements or that the provider’s experience under the PIP method shows that proper payment cannot be made under this method.

(6) Payment will be made biweekly under the PIP method unless the provider requests a longer fixed interval (not to exceed one month) between payments. The payment amount will be computed by the intermediary to approximate, on the average, the cost
of covered inpatient or home health services furnished by the provider during the period for which the payment is to be made, and each payment will be made two weeks after the end of such period of services. Upon request, the intermediary will, if feasible, compute the provider’s payments to recognize significant seasonal variation in Medicare utilization of services on a quarterly basis starting with the beginning of the provider’s reporting year.

(7) A provider’s PIP amount may be appropriately adjusted at any time if the provider presents or the intermediary otherwise obtains evidence relating to the provider’s costs or Medicare utilization that warrants such adjustment. In addition, the intermediary will recompute the payment immediately upon completion of the desk review of a provider’s cost report and also at regular intervals not less often than quarterly. The intermediary may make a retroactive lump sum interim payment to a provider, based upon an increase in its PIP amount, in order to bring past interim payments for the provider’s current cost reporting period into line with the adjusted payment amount. The objective of intermediary monitoring of provider costs and utilization is to assure payments approximating, as closely as possible, the reimbursement to be determined at settlement for the cost reporting period. A significant factor in evaluating the amount of the payment in terms of the realization of the projected Medicare utilization of services is the timely submittal to the intermediary of completed admission and billing forms. All providers must complete billings in detail under this method as under regular interim payment procedures.

(i) Bankruptcy or insolvency of provider. If on the basis of reliable evidence, the intermediary has a valid basis for believing that, with respect to a provider, proceedings have been or will shortly be instituted in a State or Federal court for purposes of determining whether such provider is insolvent or bankrupt under an appropriate State or Federal law, any payments to the provider will be adjusted by the intermediary, notwithstanding any other regulation or program instruction regarding the timing or manner of such adjustments, to a level necessary to insure that no overpayment to the provider is made.

(j) Interest payments resulting from judicial review—(1) Application. If a provider of services seeks judicial review by a Federal court of a decision furnished by the Provider Reimbursement Review Board or subsequent reversal, affirmation, or modification by the Secretary, the amount of any award of such Federal court will be increased by interest payable by the party against whom the judgment is made (see §413.153 for treatment of interest). The interest is payable for the period beginning on the first day of the first month following the 180-day period which began on either the date the intermediary made a final determination or the date the intermediary would have made a final determination had it been done on a timely basis (see §§405.1839(b) and 405.1841(a) of this chapter).

(2) Amount due. Section 1878(f) of the Act, 42 U.S.C. 1395oo(f), authorizes a court to award interest in favor of the prevailing party on any amount due as a result of the court’s decision. If the intermediary withheld any portion of the amount in controversy prior to the date the provider seeks judicial review by a Federal court, and the Medicare program is the prevailing party, interest is payable by the provider only on the amount not withheld. Similarly, if the Medicare program seeks to recover amounts previously paid to a provider, and the provider is the prevailing party, interest on the amounts previously paid to a provider is not payable by the Medicare program since that amount had been paid and is not due the provider.

(3) Rate. The amount of interest to be paid is equal to the rate of return on equity capital (see §413.157) in effect for the month in which the civil action is commenced.

Example: An intermediary made a final determination on the amount of Medicare program reimbursement on June 15, 1974, and the provider appealed that determination to the Provider Reimbursement Review Board. The Board heard the appeal and rendered a decision adverse to the provider. On October 28, 1974, the provider commenced civil action to have such decision reviewed. The rate of
§ 413.65 Requirements for a determination that a facility or an organization has provider-based status.

(a) Scope and definitions.

(1) Scope. This section applies to all facilities or organizations for which provider-based status is sought, including remote locations of hospitals, as defined in paragraph (a)(2) of this section and satellite facilities as defined in § 412.22(h)(1) and § 412.25(e)(1) of this chapter, other than ESRD facilities. Determinations for ESRD facilities are made under § 413.174 of this chapter.

(2) Definitions. In this subpart E, unless the context indicates otherwise—

Campus means the physical area immediately adjacent to the provider’s main buildings, other areas and structures that are not strictly contiguous to the main buildings but are located within 250 yards of the main buildings, and any other areas determined on an individual case basis, by the HCFA regional office, to be part of the provider’s campus.

Department of a provider means a facility or organization or a physician office that is either created by, or acquired by, a main provider for the purpose of furnishing health care services of the same type as those furnished by the main provider under the name, ownership, and financial and administrative control of the main provider, in accordance with the provisions of this section. A department of a provider may not be licensed to provide health care services in its own right, may not by itself be qualified to participate in Medicare as a provider under § 489.2 of this chapter, and Medicare conditions of participation do not apply to a department as an independent entity. For purposes of this part, the term “department of a provider” does not include an RHC or, except as specified in paragraph (m)(1) of this section, an FQHC.

Free-standing facility means an entity that furnishes health care services to Medicare beneficiaries and that is not integrated with any other entity as a main provider, a department of a provider, remote location of a hospital, satellite facility, or a provider-based entity.

Main provider means a provider that either creates, or acquires ownership of, another entity to deliver additional health care services under its name, ownership, and financial and administrative control.

Provider-based entity means a provider of health care services, or an RHC or an FQHC as defined in § 405.2401(b) of this chapter, that is either created by, or acquired by, a main provider for the purpose of furnishing health care services of a different type from those of the main provider under the name, ownership, and administrative and financial control of the main provider, in accordance with the provisions of this section.

Provider-based status means the relationship between a main provider and a provider-based entity or a department of a provider, remote location of a hospital, or satellite facility, that complies with the provisions of this section.

Remote location of a hospital means a facility or an organization that is either created by, or acquired by, a hospital that is a main provider for the purpose of furnishing inpatient hospital services under the name, ownership, and financial and administrative control of the main provider, in accordance with the provisions of this section. A remote location of a hospital may not be licensed to provide inpatient hospital services in its own right, and Medicare conditions of participation do not apply to a remote location of a hospital as an independent entity. For purposes of this part, the term “remote location of a hospital” does not include a satellite facility as defined in § 412.22(h)(1) and § 412.25(e)(1) of this chapter.

(b) Responsibility for obtaining provider-based determinations. (1) A facility or organization is not entitled to be treated as provider-based simply because it or the main provider believe it is provider-based.
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(2) A main provider or a facility or organization must contact HCFA and the facility or organization must be determined by HCFA to be provider-based before the main provider bills for services of the facility or organization as if the facility or organization were provider-based, or before it includes costs of those services on its cost report.

(3) A facility that is not located on the campus of a hospital and is used as a site of physician services of the kind ordinarily furnished in physician offices will be presumed to be a free-standing facility, unless it is determined by HCFA to have provider-based status.

(c) Reporting. (1) A main provider that creates or acquires a facility or organization for which it wishes to claim provider-based status, including any physician offices that a hospital wishes to operate as a hospital outpatient department or clinic, must report its acquisition of the facility or organization to HCFA. If the facility or organization is located off the campus of the provider, or inclusion of the costs of the facility or organization in the provider’s cost report would increase the total costs on the provider’s cost report by at least 5 percent, and must furnish all information needed for a determination as to whether the facility or organization meets the requirements in paragraph (d) of this section for provider-based status.

(2) A main provider that has had one or more facilities or organizations considered provider-based also must report to HCFA any material change in the relationship between it and any provider-based facility or organization, such as a change in ownership of the facility or organization, entry into a new or different management contract that could affect the provider-based status of the facility or organization.

(d) Requirements. An entity must meet all of the following requirements to be determined by HCFA to have provider-based status.

(1) Licensure. The department of the provider, remote location of a hospital, or satellite facility, or in States where State law does not permit licensure of the provider and the prospective department of the provider, remote location of a hospital, or satellite facility under a single license. If a State health facilities’ cost review commission or other agency that has authority to regulate the rates charged by hospitals or other providers in a State finds that a particular facility or organization is not part of a provider, HCFA will determine that the facility or organization does not have provider-based status.

(2) Operation under the ownership and control of the main provider. The facility or organization seeking provider-based status is operated under the ownership and control of the main provider, as evidenced by the following:

(i) The business enterprise that constitutes the facility or organization is 100 percent owned by the provider.

(ii) The main provider and the facility or organization seeking status as a department of the provider, remote location of a hospital, or satellite facility have the same governing body.

(iii) The facility or organization is operated under the same organizational documents as the main provider. For example, the facility or organization seeking provider-based status must be subject to common bylaws and operating decisions of the governing body of the provider where it is based.

(iv) The main provider has final responsibility for administrative decisions, final approval for contracts with outside parties, final approval for personnel actions, final responsibility for personnel policies (such as fringe benefits/code of conduct), and final approval for medical staff appointments in the facility or organization.

(3) Administration and supervision. The reporting relationship between the facility or organization seeking provider-based status and the main provider must have the same frequency, intensity, and level of accountability that exists in the relationship between the main provider and one of its departments, as evidenced by compliance with all of the following requirements:

(i) The facility or organization is under the direct supervision of the main provider.
(ii) The facility or organization is operated under the same monitoring and oversight by the provider as any other department of the provider, and is operated just as any other department of the provider with regard to supervision and accountability. The facility or organization director or individual responsible for daily operations at the entity—

(A) Maintains a reporting relationship with a manager at the main provider that has the same frequency, intensity, and level of accountability that exists in the relationship between the main provider and its departments; and

(B) Is accountable to the governing body of the main provider, in the same manner as any department head of the provider.

(iii) The following administrative functions of the facility or organization are integrated with those of the provider where the facility or organization is based: billing services, records, human resources, payroll, employee benefit package, salary structure, and purchasing services. Either the same employees or group of employees handle these administrative functions for the facility or organization and the main provider, or the administrative functions for both the facility or organization and the entity are—

(A) Contracted out under the same contract agreement; or

(B) Handled under different contract agreements, with the contract of the facility or organization being managed by the main provider.

(4) Clinical services. The clinical services of the facility or organization seeking provider-based status and the main provider are integrated as evidenced by the following:

(i) Professional staff of the facility or organization have clinical privileges at the main provider.

(ii) The main provider maintains the same monitoring and oversight of the facility or organization as it does for any other department of the provider.

(iii) The medical director of the facility or organization seeking provider-based status maintains a reporting relationship with the Chief Medical Officer or other similar official of the main provider, and is under the same type of supervision and accountability as any other director, medical or otherwise, of the main provider.

(iv) Medical staff committees or other professional committees at the main provider are responsible for medical activities in the facility or organization including quality assurance, utilization review, and the coordination and integration of services, to the extent practicable, between the facility or organization seeking provider-based status and the main provider.

(v) Medical records for patients treated in the facility or organization are integrated into a unified retrieval system (or cross reference) of the main provider.

(vi) Inpatient and outpatient services of the facility or organization and the main provider are integrated, and patients treated at the facility or organization who require further care have full access to all services of the main provider and are referred where appropriate to the corresponding inpatient or outpatient department or service of the main provider.

(5) Financial integration. The financial operations of the facility or organization are fully integrated within the financial system of the main provider, as evidenced by shared income and expenses between the main provider and the facility or organization. The costs of the facility or organization are reported in a cost center of the provider, and the financial status of the facility or organization is incorporated and readily identified in the main provider's trial balance.

(6) Public awareness. The facility or organization seeking status as a department of a provider, remote location of a hospital, or satellite facility is held out to the public and other payers as part of the main provider. When patients enter the provider-based facility or organization, they are aware that they are entering the main provider and are billed accordingly.

(7) Location in immediate vicinity. The facility or organization and the main
§413.65 Provider-based status not applicable to joint ventures.

A facility or organization cannot be considered provider-based if the entity is owned by two or more providers engaged in a joint venture. For example, where a hospital has jointly purchased or jointly created free-standing facilities under joint venture arrangements, neither party to the joint venture arrangement can claim the free-standing facility as a provider-based entity.

(f) Management contracts. Facilities and organizations that otherwise meet the requirements of paragraph (d) of this section, but are operated under management contracts, must also meet all of the following criteria:

(1) The staff of the facility or organization, other than management staff, are employed by the provider or by another organization, other than the management company, which also employs the staff of the main provider.

(2) The administrative functions of the facility or organization are integrated with those of the main provider, as determined under criteria in paragraph (d)(3)(iii) of this section.

(3) The main provider has significant control over the operations of the facility or organization as determined under criteria in paragraph (b)(3)(ii) of this section.

(4) The management contract is held by the main provider itself, not by a parent organization that has control over both the main provider and the facility or organization.

(g) Obligations of hospital outpatient departments and hospital-based entities.

(1) Hospital outpatient departments located either on or off the campus of the hospital that is the main provider must comply with the anti-dumping rules in §§489.20(l), (m), (q), and (r) and §489.24 of this chapter. If any individual comes to any hospital-based entity (including an RHC) located on the main hospital campus, and a request is made on the individual’s behalf for examination or treatment of a medical condition, as described in §489.24 of this chapter, the hospital must comply with the anti-dumping rules in §489.24 of this chapter.

(2) Physician services furnished in hospital outpatient departments or hospital-based entities (other than
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HCFA, HHS must be billed with the correct site-of-service indicator, so that applicable site-of-service reductions to physician and practitioner payment amounts can be applied.

(3) Hospital outpatient departments must comply with all the terms of the hospital’s provider agreement.

(4) Physicians who work in hospital outpatient departments or hospital-based entities are obligated to comply with the non-discrimination provisions in §489.10(b) of this chapter.

(5) Hospital outpatient departments (other than RHCS) must treat all Medicare patients, for billing purposes, as hospital outpatients. The department must not treat some Medicare patients as hospital outpatients and others as physician office patients.

(6) In the case of a patient admitted to the hospital as an inpatient after receiving treatment in the hospital outpatient department or hospital-based entity, payments for services in the hospital outpatient department or hospital-based entity are subject to the payment window provisions applicable to PPS hospitals and to hospitals and units excluded from PPS set forth at §412.2(c)(5) of this chapter and at §413.40(c)(2), respectively.

(7) When a Medicare beneficiary is treated in a hospital outpatient department or hospital-based entity (other than an RHC) that is not located on the main provider’s campus, the hospital has a duty to provide written notice to the beneficiary, prior to the delivery of services, of the amount of the beneficiary’s potential financial liability (that is, of the fact that the beneficiary will incur a coinsurance liability for an outpatient visit to the hospital as well as for the physician service, and of the amount of that liability). The notice must be one that the beneficiary can read and understand. If the beneficiary is unconscious, under great duress, or for any other reason unable to read a written notice and understand and act on his or her own rights, the notice must be provided, prior to the delivery of services, to the beneficiary’s authorized representative.

(8) Hospital outpatient departments must meet applicable hospital health and safety rules for Medicare-participating hospitals in part 482 of this chapter.

(h) Furnishing all services under arrangement. A facility or organization may not qualify for provider-based status if all patient care services furnished at the facility are furnished under arrangement.

(i) Inappropriate treatment of a facility or organization as provider-based. (1) Determination and review. If HCFA learns that a provider has treated a facility or organization as provider-based and the provider had not obtained a determination of provider-based status under this section, HCFA will—

(i) Review current payments and, if necessary, take action in accordance with the rules on inappropriate billing in paragraph (j) of this section;

(ii) Investigate and determine whether the requirements in paragraph (d) of this section (or, for periods prior to October 10, 2000, the requirements in applicable program instructions) were met; and

(iii) Review all previous payments to that provider for all cost reporting periods subject to re-opening in accordance with §405.1885 and §405.1889 of this chapter.

(2) Recovery of overpayments. If HCFA finds that payments for services at the facility or organization have been made as if the facility or organization were provider-based, even though HCFA had not previously determined that the facility or organization qualified for provider-based status, HCFA will recover the difference between the amount of payments that actually were made and the amount of payments that HCFA estimates should have been made in the absence of a determination of provider-based status. Recovery will not be made for any main provider cost reporting periods beginning before January 10, 2001 or, in the case of a facility organization paid as a provider-based entity, for that entity’s cost reporting periods beginning before January 10, 2001 if, during all of those periods, the management of the facility or organization made a good faith effort to operate it as a provider-based facility or organization, as described in paragraph (h)(3) of this section.
Exception for good faith effort. HCFA determines that the management of a facility or organization has made a good faith effort to operate it as a provider-based entity if—

(i) The requirements regarding licensure and public awareness in paragraphs (d)(1) and (d)(6) of this section are met;

(ii) All facility services were billed as if they had been furnished by a department of a provider, remote location of a hospital, satellite facility, or a provider-based entity of the main provider; and

(iii) All professional services of physicians and other practitioners were billed with the correct site-of-service indicator, as described in paragraph (g)(2) of this section.

Inappropriate billing. If HCFA finds that a facility or organization is being treated as provider-based without having obtained a determination of provider-based status under this section, HCFA will notify the provider, adjust future payments, review previous payments, determine whether the facility or organization qualifies for provider-based status under the criteria in this section. If HCFA determines that the facility or organization qualifies for provider-based status, future payment for services at or by the facility or organization will be adjusted to reflect that determination. If HCFA determines that the facility or organization does not qualify for provider-based status, future payment for services at or by the facility or organization will be made only in accordance with the rules in paragraph (i)(5) of this section.

(1) Notice to provider. If HCFA finds that inappropriate billing has occurred or is occurring since no provider-based determination has been made by HCFA, HCFA will issue written notice to the provider that payments for past cost reporting periods may be reviewed and recovered as described in paragraph (i)(1) of this section, that future payments for services in or of the facility or organization will be adjusted as described in paragraph (j)(2) of this section, and that a determination of provider-based status will be made.

(2) Adjustment of payments. If HCFA finds that inappropriate billing has occurred or is occurring since no provider-based determination has been made by HCFA, HCFA will adjust future payments to the provider, the facility or organization, or both, to approximate as closely as possible the amounts that would have been paid in the absence of a provider-based determination, if all other requirements for billing were met.

(3) Review of previous payments. If HCFA finds that inappropriate billing has occurred or is occurring since no provider-based determination has been made by HCFA, HCFA will review previous payments and, if necessary, take action in accordance with the rules on inappropriate treatment of a facility or organization as provider-based in paragraph (h) of this section.

(4) Determination regarding provider-based status. If HCFA finds that inappropriate billing has occurred or is occurring since no provider-based determination has been made by HCFA, HCFA will determine whether the facility or organization qualifies for provider-based status under the criteria in this section.

(5) Continuation of payment. The notice of denial of provider-based status sent to the provider will ask the provider to notify HCFA in writing, within 30 days of the date the notice is issued, of whether the facility or organization (or, where applicable, the practitioners who staff the facility or organization) will be seeking to enroll and meet other requirements to bill for services in a free-standing facility. If the provider notifies HCFA that it is seeking to enroll, or if HCFA does not receive a response within 30 days of the date the notice was issued, all payment under this paragraph (i)(5) will end as of the 30th day after the date of notice. If the provider indicates that the facility or organization, or its practitioners, will be seeking to meet enrollment and other requirements for billing for services in a free-standing facility, payment for services of the facility or organization will continue, at the adjusted amounts described in paragraph (j)(2) of this section for as long...
as is required for all billing requirements to be met (but not longer than 6 months) if the facility or organization, or its practitioners, submit a complete enrollment application and provide all other required information within 90 days after the date of notice; and the facility or organization, or its practitioners, furnish all other information needed by HCFA to process the enrollment application and verify that other billing requirements are met. If the necessary applications or information are not provided, HCFA will terminate all payment to the provider, facility, or organization as of the date HCFA issues notice that necessary applications or information have not been submitted.

(k) Correction of errors. HCFA may review a past determination of provider-based status for a facility or organization or may review the status of a facility or organization (that is, whether the facility or organization is provider-based) if no determination regarding provider-based status has previously been made, if HCFA believes that status may be inappropriate, based on the provisions of this section. If HCFA determines that a previous determination was in error, and the entity should not be considered provider-based, HCFA notifies the main provider. Treatment of the facility or organization as provider-based ceases with the first day of the next cost report period following notification of the redetermination, but not less than 6 months after the date of notification.

(l) Status of Indian Health Service and Tribal facilities and organizations. Facilities and organizations operated by the Indian Health Service or Tribes will be considered to be departments of hospitals operated by the Indian Health Service or Tribes if, on or before April 7, 2000, they furnished only services that were billed as if they had been furnished by a department of a provider will continue to be treated, for purposes of this section, as a department of the provider without regard to whether it complies with the criteria for provider-based status in this section, if the facility—

(1) Received a grant on or before April 7, 2000 under section 330 of the Public Health Service Act and continues to receive funding under such a grant, or is receiving funding from a grant made on or before April 7, 2000 under section 330 of the Public Health Service Act under a contract with the recipient of such a grant, and continues to meet the requirements to receive a grant under section 330 of the Public Health Service Act; or

(2) Based on the recommendation of the Public Health Service, was determined by HCFA on or before April 7, 2000 to meet the requirements for receiving a grant under section 330 of the Public Health Service Act, and continues to meet such requirements.

(m) FOHCs and "look-alikes". A facility that has, since April 7, 1995, furnished only services that were billed as if they had been furnished by a department of a provider, will continue to be treated, for purposes of this section, as a department of the provider without regard to whether it complies with the criteria for provider-based status in this section, if the facility—

(1) Owned by the Indian Health Service but leased and operated by the Tribe under the Indian Self-Determination Act (Pub. L. 93-638) in accordance with applicable regulations and policies of the Indian Health Service in consultation with Tribes; or

(3) Owned by the Indian Health Service in consultation with Tribes; or

(n) Effective date of provider-based status. Provider-based status for a facility or organization is effective on the earliest date on which a request for provider-based status has been made, and all requirements of this part have been met.

[65 FR 18538, Apr. 7, 2000, as amended at 65 FR 47677, Aug. 3, 2000]
§ 413.65 Requirements for a determination that a facility or an organization has provider-based status.

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(i) * * *

(2) Recovery of overpayments. If HCFA finds that payments for services at the facility or organization have been made as if the facility or organization were provider-based, even though HCFA had not previously determined that the facility or organization qualified for provider-based status, HCFA will recover the difference between the amount of payments that were actually made and the amount of payments that HCFA estimates should have been made in the absence of a determination of provider-based status, except that recovery will not be made for any period prior to October 10, 2000 if during all of that period the management of the entity made a good faith effort to operate it as a provider-based facility or organization, as described in paragraph (h)(3) of this section.

* * * * *

(m) FQHCs and “look-alikes”. A facility that has, since April 7, 1995, furnished only services that were billed as if they had been furnished by a department of a provider will continue to be treated, for purposes of this section, as a department of the provider without regard to whether it complies with the criteria for provider-based status in this section, if the facility—

(1) Received a grant before 1995 under section 330 of the Public Health Service Act, or is receiving funding from such a grant under a contract with the recipient of such a grant and meets the requirements to receive a grant under section 330 of the Public Health Service Act; or

(2) Based on the recommendation of the Public Health Service, was determined by HCFA before 1995 to meet the requirements for receiving such a grant.

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§ 413.70 Payment for services of a CAH.

(a) Payment for inpatient services furnished by a CAH. (1) Payment for inpatient services furnished by a CAH is the reasonable costs of the CAH in providing CAH services to its inpatients, as determined in accordance with section 1861(v)(1)(A) of the Act and the applicable principles of cost reimbursement in this part and in Part 415 of this chapter, except that the following payment principles are excluded when determining payment for CAH inpatient services:

(i) Lesser of cost or charges;

(ii) Ceilings on hospital operating costs; and

(iii) Reasonable compensation equivalent (RCE) limits for physician services to providers.

(2) Payment to a CAH for inpatient services does not include any costs of physician services or other professional services to CAH inpatients, and is subject to the Part A hospital deductible and coinsurance, as determined under subpart G of part 409 of this chapter.

(b) Payment for outpatient services furnished by a CAH—(1) General. Unless the CAH elects to be paid for services to its outpatients under the method specified in paragraph (b)(3) of this section, the amount of payment for outpatient services of a CAH is the amount determined under paragraph (b)(2) of this section.

(2) Reasonable costs for facility services.

(i) Payment for outpatient services of a CAH is the reasonable costs of the CAH in providing CAH services to its outpatients, as determined in accordance with section 1861(v)(1)(A) of the Act and the applicable principles of cost reimbursement in this part and in Part 415 of this chapter, except that the following payment principles are excluded when determining payment for CAH outpatient services:

(A) Lesser of costs or charges;

(B) RCE limits;

(C) Any type of reduction to operating or capital costs under §413.124 or §413.130(j)(7); and

(D) Blended payment amounts for ambulatory surgical services, radiology services, and other diagnostic services;

(ii) Payment to a CAH under paragraph (b)(2) of this section does not include any costs of physician services or other professional services to CAH outpatients, and is subject to the Part B deductible and coinsurance amounts, as determined under §§410.152(k), 410.160, and 410.161 of this chapter.

(iii) The following payment principles are used when determining payment for outpatient clinical diagnostic laboratory tests:

(A) The amount paid is equal to 100 percent of the least of—
(1) Charges determined under the fee schedule as set forth in section 1833(h)(1) or section 1834(d)(1) of the Act;

(2) The limitation amount for that test determined under section 1833(h)(4)(B) of the Act or the amount of the charges billed for the test; or

(3) A negotiated rate established under section 1833(h)(6) of the Act.

(B) Payment for outpatient clinical diagnostic laboratory tests is not subject to the Medicare Part B deductible and coinsurance amounts, as specified in §410.152(k) of this chapter.

(3) Election to be paid reasonable costs for facility services plus fee schedule for professional services. (i) A CAH may elect to be paid for outpatient services in any cost reporting period under the method described in paragraphs (b)(3)(ii) and (b)(3)(iii) of this section. This election must be made in writing, made on an annual basis, and delivered to the intermediary at least 60 days before the start of each affected cost reporting period. An election of this payment method, once made for a cost reporting period, remains in effect for all of that period and applies to all services furnished to outpatients during that period.

(ii) If the CAH elects payment under this method, payment to the CAH for each outpatient visit will be the sum of the following amounts:

(A) For facility services, not including any services for which payment may be made under paragraph (b)(3)(ii)(B) of this section, the reasonable costs of the services as determined under paragraph (b)(2)(i) of this section; and

(B) For professional services otherwise payable to the physician or other practitioner on a fee schedule basis, the amounts that otherwise would be paid for the services if the CAH had not elected payment under this method.

(iii) Payment to a CAH is subject to the Part B deductible and coinsurance amounts, as determined under §§410.152, 410.160, and 410.161 of this chapter.

(c) Final payment based on cost report. Final payment to the CAH for CAH facility services to inpatients and outpatients furnished during a cost reporting period is based on a cost report for that period, as required under §413.20(b).

[65 FR 47109, Aug. 1, 2000]

§ 413.74 Payment to a foreign hospital. (a) Principle. Section 1814(f) of the Act provides for the payment of emergency and nonemergency inpatient hospital services furnished by foreign hospitals to Medicare beneficiaries. Subpart H of part 424 of this chapter, together with this section, specify the conditions for payment. These conditions may result in payments only to Canadian and Mexican hospitals.

(b) Amount of payment. Effective with admissions on or after January 1, 1980, the reasonable cost for services covered under the Medicare program furnished to beneficiaries by a foreign hospital will be equal to 100 percent of the hospital's customary charges (as defined in §413.13(b)) for the services.

(c) Submittal of claims. The hospital must establish its customary charges for the services by submitting an itemized bill with each claim it files in accordance with its election under §424.104 of this chapter.

(d) Exchange rate. Payment to the hospital will be subject to the official exchange rate on the date the patient is discharged and to the applicable deductible and coinsurance amounts described in §§409.80 through 409.83.


Subpart F—Specific Categories of Costs

§ 413.80 Bad debts, charity, and courtesy allowances.

(a) Principle. Bad debts, charity, and courtesy allowances are deductions from revenue and are not to be included in allowable cost; however, except for anesthetists' services described under paragraph (h) of this section, bad debts attributable to the deductibles and coinsurance amounts are reimbursable under the program.

(b) Definitions—(1) Bad debts. Bad debts are amounts considered to be uncollectible from accounts and notes receivable that were created or acquired in providing services. "Accounts
receivable'' and "notes receivable" are designations for claims arising from the furnishing of services, and are collectible in money in the relatively near future.

(2) Charity allowances. Charity allowances are reductions in charges made by the provider of services because of the indigence or medical indigence of the patient. Cost of free care (uncompensated services) furnished under a Hill-Burton obligation are considered as charity allowances.

(3) Courtesy allowances. Courtesy allowances indicate a reduction in charges in the form of an allowance to physicians, clergy, members of religious orders, and others as approved by the governing body of the provider, for services received from the provider. Employee fringe benefits, such as hospitalization and personnel health programs, are not considered to be courtesy allowances.

(c) Normal accounting treatment: Reduction in revenue. Bad debts, charity, and courtesy allowances represent reductions in revenue. The failure to collect charges for services furnished does not add to the cost of providing the services. Such costs have already been incurred in the production of the services.

(d) Requirements for Medicare. Under Medicare, costs of covered services furnished beneficiaries are not to be borne by individuals not covered by the Medicare program. Uncollected revenue related to services furnished to beneficiaries of the program generally means the provider has not recovered the cost of services covered by that revenue. The failure of beneficiaries to pay the deductible and coinsurance amounts could result in the related costs of covered services being borne by other than Medicare beneficiaries. To assure that such covered service costs are not borne by others, the costs attributable to the deductible and coinsurance amounts that remain unpaid are added to the Medicare share of allowable costs. Bad debts arising from other sources are not allowable costs.

(e) Criteria for allowable bad debt. A bad debt must meet the following criteria to be allowable:

(1) The debt must be related to covered services and derived from deductible and coinsurance amounts.

(2) The provider must be able to establish that reasonable collection efforts were made.

(3) The debt was actually uncollectible when claimed as worthless.

(4) Sound business judgment established that there was no likelihood of recovery at any time in the future.

(f) Charging of bad debts and bad debt recoveries. The amounts uncollectible from specific beneficiaries are to be charged off as bad debts in the accounting period in which the accounts are deemed to be worthless. In some cases an amount previously written off as a bad debt and allocated to the program may be recovered in a subsequent accounting period; in such cases the income therefrom must be used to reduce the cost of beneficiary services for the period in which the collection is made.

(g) Charity allowances. Charity allowances have no relationship to beneficiaries of the Medicare program and are not allowable costs. These charity allowances include the costs of uncompensated services furnished under a Hill-Burton obligation. (Note: In accordance with section 106(b) of Pub. L. 97-248 (enacted September 3, 1982), this sentence is effective with respect to any costs incurred under Medicare except that it does not apply to costs which have been allowed prior to September 3, 1982, pursuant to a final court order affirmed by a United States Court of Appeals.) The cost to the provider of employee fringe-benefit programs is an allowable element of reimbursement.

(h) Limitations on bad debts. In determining reasonable costs for hospitals, the amount of bad debts otherwise treated as allowable costs (as defined in paragraph (e) of this section) is reduced—

(1) For cost reporting periods beginning during fiscal year 1998, by 25 percent;

(2) For cost reporting periods beginning during fiscal year 1999, by 40 percent; and
§ 413.85 Cost of educational activities.

(a) Payment—(1) General rule. Except as provided in paragraph (a)(2) of this section, a provider's allowable cost may include its net cost of approved educational activities, as calculated under paragraph (g) of this section. The net cost is subject to apportionment based on Medicare utilization as described in §413.50.

(2) Exception. For cost reporting periods beginning on or after July 1, 1985, payment to hospitals and hospital-based providers for approved residency programs in medicine, osteopathy, dentistry, and podiatry is determined as provided in §413.86.

(b) Definition—Approved educational activities. Approved educational activities means formally organized or planned programs of study usually engaged in by providers in order to enhance the quality of patient care in an institution. These activities must be licensed if required by State law. If licensing is not required, the institution must receive approval from the recognized national professional organization for the particular activity.

(c) Educational activities. Many providers engage in educational activities including training programs for nurses, medical students, interns and residents, and various paramedical specialties. These programs contribute to the quality of patient care within an institution and are necessary to meet the community's needs for medical and paramedical personnel. It is recognized that the costs of such educational activities should be borne by the community. However, many communities have not assumed responsibility for financing these programs and it is necessary that support be provided by those purchasing health care. Until communities undertake to bear these costs, the program will participate appropriately in the support of these activities. Although the intent of the program is to share in the support of educational activities customarily or traditionally carried on by providers in conjunction with their operations, it is not intended that this program should participate in increased costs resulting from redistribution of costs from educational institutions or units to patient care institutions or units.

(d) Activities not within the scope of this principle. The costs of the following activities are not within the scope of this principle but are recognized as normal operating costs and are reimbursed in accordance with applicable principles—

(1) Orientation and on-the-job training;

(2) Part-time education for bona fide employees at properly accredited academic or technical institutions (including other providers) devoted to undergraduate or graduate work;

(3) Costs, including associated travel expense, or sending employees to educational seminars and workshops that increase the quality of medical care or operating efficiency of the provider;

(4) Maintenance of a medical library;

(5) Training of a patient or patient's family in the use of medical appliances;

(6) Clinical training of students not enrolled in an approved education program operated by the provider; and

(7) Other activities that do not involve the actual operation of an approved education program including the costs of interns and residents in anesthesiology who are employed to replace anesthetists.

(e) Approved programs. Recognized professional and paramedical educational training programs now being conducted by provider institutions, and their approving bodies, include the following:

(1) Cyto-technology. Committee on Allied Health, Education, and Accreditation in collaboration with the Board of Schools of Medical Technology, American Society of Clinical Pathologists.

(2) Dietetic internships. The American Dietetic Association.
(3) Hospital administration residencies. Accrediting Commission on Education in Health Services Administration.

(4) Inhalation therapy. Committee on Allied Health, Education, and Accreditation in collaboration with the Board of Schools of Inhalation Therapy.

(5) Medical records. Committee on Allied Health, Education, and Accreditation in collaboration with the Committee on Education and Registration of the American Association of Medical Records Librarians.

(6) Medical technology. Committee on Allied Health, Education, and Accreditation in collaboration with the Board of Schools of Medical Technology, American Society of Clinical Pathologists.


(8) Professional nursing. Approved by the respective State approving authorities. Reported for the United States by the National League for Nursing.


(10) Occupational Therapy. American Society of Hospital Pharmacists.


(12) Physical therapy. Committee on Allied Health, Education, and Accreditation in collaboration with the American College of Radiology.

(13) X-ray technology. Committee on Allied Health, Education, and Accreditation in collaboration with the American Board of Radiology.

(f) Other educational programs. There may also be other educational programs not included in the foregoing in which a provider institution is engaged. Appropriate consideration will be given by the intermediary and HCFA to the costs incurred for those activities that come within the purview of the principle when determining the allowable costs for apportionment under the Medicare program.

(g) Calculating net cost. Net costs of approved educational activities are determined by deducting, from a provider’s total costs of these activities, revenues it receives from tuition. For this purpose, a provider’s total costs include trainee stipends, compensation of teachers, and other direct and indirect costs of the activities as determined under the Medicare cost-finding principles in §413.24.

(h) Medicare+Choice organizations. (1) Effective January 1, 1999, Medicare+Choice organizations may receive direct graduate medical education payments for the time that residents spend in nonhospital provider settings such as freestanding clinics, nursing homes, and physicians’ offices in connection with approved programs.

(2) Medicare+Choice organizations may receive direct graduate medical education payments if all of the following conditions are met:

(i) The resident spends his or her time in patient care activities.

(ii) The Medicare+Choice organization incurs “all or substantially all” of the costs for the training program in the nonhospital setting as defined in §413.86(b).

(iii) There is a written agreement between the Medicare+Choice organization and the nonhospital site that indicates the Medicare+Choice organization will incur the costs of the resident’s salary and fringe benefits and provide reasonable compensation to the nonhospital site for teaching activities.

(3) A Medicare+Choice organization’s allowable direct graduate medical education costs, subject to the redistribution and community support principles in §413.85(c), consist of—

(i) Residents’ salaries and fringe benefits (including travel and lodging where applicable); and

(ii) Reasonable compensation to the nonhospital site for teaching activities.
§ 413.86 Direct graduate medical education payments.

(a) Statutory basis and scope—(1) Basis. This section implements section 1886(h) of the Act by establishing the methodology for Medicare payment of the cost of direct graduate medical educational activities.

(2) Scope. This section applies to Medicare payments to hospitals and hospital-based providers for the costs of approved residency programs in medicine, osteopathy, dentistry, and podiatry for cost reporting periods beginning on or after July 1, 1985.

(b) Definitions. For purposes of this section, the following definitions apply:

Affiliated group means—

(1) Two or more hospitals located in the same urban or rural area (as those terms are defined in §412.62(f) of this subchapter) or in contiguous areas if individual residents work at each of the hospitals during the course of the program; or

(2) If the hospitals are not located in the same or a contiguous urban or rural area, the hospitals are jointly listed—

(i) As the sponsor, primary clinical site or major participating institution for one or more of the programs as these terms are used in Graduate Medical Education Directory, 1997-1998; or

(ii) As the sponsor or under "affiliations and outside rotations" for one or more programs in operation in Opportunities, Directory of Osteopathic Postdoctoral Education Programs.

(3) The hospitals are under common ownership.

All or substantially all of the costs for the training program in the nonhospital setting means the residents' salaries and fringe benefits (including travel and lodging where applicable) and the portion of the cost of teaching physicians' salaries and fringe benefits attributable to direct graduate medical education.

Approved geriatric program means a fellowship program of one or more years in length that is approved by one of the national organizations listed in §415.152 of this chapter under that respective organization's criteria for geriatric fellowship programs.

Approved medical residency program means a program that meets one of the following criteria:

(1) Is approved by one of the national organizations listed in §415.152 of this chapter.

(2) May count towards certification of the participant in a specialty or subspecialty listed in the current edition of either of the following publications:

(i) The Directory of Graduate Medical Education Programs published by the American Medical Association, and available from American Medical Association, Department of Directories and Publications, 515 North State Street, Chicago, Illinois 60610; or

(ii) The Annual Report and Reference Handbook published by the American Board of Medical Specialties, and available from American Board of Medical Specialties, One Rotary Center, suite 805, Evanston, Illinois 60201.

(3) Is approved by the Accreditation Council For Graduate Medical Education (ACGME) as a fellowship program in geriatric medicine.

(4) Is a program that would be accredited except for the accrediting agency's reliance upon an accreditation standard that requires an entity related to the training of medical residents.

(4) The direct graduate medical education payment is equal to the product of—

(i) The lower of—

(A) The Medicare+Choice organization's allowable direct graduate medical education costs per resident as defined in paragraph (h)(3) of this section; or

(B) The national average per resident amount; and

(ii) Medicare's share, which is equal to the ratio of the number of Medicare beneficiaries enrolled to the total number of individuals enrolled in the Medicare+Choice organization.

(5) Direct graduate medical education payments made to Medicare+Choice organizations under this section are made from the Federal Supplementary Medical Insurance Trust Fund.

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to perform an induced abortion or require, provide, or refer for training in the performance of induced abortions, or make arrangements for such training, regardless of whether the standard provides exceptions or exemptions.

Base period means a cost reporting period that began on or after October 1, 1983 but before October 1, 1984.

CPI–U stands for the Consumer Price Index for All Urban Consumers as compiled by the Bureau of Labor Statistics.

Foreign medical graduate means a resident who is not a graduate of a medical, osteopathy, dental, or podiatry school, respectively, accredited or approved as meeting the standards necessary for accreditation by one of the following organizations:

(1) The Liaison Committee on Medical Education of the American Medical Association.


(3) The Commission on Dental Accreditation.

(4) The Council on Podiatric Medical Education.

FMGEMS stands for the Foreign Medical Graduate Examination in the Medical Sciences (Part I and Part II).

FTE stands for full-time equivalent.

Medicare patient load means, with respect to a hospital’s cost reporting period, the total number of hospital inpatient days during the cost reporting period that are attributable to patients for whom payment is made under Medicare Part A divided by total hospital inpatient days. In calculating inpatient days, inpatient days in any distinct part of the hospital furnishing a hospital level of care are included and nursery days are excluded.

Primary care resident is a resident enrolled in an approved medical residency program in family medicine, general internal medicine, general pediatrics, preventive medicine, geriatric medicine or osteopathic general practice.

Resident means an intern, resident, or fellow who participates in an approved medical residency program, including programs in osteopathy, dentistry, and podiatry, as required in order to become certified by the appropriate specialty board.

Rural track FTE limitation means the maximum number of residents (as specified in paragraph (g)(11) of this section) training in a rural track residency program that an urban hospital may include in its FTE count and that is in addition to the number of FTE residents already included in the hospital’s FTE cap.

Rural track or integrated rural track means an approved medical residency training program established by an urban hospital in which residents train for a portion of the program at the urban hospital and then rotate for a portion of the program to a rural hospital(s) or a rural nonhospital site(s).

(c) Payment for graduate medical education costs—General rule. Beginning with cost reporting periods starting on or after July 1, 1985, hospitals, including hospital-based providers, are paid for the costs of approved graduate medical education programs as described in paragraph (d) through (h) of this section.

(d) Calculating payment for graduate medical education costs. A hospital’s Medicare payment for the costs of an approved residency program is calculated as follows:

(1) Step one. The hospital’s updated per resident amount (as determined under paragraph (e) of this section) is multiplied by the actual number of FTE residents (as determined under paragraph (g) of this section). This result is the aggregate approved amount for the cost reporting period.

(2) Step two. The product derived in step one is multiplied by the hospital’s Medicare patient load.

(3) Step Three. For portions of cost reporting periods occurring on or after January 1, 1998, the product derived in step one is multiplied by the proportion of the hospital’s inpatient days attributable to individuals who are enrolled under a risk-sharing contract with an eligible organization under section 1876 of the Act and who are entitled to Medicare Part A or with a Medicare+Choice organization under Title XVIII, Part C of the Act. This amount is multiplied by an applicable payment percentage equal to—

(i) 20 percent for 1998;

(ii) 40 percent for 1999;

(iii) 60 percent in 2000.
(iv) 80 percent in 2001; and
(v) 100 percent in 2002 and subsequent years.

(4) Step four. Effective for cost reporting periods beginning on or after January 1, 2000, the product derived from step three is reduced in accordance with the provisions of §413.87(f).

(5) Step five. (i) For portions of cost reporting periods beginning on or after January 1, 1998 and before January 1, 2000, add steps two and three.

(ii) Effective for portions of cost reporting periods beginning on or after January 1, 2000, add the results of steps two and four.

(6) Step six. The product derived in step two is apportioned between Part A and Part B of Medicare based on the ratio of Medicare's share of reasonable costs excluding graduate medical education costs attributable to each part as determined through the Medicare cost report.

(e) Determining per resident amounts for the base period—(1) For the base period. (i) Except as provided in paragraph (e)(4) of this section, the intermediary determines a base-period per resident amount for each hospital as follows:

(A) Determine the allowable graduate medical education costs for the cost reporting period beginning on or after October 1, 1983 but before October 1, 1984. In determining these costs, graduate medical education costs allocated to the nursery cost center, research and other nonreimbursable cost centers, and hospital-based providers that are not participating in Medicare are excluded and graduate medical education costs allocated to distinct-part hospital units and hospital-based providers that participate in Medicare are included.

(B) Divide the costs calculated in paragraph (e)(1)(i)(A) of this section by the average number of FTE residents working in all areas of the hospital complex (including those areas whose costs were excluded under paragraph (e)(1)(i)(A) of this section) for its cost reporting period beginning on or after October 1, 1983 but before October 1, 1984.

(ii) In determining the base-period per resident amount under paragraph (e)(1)(i) of this section, the intermediary—

(A) Verifies the hospital's base-period graduate medical education costs and the hospital's average number of FTE residents;

(B) Excludes from the base-period graduate medical education costs any nonallowable or misclassified costs, including those previously allowed under §412.113(b)(3) of this chapter; and

(C) Upon a hospital's request, includes graduate medical education costs that were misclassified as operating costs during the hospital's prospective payment base year and were not allowable under §412.113(b)(3) of this chapter during the graduate medical education base period. These costs may be included only if the hospital requests an adjustment of its prospective payment hospital-specific rate or target amount as described in paragraph (k)(1) of this section.

(iii) If the hospital's cost report for its GME base period is no longer subject to reopening under §405.1885 of this chapter, the intermediary may modify the hospital's base-period costs solely for purposes of computing the per resident amount.

(iv) If the intermediary modifies a hospital's base-period graduate medical education costs as described in paragraph (e)(1)(iii)(B) of this section, the hospital may request an adjustment of its prospective payment hospital-specific rate or target amount as described in paragraph (k)(1) of this section.

(v) The intermediary notifies each hospital that either had direct graduate medical education costs or received indirect education payment in its cost reporting period beginning on or after October 1, 1984 and before October 1, 1985 of its base-period average per resident amount. A hospital may appeal this amount within 180 days of the date of that notice.

(2) For cost reporting periods beginning on or after July 1, 1985 and before July 1, 1986. For cost reporting periods beginning on or after July 1, 1985 and before July 1, 1986, a hospital's base-period per resident amount is adjusted as follows:

(i) If a hospital's base period began on or after October 1, 1983 and before July 1, 1984, the amount is adjusted by the percentage change in the CPI-U-
that occurred between the hospital's base period and the first cost reporting period to which the provisions of this section apply. The adjusted amount is then increased by one percent.

(ii) If a hospital's base period began on or after July 1, 1984 and before October 1, 1984, the amount is increased by one percent.

(3) For cost reporting periods beginning on or after July 1, 1986. Subject to the provisions of paragraph (e)(4) of this section, for cost reporting periods beginning on or after July 1, 1986, a hospital's base-period per resident amount is adjusted as follows:

(i) Except as provided in paragraph (e)(3)(ii) of this section, each hospital's per resident amount for the previous cost reporting period will be adjusted for projected change in the CPI-U for the 12-month cost reporting period. This adjustment is subject to revision during the settlement of the cost report to reflect actual changes in the CPI-U that occurred during the cost reporting period.

(ii) For cost reporting periods beginning on or after October 1, 1993 through September 30, 1995, each hospital's per resident amount for the previous cost reporting period will not be adjusted for any resident FTEs who are not either a primary care resident or an obstetrics and gynecology resident.

(4) For cost reporting periods beginning on or after October 1, 2000 and ending on or before September 30, 2005. For cost reporting periods beginning on or after October 1, 2000 and ending on or before September 30, 2005, a hospital’s per resident amount for each fiscal year is adjusted in accordance with the following provisions:

(i) General provisions. For purposes of §413.86(e)(4)—

(A) Weighted average per resident amount. The weighted average per resident amount is established as follows:

(1) Using data from hospitals' cost reporting periods ending during FY 1997, HCFA calculates each hospital's single per resident amount by adding each hospital's primary care and non-primary care per resident amounts, weighted by its respective FTEs, and dividing by the sum of the FTEs for primary care and non-primary care residents.

(2) Each hospital’s single per resident amount calculated under paragraph (e)(4)(i)(A) of this section is standardized by the 1999 geographic adjustment factor for the physician fee schedule area (as determined under §414.26 of this chapter) in which the hospital is located.

(3) HCFA calculates an average of all hospitals’ standardized per resident amounts that are determined under paragraph (e)(4)(i)(A) of this section. The resulting amount is the weighted average per resident amount.

(B) Primary care/obstetrics and gynecology and non-primary care per resident amounts. A hospital’s per resident amount is an amount inclusive of any CPI-U adjustments that the hospital may have received since the hospital’s base year, including any CPI-U adjustments the hospital may have received because the hospital trains primary care/obstetrics and gynecology residents and non-primary care residents as specified under paragraph (e)(3)(ii) of this section.

(ii) Adjustment beginning in FY 2001 and ending in FY 2005. For cost reporting periods beginning on or after October 1, 2000 and ending on or before September 30, 2005, a hospital’s per resident amount is adjusted in accordance with paragraphs (e)(4)(ii)(A) through (e)(4)(ii)(C) of this section, in that order:

(A) Updating the weighted average per resident amount for inflation. The weighted average per resident amount (as determined under paragraph (e)(4)(ii)(A) of this section) is updated by the estimated percentage increase in the CPI-U during the period beginning with the month that represents the midpoint of the cost reporting periods ending during FY 1997 (that is, October 1, 1996) and ending with the midpoint of the hospital’s cost reporting period that begins in FY 2001.

(B) Adjusting for locality. The updated weighted average per resident amount determined under paragraph (e)(4)(ii)(A) of this section (the national average per resident amount) is adjusted for the locality of each hospital by multiplying the national average per resident amount by the 1999 geographic adjustment factor for the physician fee schedule area in which each hospital is located.
hospital is located, established in accordance with §414.26 of this subchapter.

(C) Determining necessary revisions to the per resident amount. The locality-adjusted national average per resident amount, as calculated in accordance with paragraph (e)(4)(ii)(B) of this section, is compared to the hospital's per resident amount. If the hospital's per resident amount is greater than 140 percent of the locality-adjusted national average per resident amount, then, subject to the provision stated in paragraph (e)(4)(ii)(C) of this section, the hospital's per resident amount is revised, if appropriate, according to the following three categories:

(1) Floor. For cost reporting periods beginning on or after October 1, 2000 and on or before September 30, 2001, the hospital's per resident amount would otherwise be less than 70 percent of the locality-adjusted national average per resident amount for FY 2001 (as determined under paragraph (e)(4)(ii)(B) of this section), the per resident amount is equal to 70 percent of the locality-adjusted national average per resident amount for FY 2001.

For subsequent cost reporting periods, the hospital's per resident amount is updated using the methodology specified under paragraph (e)(3)(i) of this section.

(2) Ceiling. If the hospital's per resident amount is greater than 140 percent of the locality-adjusted national average per resident amount for FY 2001, the per resident amount is adjusted as follows for FY 2001 through FY 2005:

(i) FY 2001. For cost reporting periods beginning on or after October 1, 2000 and on or before September 30, 2001, if the hospital's FY 2000 per resident amount exceeds 140 percent of the FY 2001 locality-adjusted national average per resident amount (as calculated under paragraph (e)(4)(ii)(B) of this section), then, subject to the provision stated in paragraph (e)(4)(ii)(C) of this section, the hospital's per resident amount is frozen at the FY 2000 per resident amount and is not updated for FY 2002 by the CPI-U factor.

(ii) FY 2002. For cost reporting periods beginning on or after October 1, 2001, and on or before September 30, 2002, if the hospital's FY 2001 per resident amount exceeds 140 percent of the FY 2002 locality-adjusted national average per resident amount, then, subject to the provision stated in paragraph (e)(4)(ii)(C) of this section, the hospital's per resident amount is frozen at the FY 2001 per resident amount and is not updated for FY 2002 by the CPI-U factor.

(iii) FY 2003 through FY 2005. For cost reporting periods beginning on or after October 1, 2002 and on or before September 30, 2005, if the hospital's per resident amount for the previous cost reporting period is greater than 140 percent of the locality-adjusted national average per resident amount for the same previous cost reporting period (for example, for cost reporting periods beginning in FY 2003, compare the hospital's per resident amount from the FY 2002 cost report to the hospital's locality-adjusted national average per resident amount from FY 2002), then, subject to the provision stated in paragraph (e)(4)(ii)(C) of this section, the hospital's per resident amount is adjusted using the methodology specified in paragraph (e)(3)(i) of this section, except that the CPI-U applied for a 12-month period is reduced (but not below zero) by 2 percentage points.

(iv) General rule for hospitals that exceed the ceiling. For cost reporting periods beginning on or after October 1, 2000 and on or before September 30, 2005, if a hospital's per resident amount exceeds 140 percent of the hospital's locality-adjusted national average per resident amount and it is adjusted under any of the criteria (e)(4)(ii)(C)(2)(i) through (iii) of this section, the current year per resident amount cannot be reduced below 140 percent of the locality-adjusted national average per resident amount.

(3) Per resident amounts greater than or equal to the floor and less than or equal to the ceiling. For cost reporting periods beginning on or after October 1, 2000 and on or before September 30, 2005, if a hospital's per resident amount is greater than or equal to 70 percent and less than or equal to 140 percent of the hospital's locality-adjusted national average per resident amount, for each respective fiscal year, the hospital's per resident amount is frozen at the FY 2000 per resident amount and is not updated for FY 2001 by the CPI-U factor.

(4) Per resident amounts less than 70 percent of the locality-adjusted national average per resident amount. For subsequent cost reporting periods, the hospital's per resident amount is updated using the methodology specified in paragraph (e)(3)(i) of this section.

(5) Exceptions—(i) Base period for certain hospitals. If a hospital did not have any approved medical residency training programs or did not participate in...
Medicare during the base period, but either condition changes in a cost reporting period beginning on or after July 1, 1985, the intermediary establishes a per resident amount for the hospital using the information from the first cost reporting period during which the hospital participates in Medicare and the residents are on duty during the first month of that period. Any graduate medical education program costs incurred by the hospital before that cost reporting period are reimbursed on a reasonable cost basis. The per resident amount is based on the lower of the following:

(A) The hospital’s actual costs, incurred in connection with the graduate medical education program for the hospital’s first cost reporting period in which residents were on duty during the first month of the cost reporting period.

(B) The weighted mean value of per resident amounts of hospitals located in the same geographic wage area, as that term is used in the prospective payment system under part 412 of this chapter, for cost reporting periods beginning in the same fiscal years. If there are fewer than three amounts that can be used to calculate the weighted mean value, the calculation of the per resident amounts includes all hospitals in the hospital’s region as that term is used in §412.62(f)(1)(i) of this chapter.

(ii) Short or long base-period cost reporting periods. If a hospital’s base-period cost reporting period reflects graduate medical education costs for a period that is shorter than 50 weeks or longer than 54 weeks, the intermediary converts the allowable costs for the base period into a daily figure. The daily figure is then multiplied by 365 or 366, as appropriate, to derive the approved per resident amount for a 12-month base-period cost reporting period. If a hospital has two cost reporting periods beginning in the base period, the later period serves as the base-period cost reporting period.

(iii) Short or long cost reporting periods beginning on or after July 1, 1985. If a hospital’s cost reporting period is shorter than 50 weeks or longer than 54 weeks, the hospital’s intermediary should contact HCFA Central Office to receive a special CPI-U adjustment factor.

(iv) Effective October 1, 2000, the per resident amounts established under paragraphs (e)(5)(i) through (iii) of this section are subject to the provisions of paragraph (e)(4) of this section.

(f) Determining the total number of FTE residents. Subject to the weighting factors in paragraphs (g) and (h) of this section, the count of FTE residents is determined as follows:

1. Residents in an approved program working in all areas of the hospital complex may be counted.

2. No individual may be counted as more than one FTE. Except as provided in paragraphs (f)(1)(3) and (4) of this section, if a resident spends time in more than one hospital or, in a nonprovider setting, the resident counts as partial FTE based on the proportion of time worked at the hospital to the total time worked. A part-time resident counts as a partial FTE based on the proportion of allowable time worked compared to the total time necessary to fill a full-time internship or residency slot.

3. On or after July 1, 1987 and for portions of cost reporting periods occurring before January 1, 1999, the time residents spend in nonprovider settings such as freestanding clinics, nursing homes, and physicians’ offices in connection with approved programs is not excluded in determining the number of FTE residents in the calculation of a hospital’s resident count if the following conditions are met—

(i) The resident spends his or her time in patient care activities.

(ii) There is a written agreement between the hospital and the outside entity that states that the resident’s compensation for training time spent outside of the hospital setting is to be paid by the hospital.

4. For portions of cost reporting periods occurring on or after January 1, 1999, the time residents spend in nonprovider settings such as freestanding clinics, nursing homes, and physicians’ offices in connection with approved programs may be included in determining the number of FTE residents in the calculation of a hospital’s resident count if the following conditions are met—
(i) The resident spends his or her time in patient care activities.

(ii) The written agreement between the hospital and the nonhospital site must indicate that the hospital will incur the cost of the resident’s salary and fringe benefits while the resident is training in the nonhospital site and the hospital is providing reasonable compensation to the nonhospital site for supervisory teaching activities. The agreement must indicate the compensation the hospital is providing to the nonhospital site for supervisory teaching activities.

(iii) The hospital must incur all or substantially all of the costs for the training program in the nonhospital setting in accordance with the definition in paragraph (b) of this section.

(g) Determining the weighted number of FTE residents. Subject to the provisions in paragraph (h) of this section, HCFA determines a hospital’s number of FTE residents by applying a weighting factor to each resident and then summing the resulting numbers that represent each resident. The weighting factor is determined as follows:

(1) Generally, for purposes of this section, effective July 1, 1995, an initial residency period is defined as the minimum number of years required for board eligibility. Prior to July 1, 1995, the initial residency period equals the minimum number of years required for board eligibility in a specialty or subspecialty plus 1 year. An initial residency period may not exceed 5 years in order to be counted toward determining FTE status except in the case of fellows in an approved geriatric program whose initial residency period may last up to 2 additional years. Effective June 1, 2000, for residency programs that began before, on, or after November 29, 1999, the period of board eligibility and the initial residency period for a resident in an approved child neurology program is the period of board eligibility for pediatrics plus 2 years. Effective August 10, 1993, residents or fellows in an approved preventive medicine residency or fellowship program also may be counted as a full FTE resident for up to 2 additional years beyond the initial residency period limitation. For combined residency programs, an initial residency period is defined as the time required for individual certification in the longer of the programs. If the resident is enrolled in a combined medical residency training program in which all of the individual programs (that are combined) are for training primary care residents (as defined in paragraph (b) of this section) or obstetrics and gynecology residents, the initial residency period is the time required for individual certification in the longer of the programs plus 1 year.

(ii) For residency programs other than those specified in paragraphs (g)(1)(ii) and (g)(1)(iii) of this section, the initial residency period is the minimum number of years of formal training necessary to satisfy the requirements for initial board eligibility in the particular specialty for which the resident is training, as specified in the most recently published edition of the Graduate Medical Education Directory.

(iii) For residency programs in osteopathy, dentistry, and podiatry, the minimum requirement for certification in a specialty or subspecialty is the minimum number of years of formal training necessary to satisfy the requirements of the appropriate approving body listed in §415.152 of this chapter.

(iv) For residency programs in geriatric medicine, accredited by the appropriate approving body listed in 415.152 of this chapter, these programs are considered approved programs on the later of—

(A) The starting date of the program within a hospital; or

(B) The hospital’s cost reporting periods beginning on or after July 1, 1985.

(iv) The time spent in residency programs that do not lead to certification in a specialty or subspecialty, but that otherwise meet the definition of approved programs, as described in paragraph (b) of this section, is counted toward the initial residency period limitation.

(2) If the resident is in an initial residency period, the weighting factor is one.

(3) If the resident is not in an initial residency period, the weighting factor is 1.00 during the period beginning on or after July 1, 1985 and before July 1, 1986, .75 during the period beginning on or after July 1, 1986 and before July 1,
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1987 and is .50 thereafter without re-
gard to the hospital's cost reporting
period.

(4) For purposes of determining di-
rect graduate medical education pay-
ment, for cost reporting periods begin-
ing on or after October 1, 1997, a hos-
pital's unweighted FTE count for resi-
dents in allopathic and osteopathic
medicine may not exceed the hospital's
unweighted FTE count (or, effective for
cost reporting periods beginning on or
after April 1, 2000, 130 percent of the
unweighted FTE count for a hospital
located in a rural area) for these resi-
dents for the most recent cost report-
ing period ending on or before Decem-
ber 31, 1996. If the hospital's number of
FTE residents in a cost reporting pe-
riod beginning on or after October 1,
1997, exceeds the limit described in this
paragraph (g), the hospital's weighted
FTE count (before application of the
limit) will be reduced in the same pro-
portion that the number of FTE resi-
dents for that cost reporting period ex-
ceds the number of FTE residents for
the most recent cost reporting period
ending on or before December 31, 1996.
Hospitals that are part of the same af-
iliated group may elect to apply the
limit on an aggregate basis. The fiscal
intermediary may make appropriate
modifications to apply the provisions
of this paragraph (g)(4) based on the
equivalent of a 12-month cost reporting
period.

(5) For purposes of determining di-
rect graduate medical education pay-
ment, for the hospital's first cost re-
porting period beginning on or after
October 1, 1997, the hospital's weighted
FTE count is equal to the average of the
weighted FTE count for the pay-
ment year cost reporting period and
the preceding cost reporting period.
For cost reporting periods beginning on
or after October 1, 1998, the hospital's
weighted FTE count is equal to the av-
erage of the weighted FTE count for the
payment year cost reporting period
and the preceding two cost reporting
periods. The fiscal intermediary may
make appropriate modifications to
apply the provisions of this paragraph
based on the equivalent of 12-month
cost reporting periods. If a hospital
qualifies for an adjustment to the limit
established under paragraph (g)(4) of
this section for new medical residency
programs created under paragraph
(g)(6) of this section, the count of resi-
dents participating in new medical
residency training programs above the
number included in the hospital's FTE
count for the cost reporting period end-
ing during calendar year 1996 is added
after applying the averaging rules in
this paragraph for a period of years.
Residents participating in new medical
residency training programs are in-
cluded in the hospital's FTE count be-
fore applying the averaging rules after
the period of years has expired. For
purposes of this paragraph, the period
of years equals the minimum accred-
dited length for the type of program.
The period of years begins when the
first resident begins training.

(6) If a hospital establishes a new
medical residency training program as
defined in paragraph (g)(9) of this sec-
tion on or after January 1, 1995, the
hospital's FTE cap described under
paragraph (g)(4) of this section may be
adjusted as follows:

(i) If a hospital had no allopathic or
osteopathic residents in its most re-
cent cost reporting period ending on or
before December 31, 1996, and it estab-
lishes a new medical residency training
program on or after January 1, 1995,
the hospital's unweighted FTE resident
cap under paragraph (g)(4) of this sec-
tion may be adjusted based on the
product of the highest number of resi-
dents in any program year during the
third year of the first program's exist-
ence and the number of years in
which residents are expected to com-
plete the program based on the min-
imum accredited length for the type of
program. The adjustment to the cap
may not exceed the number of accred-
dited slots available to the hospital for
the new program.

(A) If the residents are spending an
entire program year (or years) at one
hospital and the remainder of the pro-
gram at another hospital, the adjust-
ment to each respective hospital's cap
is equal to the product of the highest
number of residents in any program
year during the third year of the first
program's existence and the number of
years the residents are training at each
respective hospital.
(B) Prior to the implementation of the hospital’s adjustment to its FTE cap beginning with the fourth year of the hospital’s residency program, the hospital’s cap may be adjusted during each of the first 3 years of the hospital’s new residency program using the actual number of residents participating in the new program. The adjustment may not exceed the number of accredited slots available to the hospital for each program year.

(C) Except for rural hospitals, the cap will not be adjusted for new programs established more than 3 years after the first program begins training residents.

(D) An urban hospital that qualifies for an adjustment to its FTE cap under paragraph (g)(6)(i) of this section is not permitted to be part of an affiliated group for purposes of establishing an aggregate FTE cap.

(E) A rural hospital that qualifies for an adjustment to its FTE cap under paragraph (g)(6)(i) of this section is permitted to be part of an affiliated group for purposes of establishing an aggregate FTE cap.

(ii) If a hospital with allopathic or osteopathic residents in its most recent cost reporting period ending on or before December 31, 1996, is located in a rural area (or other hospitals located in rural areas that added residents under paragraph (g)(6)(i) of this section), the hospital’s unweighted FTE limit may be adjusted in the same manner described in paragraph (g)(6)(ii) of this section to reflect the increase for residents in the new medical residency training programs established after August 5, 1997. For these hospitals, the limit will be adjusted for additional new programs but not for expansions of existing or previously existing programs.

(iv) A hospital seeking an adjustment to the limit on its unweighted resident count policy must provide documentation to its fiscal intermediary justifying the adjustment.

(7) A hospital that began construction of its facility prior to August 5, 1997, and sponsored new medical residency training programs on or after January 1, 1995 and on or before August 5, 1997, that either received initial accreditation by the appropriate accrediting body or temporarily trained residents at another hospital(s) until the facility was completed, may receive an adjustment to its FTE cap.

(i) The newly constructed hospital’s FTE cap is equal to the lesser of:

(A) The product of the highest number of residents in any program year during the third year of the newly established program and the number of years in which residents are expected to complete the programs based on the minimum accredited length for the type of program.

(B) Prior to the implementation of the hospital’s adjustment to its FTE cap beginning with the fourth year of the hospital’s residency program, the hospital’s cap may be adjusted during each of the first 3 years of the hospital’s new residency program, using the actual number of residents in the new programs. The adjustment may not exceed the number of accredited slots available to the hospital for each program year.

(iii) If a hospital with allopathic or osteopathic residents in its most recent cost reporting period ending on or before December 31, 1996, is located in a rural area (or other hospitals located in rural areas that added residents under paragraph (g)(6)(i) of this section), the hospital’s unweighted FTE limit may be adjusted in the same manner described in paragraph (g)(6)(ii) of this section to reflect the increase for residents in the new medical residency training programs established after August 5, 1997. For these hospitals, the limit will be adjusted for additional new programs but not for expansions of existing or previously existing programs.

(iv) A hospital seeking an adjustment to the limit on its unweighted resident count policy must provide documentation to its fiscal intermediary justifying the adjustment.

(7) A hospital that began construction of its facility prior to August 5, 1997, and sponsored new medical residency training programs on or after January 1, 1995 and on or before August 5, 1997, that either received initial accreditation by the appropriate accrediting body or temporarily trained residents at another hospital(s) until the facility was completed, may receive an adjustment to its FTE cap.

(i) The newly constructed hospital’s FTE cap is equal to the lesser of:

(A) The product of the highest number of residents in any program year during the third year of the newly established program and the number of years in which residents are expected to complete the programs based on the minimum accredited length for each type of program; or

(B) The number of accredited slots available to the hospital for each year of the programs.

(ii) If the new medical residency training programs sponsored by the
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newly constructed hospital have been in existence for 3 years or more by the time the residents begin training at the newly constructed hospital, the newly constructed hospital’s cap will be based on the number of residents training in the third year of the programs begun at the temporary training site.

(iii) If the new medical residency training programs sponsored by the newly constructed hospital have been in existence for less than 3 years by the time the residents begin training at the newly constructed hospital, the newly constructed hospital’s cap will be based on the number of residents training at the newly constructed hospital in the third year of the programs (including the years at the temporary training site).

(iv) A hospital that qualifies for an adjustment to its FTE cap under paragraph (g)(7) of this section may be part of an affiliated group for purposes of establishing an aggregate FTE cap.

(v) The provisions of this paragraph (g)(7) are applicable during portions of cost reporting periods occurring on or after October 1, 1999.

(8) A hospital may receive a temporary adjustment to its FTE cap to reflect residents added because of another hospital’s closure if the hospital meets the following criteria:

(i) The hospital is training additional residents from a hospital that closed on or after July 1, 1996.

(ii) No later than 60 days after the hospital begins to train the residents, the hospital submits a request to its fiscal intermediary for a temporary adjustment to its FTE cap, documents that the hospital is eligible for this temporary adjustment by identifying the residents who have come from the closed hospital and have caused the hospital to exceed its cap, and specifies the length of time the adjustment is needed.

(iii) For purposes of paragraph (g)(8) of this section, “closure” means the hospital terminates its Medicare agreement under the provisions of §489.52 of this chapter.

(9) Effective for cost reporting periods beginning on or after November 29, 1999, a hospital may receive an adjustment to its FTE cap of up to three additional resident FTEs, if the hospital meets the following criteria:

(i) The additional residents are residents of a primary care program that would have been counted by the hospital as residents for purposes of the hospital’s FTE cap but for the fact that the additional residents were on maternity or disability leave or a similar approved leave of absence during the hospital’s most recent cost reporting period ending on or before December 31, 1996;

(ii) The leave of absence was approved by the residency program director to allow the residents to be absent from the program and return to the program after the leave of absence; and

(iii) No later than 6 months after August 1, 2000, the hospital submits to the fiscal intermediary a request for an adjustment to its FTE cap, and provides contemporaneous documentation of the approval of the leave of absence by the residency director, specific to each additional resident that is to be counted for purposes of the adjustment.

(10) For cost reporting periods beginning on or after October 1, 1997, a non-Veterans Affairs (VA) hospital may receive a temporary adjustment to its FTE cap to reflect residents who had previously trained at a VA hospital and were subsequently transferred to the non-VA hospital, if that hospital meets the following criteria:

(i) The transferred residents had been training previously at a VA hospital in a program that would have lost its accreditation by the ACGME if the residents continued to train at the VA hospital;

(ii) The residents were transferred to the hospital from the VA hospital on or after January 1, 1997, and before July 31, 1998; and

(iii) The hospital submits a request to its fiscal intermediary for a temporary adjustment to its FTE cap, documents that it is eligible for this temporary adjustment by identifying the residents who have come from the VA hospital, and specifies the length of time those residents will be trained at the hospital.

(11) For cost reporting periods beginning on or after April 1, 2000, an urban
hospital that establishes a new residency program, or has an existing residency program, with a rural track (or an integrated rural track) may include in its FTE count residents in those rural tracks, in addition to the residents subject to its FTE cap specified under paragraph (g)(4) of this section. An urban hospital with a rural track residency program may count residents in those rural tracks up to a rural track FTE limitation if the hospital complies with the conditions specified in paragraphs (g)(11)(i) through (g)(11)(vi) of this section.

(i) If an urban hospital rotates residents in the rural track program to a rural hospital(s) for at least two-thirds of the duration of the program, the urban hospital may include those residents in its FTE count for the time the rural track residents spend at the urban hospital. The urban hospital may include in its FTE count those residents in the rural track training at the urban hospital, not to exceed its rural track FTE limitation, determined as follows:

(A) For the first 3 years of the rural track’s existence, the rural track FTE limitation for each urban hospital will be the actual number of FTE residents training in the rural track at the urban hospital.

(B) Beginning with the fourth year of the rural track’s existence, the rural track FTE limitation is equal to the product of—

(1) The highest number of residents in any program year who, during the third year of the rural track’s existence, are training in the rural track at—

(i) The urban hospital and are designated at the beginning of their training to be rotated to a rural nonhospital site(s) for at least two-thirds of the duration of the program; and

(ii) The rural nonhospital site(s); and

(2) The number of years in which the residents are expected to complete each program based on the minimum accredited length for the type of program.

(ii) If an urban hospital rotates residents in the rural track program to a rural hospital(s) for periods of time that are less than two-thirds of the duration of the program, the urban hospital may not include those residents in its FTE count (if the urban hospital’s FTE count exceeds that hospital’s FTE cap), nor may the urban hospital include those residents when calculating its rural track FTE limitation.

(iv) If an urban hospital rotates residents in the rural track program to a rural nonhospital site(s) for periods of time that are less than two-thirds of the duration of the program, the urban hospital may include those residents in its FTE count, subject to the requirements under paragraph (f)(4) of this section. The urban hospital may include in its FTE count those residents in the rural track, not to exceed its rural track FTE limitation, determined as follows:

(A) For the first 3 years of the rural track’s existence, the rural track FTE limitation for the urban hospital will be the actual number of FTE residents training in the rural track at the rural nonhospital site(s).
(B) Beginning with the fourth year of the rural track’s existence, the rural track FTE limitation is equal to the product of—

(1) The highest number of residents in any program year who, during the third year of the rural track’s existence, are training in the rural track at the rural nonhospital site(s) or are designated at the beginning of their training to be rotated to the rural nonhospital site(s) for a period that is less than two-thirds of the duration of the program; and

(2) The length of time in which the residents are being training at the rural nonhospital site(s) only.

(v) All urban hospitals that wish to count FTE residents in rural tracks, not to exceed their respective rural track FTE limitation, must also comply with all of the following conditions:

(A) An urban hospital may not include in its rural track FTE limitation or (assuming the urban hospital’s FTE count exceeds its FTE cap) FTE count residents who are training in a rural track residency program that were already included as part of the hospital’s FTE cap.

(B) The hospital must base its count of residents in a rural track on written contemporaneous documentation that each resident enrolled in a rural track program at the hospital intends to rotate for a portion of the residency program to a rural area.

(C) All residents that are included by the hospital as part of its FTE count (not to exceed its rural track FTE limitation) must ultimately train in the rural area.

(vi) If HCFA finds that residents who are included by the urban hospital as part of its FTE count did not actually complete the training in the rural area, HCFA will reopen the urban hospital’s cost report within the 3-year reopening period as specified in §405.1885 of this chapter and adjust the hospital’s Medicare GME payments (and, where applicable, the hospital’s rural track FTE limitation).

(12) For purposes of paragraph (g) of this section, a new medical residency training program means a medical residency training program that receives initial accreditation by the appropriate accrediting body or begins training residents on or after January 1, 1995.

(h) Determination of weighting factors for foreign medical graduates. (1) The weighting factor for a foreign medical graduate is determined under the provisions of paragraph (g) of this section if the foreign medical graduate—

(i) Has passed FMGEMS; or

(ii) Before July 1, 1986, received certification from, or passed an examination of, the Educational Committee for Foreign Medical Graduates.

(2) Before July 1, 1986, the weighting factor for a foreign medical graduate is 1.0 times the weight determined under the provisions of paragraph (g) of this section. On or after July 1, 1986, and before July 1, 1987, the weighting factor for a graduate of a foreign medical school who was in a residency program both before and after July 1, 1986 but who does not meet the requirements set forth in paragraph (h)(1) of this section is .50 times the weight determined under the provisions of paragraph (g) of this section.

(3) On or after July 1, 1987, these foreign medical graduates are not counted in determining the number of FTE residents.

(4) During the cost reporting period in which a foreign medical graduate passes FMGEMS, the weighting factor for that resident is determined under the provisions of paragraph (g) of this section for the part of the cost reporting period beginning with the month the resident passes the test.

(5) On or after September 1, 1989, the National Board of Medical Examiners Examination, Parts I and II, may be substituted for FMGEMS for purposes of the determination made under paragraphs (h)(1) and (h)(4) of this section.

(6) On or after June 1, 1992, the United States Medical Licensing Examination may be substituted for the FMGEMS for purposes of the determination made under paragraphs (h)(1) and (h)(4) of this section. On or after July 1, 1993 only the results of steps I and II of the United States Medical Licensing Examination shall be accepted for purposes of making this determination.

(i) To include a resident in the FTE count for a particular cost reporting
period, the hospital must furnish the following information. The information must be certified by an official of the hospital and, if different, an official responsible for administering the residency program.

(1) The name and social security number of the resident.

(2) The type of residency program in which the individual participates and the number of years the resident has completed in all types of residency programs.

(3) The dates the resident is assigned to the hospital and any hospital-based providers.

(4) The dates the resident is assigned to other hospitals, or other free-standing providers, and any nonprovider setting during the cost reporting period, if any.

(5) The name of the medical, osteopathic, dental, or podiatric school from which the resident graduated and the date of graduation.

(6) If the resident is an FMG, documentation concerning whether the resident has satisfied the requirements of paragraph (h) of this section.

(7) The name of the employer paying the resident’s salary.

(1) Special rules for States that formerly had a waiver from Medicare reimbursement principles. (1) Effective for cost reporting periods beginning on or after January 1, 1986, hospitals in States that, prior to becoming subject to the prospective payment system, had a waiver for the operation of a State reimbursement control system under section 1836(c) of the Act, section 402 of the Social Security Amendments of 1967 (42 U.S.C. 1395b-1 or section 222(a) of the Social Security Amendment of 1972 (42 U.S.C. 1395b-1 (note)) are permitted to change the order in which they allocate administrative and general costs to the order specified in the instructions for the Medicare cost report.

(2) For hospitals making this election, the base-period costs for the purpose of determining the per resident amount are adjusted to take into account the change in the order by which they allocate administrative and general costs to interns and residents in approved program cost centers.

(3) Per resident amounts are determined for the base period and updated as described in paragraph (e) of this section. For cost reporting periods beginning on or after January 1, 1986, payment is made based on the methodology described in paragraph (d) of this section.

(k) Adjustment of a hospital’s target amount or prospective payment hospital-specific rate—(1) Misclassified operating costs—(i) General rule. If a hospital has its base-period graduate medical education costs reduced under paragraph (e)(1) of this section because those costs included misclassified operating costs, the hospital may request that the intermediary review the classification of the affected costs in its rate-of-increase ceiling or prospective payment base year for purposes of adjusting the hospital’s target amount or hospital-specific rate. For those cost reports that are not subject to reopening under §405.1885 of this chapter, the hospital’s reopening request must explicitly state that the review is limited to this one issue.

(ii) Request for review. The hospital must request review of the classification of its rate of increase ceiling or prospective payment base year costs no later than 180 days after the date of the notice by the intermediary of the hospital’s base-period average per resident amount. A hospital’s request for review must include sufficient documentation to demonstrate to the intermediary that adjustment of the hospital’s hospital-specific rate or target amount is warranted.

(iii) Effect of intermediary’s review. If the intermediary, upon review of the hospital’s costs, determines that the hospital’s hospital-specific rate or target amount should be adjusted, the adjustment of the hospital-specific rate or the target amount is effective for the hospital’s cost reporting periods subject to the prospective payment system or the rate-of-increase ceiling that are still subject to reopening under §405.1885 of this chapter.

(2) Misclassification of graduate medical education costs—(i) General rule. If costs that should have been classified as graduate medical education costs were treated as operating costs during both the graduate medical education base
period and the rate-of-increase ceiling base year or prospective payment base year and the hospital wishes to receive benefit for the appropriate classification of these costs as graduate medical education costs in the graduate medical education base period, the hospital must request that the intermediary review the classification of the affected costs in the rate-of-increase ceiling or prospective payment base year for purposes of adjusting the hospital’s target amount or hospital-specific rate. For those cost reports that are not subject to reopening under §405.1885 of this chapter, the hospital’s reopening request must explicitly state that the review is limited to this one issue.

(ii) Request for review. The hospital must request review of the classification of its costs no later than 180 days after the date of the intermediary’s notice of the hospital’s base-period average per resident amount. A hospital’s request for review must include sufficient documentation to demonstrate to the intermediary that modification of the adjustment of the hospital’s hospital-specific rate or target amount is warranted.

(iii) Effect of intermediary’s review. If the intermediary, upon review of the hospital’s costs, determines that the hospital’s hospital-specific rate or target amount should be adjusted, the adjustment of the hospital-specific rate and the adjustment of the target amount is effective for the hospital’s cost reporting periods subject to the prospective payment system or the rate-of-increase ceiling that are still subject to reopening under §405.1885 of this chapter.

§ 413.87 Payments for Medicare+Choice nursing and allied health education programs.

(a) Statutory basis. This section implements section 1886(l) of the Act, which provides for additional payments to hospitals that operate and receive Medicare reasonable cost reimburse-

(b) Scope. This section sets forth the rules for determining an additional payment amount to hospitals that receive payments for the costs of operating approved nursing or allied health education programs under §413.85.

(c) Qualifying conditions for payment. For portions of cost reporting periods occurring on or after January 1, 2000, a hospital that operates and receives payment for a nursing or allied health education program under §413.85 may receive an additional payment amount. The hospital may receive the additional payment amount, which is calculated in accordance with the provisions of paragraph (d) of this section, if both of the conditions specified in paragraph (c)(1) and (c)(2) of this section are met.

(1) The hospital must have received Medicare reasonable cost payment for an approved nursing or allied health education program under §413.85 in its cost reporting period(s) ending in the fiscal year that is 2 years prior to the current calendar year. (For example, if the current year is calendar year 2000, the fiscal year that is 2 years prior to calendar year 2000 is FY 1998.) For a hospital that first establishes a nursing or allied health education program and receives reasonable cost payment for the program as specified under §413.85 after FY 1998, the hospital is eligible to receive an additional payment amount in a calendar year that is 2 years after the respective fiscal year so long as the hospital also meets the condition under paragraph (c)(2) of this section.

(2) The hospital must be receiving reasonable cost payment for an approved nursing or allied health education program under §413.85 in the current calendar year.

(d) Calculating the additional payment amount. Subject to the provisions of paragraph (f) of this section relating to calculating a proportional reduction in Medicare+Choice direct GME payments, the additional payment amount specified in paragraph (c) of this section is calculated according to the following steps:
§ 413.88 Incentive payments under plans for voluntary reduction in number of medical residents.

(a) Statutory basis. This section implements section 1886(h)(6) of the Act, which establishes a program under which incentive payments may be made to qualifying entities that develop and implement approved plans to voluntarily reduce the number of residents in medical residency training.

(b) Qualifying entity defined. “Qualifying entity” means:

(1) An individual hospital that is operating one or more approved medical residency training programs as defined in §413.86(b) of this chapter; or

(2) Two or more hospitals that are operating approved medical residency training programs as defined in §413.86(b) of this chapter and that submit a residency reduction application as a single entity.

(c) Conditions for payments. (1) A qualifying entity must submit an application for a voluntary residency reduction plan that meets the requirements and conditions of this section in order to receive incentive payments for reducing the number of residents in its medical residency training programs.

(2) The incentive payments will be determined as specified under paragraph (g) of this section.

(d) Requirements for voluntary plans. In order for a qualifying entity to receive incentive payments under a voluntary residency reduction plan, the qualifying entity must submit an application that contains the following information, documents, and agreements—

(1) A description of the operation of a plan for reducing the full-time equivalent (FTE) residents in its approved medical residency training programs, consistent with the percentage reduction requirements specified in paragraphs (g)(2) and (g)(3) of this section;

(2) An election of the period of residency training years during which the reductions will occur. The reductions must be fully implemented by not later than the fifth residency training year in which the plan is effective;

(3) FTE counts for the base number of residents, as defined in paragraph (g)(1) of this section, with a breakdown of the number of primary care residents compared to the total number of residents; and the direct and indirect FTE counts of the entity on June 30, 1997. For joint applicants, these counts must be provided individually and collectively;

(4) Data on the annual and cumulative targets for reducing the number...
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of FTE residents and the ratios of the number of primary care residents to the total number of residents for the base year and for each year in the 5-year reduction period. For joint applicants, these data must be provided individually and collectively;

(5) An agreement to not reduce the proportion of its primary care residents to its total number of residents below the proportion that exists in the base year, as specified in paragraph (g)(1) of this section;

(6) An agreement to comply with data submission requirements deemed necessary by HCFA to make annual incentive payments during the 5-year residency reduction plan, and to fully cooperate with additional audit and monitoring activities deemed necessary by HCFA;

(7) For a qualifying entity that is a member of an affiliated group as defined in §413.86(b), a statement that all members of the group agree to an aggregate FTE cap that reflects—

(i) The reduction in the qualifying entity’s FTE count as specified in the plan during each year of the plan; and

(ii) The 1996 FTE count of the other hospital(s) in the affiliated group;

(8) A statement indicating voluntary participation in the plan under the terms of this section, signed by each hospital that is part of the applying entity.

(e) Deadline for applications. A qualifying entity must submit an application that meets the requirements of paragraph (d) of this section at least one day prior to the first day of the period to which the plan would be effective but no later than November 1, 1999. The application must be submitted to the fiscal intermediary, with a copy to HCFA.

(f) Effective dates of plans. Residency reduction plans that are submitted to the fiscal intermediary on or after September 17, 1999 but on or before November 1, 1999, may be effective for portions of cost reporting periods beginning no earlier than the day after the date of the application.

(g) Residency reduction requirements—

(1) Base number of residents defined. (i) "Base number of residents" means the lesser of—

(A) The number of FTE residents in all approved medical residency training programs of the qualifying entity (before application of weighting factors under §413.86(g)) for the most recent residency training year ending June 30, 1996; or

(B) The number of FTE residents in all approved medical residency training programs of the qualifying entity (before application of weighting factors under §413.86(g)) for any subsequent residency training year that ends before the date the entity submits its plan to the fiscal intermediary and HCFA.

(ii) The residency training year used to determine the base number of residents is the "base year" for determining reduction requirements.

(iii) The qualifying entity’s base number of residents may not be adjusted to reflect adjustments that may otherwise be made to the entity’s FTE caps for new medical residency training programs.

(2) Qualifying entity consisting of individual hospital. The base number of FTE residents in all the approved medical residency training programs operated by or through a qualifying entity consisting of a single hospital must be reduced as follows:

(i) If the base number of residents exceeds 750, residents, by at least 20 percent of the base number.

(ii) If the base number of residents exceeds 600 but is less than or equal to 750 residents—

(A) By 150 residents; or

(B) By 20 percent, if the qualifying entity increases the number of primary care residents included in the base number by at least 20 percent.

(iii) If the base number of residents is 600 or less residents—

(A) By 25 percent; or

(B) By 20 percent, if the qualifying entity increases the number of primary care residents included in the base number by at least 20 percent.

(3) Qualifying entity consisting of two or more hospitals. The base number of FTE residents in the aggregate for all the approved medical residency training programs operated by or through a qualifying entity consisting of two or more hospitals must be reduced—

(A) By 150 residents; or

(B) By 20 percent, if the qualifying entity increases the number of primary care residents included in the base number by at least 20 percent.
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(i) By 25 percent; or
(ii) By 20 percent, if the qualifying entity increases the number of primary care residents included in the base number of residents by at least 20 percent.

(4) Treatment of rotating residents. A qualifying entity will not be eligible for incentive payments for a reduction in the base number of residents if the reduction is a result of the entity rotating residents to another hospital that is not a part of its voluntary residency reduction plan.

(5) Updates to annual and cumulative targets

(i) Except as provided in paragraph (g)(5)(ii) of this section an entity with an approved voluntary residency reduction plan may not change the annual and cumulative reduction targets that are specified in its plan in accordance with paragraphs (g)(2) and (g)(3) of this section.

(ii) An entity may update annual reduction targets specified in its plan only if—

(A) It has failed to meet a specified annual target for a plan year in the 5-year period; and

(B) It wishes to adjust future annual targets for the remaining years of the plan in order to comply with its cumulative target.

(iii) An updated plan allowed under paragraph (g)(5)(ii) of this section must be submitted prior to the beginning of each July 1 medical residency training year during the plan years.

(h) Computation of incentive payment amount. (1) Incentive payments to qualifying entities that meets the requirements and conditions of paragraphs (d) and (g) of this section will be computed as follows:

(i) Step 1. Determine the amount (if any) by which the payment amount that would have been made under §413.86(d) if there had been a 5-percent reduction in the number of FTE residents in the approved medical education training programs of the hospital as of June 30, 1997, exceeds the payment amount made under §412.105 of this chapter in each year under the voluntary residency reduction plan, taking into account the actual reduction in the number of FTE residents.

(ii) Step 2. Determine the amount (if any) by which the payment amount that would have been made under §412.322 of this chapter if there had been a 5-percent reduction in the number of FTE residents in the approved medical education training programs of the hospital as of June 30, 1997, exceeds the payment amount made under §412.322 of this chapter in each year under the voluntary residency reduction plan, taking into account the actual reduction in the number of FTE residents.

(iii) Step 3. Determine the amount (if any) by which the payment amount that would have been made under §412.105 of this chapter if there had been a 5-percent reduction in the number of FTE residents in the approved medical education training programs of the hospital as of June 30, 1997, exceeds the payment amount made under §412.322 of this chapter in each year under the voluntary residency reduction plan, taking into account the actual reduction in the number of FTE residents.

(iv) Step 4. Multiply the sum of the amounts determined under paragraph (h)(1), (ii), and (iii) of this section by the applicable hold harmless percentages specified in paragraph (i) of this section.

(i) Applicable hold-harmless percentage. The applicable hold-harmless percentages for each year in which the residency reduction plan is in effect as follows:

(1) 100 percent for the first and second residency training years;
(2) 75 percent for the third year;
(3) 50 percent for the fourth year; and
(4) 25 percent for the fifth year.

(j) Payments to qualifying entities. Annual incentive payments through cost reports will be made to each hospital that is or is part of a qualifying entity over the 5-year reduction period if the qualifying entity meets the annual and cumulative reduction targets specified in its voluntary reduction plan.
§ 413.90 Research costs.

(a) Principle. Costs incurred for research purposes, over and above usual patient care, are not includable as allowable costs.

(b) Application. (1) There are numerous sources of financing for health-related research activities. Funds for this purpose are provided under many Federal programs and by other tax-supported agencies. Also, many foundations, voluntary health agencies, and other private organizations, as well as individuals, sponsor or contribute to the support of medical and related research. Funds available from such sources are generally ample to meet basic medical and hospital research needs. A further consideration is that quality review should be assured as a condition of governmental support for research. Provisions for such review would introduce special difficulties in the Medicare programs.

(2) If research is conducted in conjunction with, and as a part of, the care of patients, the costs of usual patient care and studies, analyses, surveys, and related activities to serve the provider's administrative and program needs are allowable costs in the determination of payment under Medicare.

§ 413.92 Costs of surety bonds.

Costs incurred by a provider to obtain a surety bond required by part 489, subpart F of this chapter are not includable as allowable costs.

§ 413.94 Value of services of nonpaid workers.

(a) Principle. The value of services in positions customarily held by full-time employees performed on a regular, scheduled basis by individuals as nonpaid members of organizations under arrangements between such organizations and a provider for the performance of such services without direct remuneration from the provider to such individuals is allowable as an operating expense for the determination of allowable cost subject to the limitation contained in paragraph (b) of this section. The amounts allowed are not to exceed those paid others for similar work. Such amounts must be identifiable in the records of the institutions as a legal obligation for operating expenses.

(b) Limitations: Services of nonpaid workers. The services must be performed on a regular, scheduled basis in positions customarily held by full-time employees and necessary to enable the provider to carry out the functions of normal patient care and operation of
the institution. The value of services of a type for which providers generally do not remunerate individuals performing such services is not allowable as a reimbursable cost under the Medicare program. For example, donated services of individuals in distributing books and magazines to patients, or in serving in a provider canteen or cafeteria or in a provider gift shop, would not be reimbursable.

(c) Application. The following illustrates how a provider would determine an amount to be allowed under this principle: The prevailing salary for a lay nurse working in Hospital A is $5,000 for the year. The lay nurse receives no maintenance or special perquisites. A sister working as a nurse engaged in the same activities in the same hospital receives maintenance and special perquisites which cost the hospital $2,000 and are included in the hospital’s allowable operating costs. The hospital would then include in its records an additional $3,000 to bring the value of the services rendered to $5,000. The amount of $3,000 would be allowable if the provider assumes obligation for the expense under a written agreement with the sisterhood or other religious order covering payment by the provider for the services.

§ 413.98 Purchase discounts and allowances, and refunds of expenses.

(a) Principle. Discounts and allowances received on purchases of goods or services are reductions of the costs to which they relate. Similarly, refunds of previous expense payments are reductions of the related expense.

(b) Definitions—(1) Discounts. Discounts, in general, are reductions granted for the settlement of debts.

(2) Allowances. Allowances are deductions granted for damage, delay, shortage, imperfection, or other causes, excluding discounts and returns.

(3) Refunds. Refunds are amounts paid back or a credit allowed on account of an overcollection.

(c) Normal accounting treatment—Reduction of costs. All discounts, allowances, and refunds of expenses are reductions in the cost of goods or services purchased and are not income. If they are received in the same accounting period in which the purchases were made or expenses were incurred, they will reduce the purchases or expenses of that period. However, if they are received in a later accounting period, they will reduce the comparable purchases or expenses in the period in which they are received.

(d) Application. (1) Purchase discounts have been classified as cash, trade, or quantity discounts. Cash discounts are reductions granted for the settlement of debts before they are due. Trade discounts are reductions from list prices granted to a class of customers before consideration of credit terms. Quantity discounts are reductions from list prices granted because of the size of individual or aggregate purchase transactions. Whatever the classification of purchase discounts, like treatment in reducing allowable costs is required. In the past, purchase discounts were considered as financial management income. However, modern accounting theory holds that income is not derived from a purchase but rather from a sale or an exchange and that purchase discounts are reductions in the cost of whatever was purchased. The true cost of the goods or services is the net amount actually paid for them. Treating purchase discounts as income would result in an overstatement of costs to the extent of the discount.

(2) As with discounts, allowances, and rebates received from purchases of goods or services, refunds of previous expense payments are clearly reductions in costs and must be reflected in the determination of allowable costs. This treatment is equitable and is in accord with that generally followed by other governmental programs and third-party payment organizations paying on the basis of cost.

§ 413.100 Special treatment of certain accrued costs.

(a) Principle. As described in §413.24(b)(2), under the accrual basis of accounting, revenue is reported in the period in which it is earned and expenses are reported in the period in which they are incurred. In the case of accrued costs described in this section, for Medicare payment purposes the costs are allowable in the year in which the costs are accrued and claimed for Medicare payment only.
under the conditions set forth in paragraph (c) of this section.

(b) Definitions—(1) All-inclusive paid days off benefit. An all-inclusive paid days off benefit replaces other vacation and sick pay plans. It is a formal plan under which, based on actual hours worked, all employees accrue vested
leave or payment in lieu of vested leave for any combination of types of leave, such as illness, medical appointments, holidays, and vacations.

(2) Self-insurance. Self-insurance is a means by which a provider independently or as part of a group undertakes the risk of protecting itself against anticipated liabilities by providing funds in an amount equal to anticipated liabilities, rather than by purchasing insurance coverage.

(c) Recognition of accrued costs—(1) General. Although Medicare recognizes, in the year of accrual, the accrual of costs for which a provider has not actually expended funds during the current cost reporting period, for purposes of payment Medicare does not recognize the accrual of costs unless the related liabilities are liquidated timely.

(2) Requirements for liquidation of liabilities. For accrued costs to be recognized for Medicare payment in the year of the accrual, the requirements set forth below must be met with respect to the liquidation of related liabilities. If liquidation does not meet these requirements, the cost is disallowed, generally within 1 year after the end of the cost reporting period in which the liability was incurred.

(i) A short-term liability. (A) Except as provided in paragraph (c)(2)(ii)(B) of this section, a short-term liability, including the current portion of a long-term liability (for example, mortgage interest payments due to be paid in the current year), must be liquidated within 1 year after the end of the cost reporting period in which the liability is incurred.

(B) If, within the 1-year time limit, the provider furnishes to the intermediary sufficient written justification (based upon documented evidence) for nonpayment of the liability, the intermediary may grant an extension for good cause. The extension may not exceed 3 years beyond the end of the cost reporting year in which the liability was incurred.

(ii) Vacation pay and all-inclusive paid days off. (A) If the provider’s vacation policy, or its policy for all-inclusive paid days off, is consistent for all employees, liquidation of the liability must be made within the period provided for by that policy.

(B) If the provider’s vacation policy, or its policy for all-inclusive paid days off, is not consistent for all employees, liquidation of the liability must be made within 2 years after the close of the cost reporting period in which the liability is accrued.

(C) If payment is not made within the required time period or if benefits are forfeited by the employee, an adjustment to disallow the accrued cost is made in the current period (that is, the latest year in which payment should have been made or the year in which the benefits are forfeited) rather than in the period in which the cost was accrued and claimed for Medicare payment. However, an intermediary may choose to require the adjustment in the period in which the cost was accrued and claimed for Medicare payment if the cost report for that period is open or can be reopened as provided in §405.1885 of this chapter, and if the intermediary believes the adjustment is more appropriate in that period.

(iii) Sick pay. (A) If sick leave is vested and funded in a deferred compensation plan, liabilities related to the contributions to the fund must be liquidated, generally within 1 year after the end of the cost reporting period in which the liability is incurred. If, within the 1-year time limit, the provider furnishes to the intermediary sufficient written justification (based upon documented evidence) for nonpayment of the liability, the intermediary may grant an extension for good cause. The extension may not exceed 3 years beyond the end of the cost reporting period in which the liability was incurred. Contributions to the deferred compensation plan must be reduced to reflect estimated forfeitures. Actual forfeitures above or below estimated forfeitures must be used to adjust annual contributions to the fund.
(B) If the sick leave plan grants employees the nonforfeitable right to demand cash payment for unused sick leave at the end of each year, sick pay is includable in allowable costs, without funding, in the cost reporting period in which it is earned.

(C) Sick pay paid on any basis other than that specified in paragraphs (c)(2)(iii) (A) or (B) of this section can be claimed for Medicare payment only on a cash basis for the year in which the benefits are paid.

(iv) Compensation of owners. Accrued liability related to compensation of owners other than sole proprietors and partners must be liquidated within 75 days after the close of the cost reporting period in which the liability occurs.

(v) Nonpaid workers. Obligations incurred under a legally-enforceable agreement to remunerate an organization of nonpaid workers must be discharged no later than the end of the provider's cost reporting period following the period in which the services were furnished.

(vi) FICA and other payroll taxes—(A) General rule. The provider's share of FICA and other payroll taxes that the provider becomes obligated to remit to governmental agencies is included in allowable costs only during the cost reporting period in which payment (upon which the payroll taxes are based) is actually made to the employee. For example, payroll taxes applicable to vacation benefits are not to be accrued in the period in which the vacation benefits themselves are accrued but rather are allowable only in the period in which the employee takes the vacation.

(B) Exception. If payment would be made to an employee during a cost reporting period but for the fact the regularly scheduled payment date is after the end of the period, costs of accrued payroll taxes related to the portion of payroll accrued through the end of the period, but paid to the employee after the beginning of the new period, are allowable costs in the year of accrual, subject to the liquidation requirements specified in paragraph (c)(2)(i) of this section.

(vii) Deferred compensation. (A) Reasonable provider payments made under unfunded deferred compensation plans are included in allowable costs only during the cost reporting period in which actual payment is made to the participating employee.

(B) Accrued liability related to contributions to a funded deferred compensation plan must be liquidated within 1 year after the end of the cost reporting period in which the liability is incurred. An extension, not to exceed 3 years beyond the end of the cost reporting year in which the liability was incurred, may be granted by the intermediary for good cause if the provider, within the 1-year time limit, furnishes to the intermediary sufficient written justification for non-payment of the liability.

(C) Postretirement benefit plans (including those addressed in Statement of Financial Accounting Standards No. 106 (December 1990)) are deferred compensation arrangements and thus are subject to the provisions of this section regarding deferred compensation and to applicable program instructions for determining Medicare payment for deferred compensation.

(viii) Self-insurance. Accrued liability related to contributions to a self-insurance program that are systematically made to a funding agency and that cover malpractice and comprehensive general liability, unemployment compensation, workers' compensation insurance losses, or employee health benefits, must be liquidated within 75 days after the close of the cost reporting period.

§ 413.102 Compensation of owners.

(a) Principle. A reasonable allowance of compensation for services of owners is an allowable cost provided that the services are actually performed in a necessary function.

(b) Definitions. (1) Compensation. Compensation means the total benefit received by the owner for the services he furnishes to the institution. It includes the following items:

(i) Salary amounts paid for managerial, administrative, professional, and other services.

(ii) Amounts paid by the institution for the personal benefit of the proprietor.
§ 413.106 Reasonable cost of physical and other therapy services furnished under arrangements.

(a) Principle. The reasonable cost of the services of physical, occupational, speech, and other therapists, and services of other health specialists (other than physicians), furnished under arrangements (as defined in section 1861(w) of the Act) with a provider of services, a clinic, a rehabilitation agency or a public health agency, may not exceed an amount equivalent to the prevailing salary and additional costs that would reasonably have been incurred by the provider or other organization had such services been performed by such person in an employment relationship, plus the cost of other reasonable expenses incurred by such person in furnishing services under such an arrangement. However, if the services of a therapist are required on a limited part-time basis, or to perform intermittent services, payment may be made on the basis of a reasonable rate per unit of service, even though this rate may be greater per unit of time than salary-related amounts, if the greater payment is, in the aggregate, less than the amount that would have been paid had a therapist been employed on a full-time or regular part-time salaried basis. Pursuant to section 17(a) of Public Law 93-233 (87 Stat. 967), the provisions of this section are effective for cost reporting periods beginning after March, 1975.

(b) Definitions—(1) Prevailing salary. The prevailing salary is the hourly salary rate based on the 75th percentile of salary ranges paid by providers in the geographical area, by type of therapy, to therapists working full time in an employment relationship.

(2) Fringe benefit and expense factor. The standard fringe benefit and expense factor is an amount that takes account of fringe benefits, such as vacation pay, insurance premiums, pension payments, allowances for job-related training, meals, etc., generally received by an employee therapist, as well as expenses, such as maintaining an office, appropriate insurance, etc., an individual not working as an employee might incur in furnishing services under arrangements.

(3) Adjusted hourly salary equivalency amount. The adjusted hourly salary equivalency amount is the prevailing hourly salary rate plus the standard fringe benefit and expense factor. This amount is determined on a periodic...
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(4) Travel allowance. A standard travel allowance is an amount that is recognized, in addition to the adjusted hourly salary equivalency amount.

(5) Limited part-time or intermittent services. Therapy services are considered to be on a limited part-time or intermittent basis if the provider or other organization furnishing the services under arrangements requires the services of a therapist or therapists on an average of less than 15 hours per week. This determination is made by dividing the total hours of services furnished during the cost reporting period by the number of weeks in which the services were furnished in the cost reporting period regardless of the number of days in each week in which services were performed.

(6) Guidelines. Guidelines are the amounts published by HCFA reflecting the application of paragraphs (b) (1) through (4) of this section to an individual therapy service and a geographical area. Other statistically valid data may be used to establish guidelines for a geographical area, provided that the study designs, questionnaires and instructions, as well as the resultant survey data for determining the guidelines are submitted to and approved in advance by HCFA. Such data must be arrayed so as to permit the determination of the 75th percentile of the range of salaries paid to full-time employee therapists.

(7) Administrative responsibility. Administrative responsibility is the performance of those duties that normally fall within the purview of a department head or other supervisor. This term does not apply to directing aides or other assistants in furnishing direct patient care.

(c) Application. (1) Under this provision, HCFA will establish criteria for use in determining the reasonable cost of physical, occupational, speech, and other therapy services and the services of other health specialists (other than physicians) furnished by individuals under arrangements with a provider of services, a clinic, a rehabilitation agency, or public health agency. It is recognized that providers have a wide variety of arrangements with such individuals. These individuals may be independent practitioners or employees of organizations furnishing various health care specialists. This provision does not require change in the substance of these arrangements.

(2) If therapy services are performed under arrangements at a provider site on a full-time or regular part-time basis, the reasonable cost of such services may not exceed the amount determined by taking into account the total number of hours of services furnished by the therapist, the adjusted hourly salary equivalency amount appropriate for the particular therapy in the geographical area in which the services are furnished and a standard travel allowance.

(3) If therapy services are performed under arrangements on a limited part-time or intermittent basis at the provider site, the reasonable cost of such services is evaluated on a reasonable rate per unit of service basis, except that payment for these services, in the aggregate, during the cost reporting period, may not exceed the amount that would be determined to be reasonable under paragraph (c)(2) of this section, had a therapist furnished the provider or other organization furnishing the services under arrangements 15 hours of service per week on a regular part-time basis for the weeks in which services were furnished by the non-employee therapist.

(4) If an HHA furnishes services under arrangements at the patient's residence or in other situations in which therapy services are not performed at the provider's site, the reasonable cost of such services is evaluated as follows:

(i) Time records available. If time records of HHA visits are maintained by the provider, the reasonable cost of such services is evaluated on a unit-of-time basis, by taking into account the total number of hours of service furnished by the therapist, the adjusted hourly salary equivalency amount appropriate for the particular therapy in the geographical area in which the services are furnished, and a standard travel allowance for each visit. However, if the travel time of the therapist is accurately recorded by the therapist, and approved and maintained by the provider, the reasonable cost of such
services may be evaluated, at the option of the provider, by taking into account the total number of hours of service furnished by the therapist, including travel time, and the adjusted hourly salary equivalency amount appropriate for the particular therapy in the geographical area in which the services are furnished. This option does not apply to services furnished by HHAs under arrangements with providers other than HHAs.

(ii) No time records available. If time records are unavailable or found to be inaccurate, each HHA visit is considered the equivalent of one hour of service. In such cases, the reasonable cost of such services is determined by taking into account the number of visits made by the therapist under arrangements with such agency, the adjusted hourly salary equivalency amount appropriate for the particular therapy in the geographical area in which the services are furnished, and a standard travel allowance.

(iii) Limited part-time or intermittent services. If under paragraph (c)(4)(i) or (ii) of this section, the provider required therapy services on an average of less than 15 hours per week, the services are considered limited part-time or intermittent services, and the reasonable cost of such services is evaluated on a reasonable rate per unit of service basis as described in paragraph (c)(3) of this section.

(5) If therapy services are performed in situations where compensation to a therapist employed by the provider is based, at least in part, on a fee-for-service or on a percentage of income (commission), the guidelines will apply. The entire compensation will be subject to the guidelines in cases where the nature of the arrangements is most like an "arrangement" situation, although technically the provider may treat the therapists as employees. The intent of this section is to prevent an employment relationship from being used to circumvent the guidelines.

(6) These provisions are applicable to individual therapy services or disciplines by means of separate guidelines by geographical area and apply to costs incurred after issuance of the guidelines but no earlier than the beginning of the provider's cost reporting period described in paragraph (a) of this section. Until a guideline is issued for a specific therapy or discipline, costs are evaluated so that such costs do not exceed what a prudent and cost-conscious buyer would pay for the given service.

(d) Notice of guidelines to be imposed. Prior to the beginning of a period to which a guideline will be applied, a notice will be published in the FEDERAL REGISTER establishing the guideline amounts to be applied to each geographical area by type of therapy.

(e) Additional allowances. (1) If a therapist supervises other therapists or has administrative responsibility for operating a provider's therapy department, a reasonable allowance may be added to the adjusted hourly salary equivalency amount by the intermediary based on its knowledge of the differential between therapy supervisors' and therapists' salaries in similar provider settings in the area.

(2) If a therapist performing services under arrangements furnishes equipment and supplies used in furnishing therapy services, the guideline amount may be supplemented by the cost of the equipment and supplies, provided the cost does not exceed the amount the provider, as a prudent and cost-conscious buyer, would have been able to include as allowable costs.

(f) Exceptions. The following exceptions may be granted but only upon the provider's demonstration that the conditions indicated are present:

(1) Exception because of unique circumstances or special labor market conditions. An exception may be granted under this section by the intermediary if a provider demonstrates that the costs for therapy services established by the guideline amounts are inappropriate to a particular provider because of some unique circumstances or special labor market conditions in the area.

(2) Exception for services furnished by risk-basis HMO providers. For special rules concerning services furnished to an HMO's enrollees who are Medicare beneficiaries by a provider owned or operated by a risk-basis HMO (see §417.201(b) of this chapter) or related to
a risk-basis HMO by common ownership or control (see §417.250(c) of this chapter).

(3) Exception for inpatient hospital services. Effective with cost reporting periods beginning on or after October 1, 1983, the costs of therapy services furnished under arrangements to a hospital inpatient are excepted from the guidelines issued under this section if such costs are subject to the provisions of §413.40 or part 412 of this chapter. The intermediary will grant the exception without request from the provider.

(g) Appeals. A request by a provider for a hearing on the determination of an intermediary concerning the therapy costs determined to be allowable based on the provisions of this section, including a determination with respect to an exception under paragraph (f) of this section, is made to the intermediary only after submission of its cost report and receipt of the notice of amount of program reimbursement reflecting such determination, in accordance with the provisions of subpart R of part 405 of this chapter.


§ 413.114 Payment for posthospital SNF care furnished by a swing-bed hospital.

(a) Purpose and basis. This section implements section 1883 of the Act, which provides for payment for posthospital SNF care furnished by rural hospitals having a swing-bed approval. Payments to these hospitals for posthospital SNF care furnished in general routine inpatient beds are based on the reasonable cost of posthospital SNF care in accordance with paragraph (c) of this section. Swing-bed hospitals approved after March 31, 1988 with more than 49 beds must meet additional payment requirements as set forth in paragraph (d) of this section.

(b) Definitions. For purposes of this section—

Availability date means with respect to a posthospital SNF care patient in a swing-bed hospital, the later of—

(i) Any date on which a bed is available for the patient in a Medicare-participating SNF located within the hospital’s geographic region;

(ii) The date that a hospital learns that a bed is available in a Medicare-participating SNF; or

(iii) If the notice is prospective, the date that a bed will become available in a Medicare-participating SNF.

Geographic region means an area that includes the SNFs with which a hospital has traditionally arranged transfers and all other SNFs within the same proximity to the hospital. In the case of a hospital without existing transfer practices upon which to base a determination, the geographic region is an area that includes all the SNFs within 50 miles (as defined in §412.92(c)(1) of this chapter) of the hospital unless the hospital can demonstrate that the SNFs are inaccessible to its patients. In the event of a dispute as to whether an SNF is within a hospital’s geographic region or the SNF is inaccessible to hospital patients, the HCFA Regional Office makes a determination.

Swing-bed hospital means a hospital or CAH participating in Medicare that has an approval from HCFA to provide posthospital SNF care as defined in §409.20 of this chapter, and meets the requirements specified in §482.66 or §485.645 of this chapter, respectively.

(c) Principle. The reasonable cost of posthospital SNF care furnished by a swing-bed hospital is determined as follows:

(1) The reasonable cost of routine SNF services is based on the average Medicare rate per patient day for routine services provided in freestanding SNFs in the region where the swing-bed hospital is located. The rates are calculated using the regions as defined in section 1886(d)(2)(D) of the Social Security Act. The rates are based on the most recent year for which settled cost reporting period data are available, increased in a compounded manner, using the increase applicable to the SNF routine cost limits, up to and including the calendar year for which the rates are in effect. If the current Medicare swing-bed rate for routine extended care services furnished by a swing-bed hospital during a calendar year is less than the rate for the prior calendar year, payment is made based on the prior calendar year’s rate.
§413.118 Payment for facility services related to covered ASC surgical procedures performed in hospitals on an outpatient basis.

(a) Basis and scope. This section implements section 1833(a)(4) and (i)(3) of the Act and establishes the method for determining Medicare payments for services related to covered ambulatory surgical center (ASC) procedures performed in a hospital on an outpatient basis. It does not apply to services furnished by an ASC operated by a hospital that has an agreement with HCFA to be paid in accordance with §416.30 of this chapter. (For regulations governing ASCs see part 416 of this chapter.)

(b) Definitions. For purposes of this section—

Facility services are those items and services, as specified in §416.61 of this chapter, that are furnished by a hospital on an outpatient basis in connection with covered ASC surgical procedures, as described in §416.65 of this chapter.

Standard overhead amount means an amount equal to the prospectively determined payment rate that would be paid for the procedure if it had been furnished by an ASC in the same geographic area.

(c) Payment principle. The aggregate amount of payments for facility services, furnished in a hospital on an outpatient basis, that are related to covered ASC surgical procedures (covered under §416.65 of this chapter) is equal to the lesser of—

(1) The hospital's reasonable cost or customary charges, as determined in
accordance with §413.13, reduced by deductibles and coinsurance; or

(2) The blended payment amount as described in paragraph (d) of this section, which is based on hospital-specific cost and charge data and rates paid to free-standing ASCs.

(d) Blended payment amount. (1) For cost reporting periods beginning on or after October 1, 1987 but before October 1, 1988, the blended payment amount is equal to the sum of—

(i) 75 percent of the hospital-specific amount (the lesser of the hospital's reasonable cost or customary charges, reduced by deductibles and coinsurance); and

(ii) 25 percent of the ASC payment amount (that is, 80 percent of the result obtained by subtracting the deductibles from the sum of the standard overhead amounts.)

(2) For the period of time beginning with the first day of a hospital's cost reporting period that begins on or after October 1, 1988 and ends on December 31, 1990, the blended payment amount is equal to 50 percent of the hospital-specific amount and 50 percent of the ASC payment amount.

(3) For portions of cost reporting periods beginning on or after January 1, 1991, the blended payment amount is equal to 42 percent of the hospital-specific amount and 58 percent of the ASC payment amount.

(4) For cost reporting periods beginning on or after October 1, 1988 and before January 1, 1995, the blended payment amount is equal to the sum of 75 percent of the hospital-specific amount and 25 percent of the ASC payment amount for a hospital that makes an application to its fiscal intermediary and meets the following requirements.

(i) More than 60 percent of the hospital's inpatient hospital discharges, as described in §412.60 of this chapter, occurring during its cost reporting period beginning on or after October 1, 1986 and before October 1, 1987, are classified in diagnosis related groups 36 through 74.

(ii) During its cost reporting period beginning on or after October 1, 1986 and before October 1, 1987, more than 30 percent of the hospital's total revenues is derived from outpatient services.

(5) For portions of cost reporting periods beginning on or after October 1, 1997, for purposes of calculating the blended payment amount under paragraph (d)(4) of this section, the ASC payment amount is the sum of the standard overhead amounts reduced by deductibles and coinsurance as defined in section 1866(a)(2)(ii) of the Act.

(e) Aggregation of cost, charges, and the blended amount. For purposes of determining the correct payment amount under paragraphs (c) and (d) of this section, all reasonable costs and customary charges attributable to facility services furnished during a cost reporting period are aggregated and treated separately from the reasonable costs and customary charges attributable to all other services furnished in the hospital.

§413.122 Payment for hospital outpatient radiology services and other diagnostic procedures.

(a) Basis and purpose. (1) This section implements section 1833(n) of the Act and establishes the method for determining Medicare payments for radiology services and other diagnostic procedures performed by a hospital on an outpatient basis.

(2) For purposes of this section—

(i) Radiology services include diagnostic and therapeutic radiology, nuclear medicine, CAT scan procedures, magnetic resonance imaging, ultrasound and other imaging services; and

(ii) Other diagnostic procedures are those identified by HCFA, and do not include diagnostic radiology procedures or diagnostic laboratory tests.

(b) Payment for hospital outpatient radiology services. (1) The aggregate payment for hospital outpatient radiology services furnished on or after October 1, 1988 is equal to the lesser of the following:

(i) The hospital's reasonable cost or customary charges, as determined in accordance with §413.13, reduced by the applicable Part B annual deductible and coinsurance amounts.
(ii) The blended payment amount described in paragraph (b)(2) of this section.

(2) The blended payment amount for hospital outpatient radiology services furnished on or after October 1, 1988, but before October 1, 1989, is equal to the sum of—

(i) 65 percent of the hospital-specific amount (the hospital’s reasonable cost or customary charges, whichever is less, reduced by the applicable Part B annual deductible and coinsurance amounts); and

(ii) 35 percent of a prevailing charge or fee schedule amount that is calculated as 80 percent of the amount determined by subtracting the applicable Part B annual deductible from 62 percent of the prevailing charges (or for services furnished on or after January 1, 1989, the fee schedule amount established for the same services when furnished by participating physicians in their offices in the same locality).

(3) For hospital outpatient radiology services furnished on or after October 1, 1989, the blended payment amount is equal to the sum of 50 percent of the hospital-specific amount and 50 percent of the fee schedule amount.

(4) For hospital outpatient radiology services furnished on or after January 1, 1991, the blended payment amount is equal to the sum of 42 percent of the hospital-specific amount and 58 percent of the fee schedule amount.

(5) For hospital outpatient radiology services furnished on or after October 1, 1997, the blended payment amount is equal to the sum of—

(i) 42 percent of the hospital-specific amount; and

(ii) 58 percent of the fee schedule amount calculated as 62 percent of the sum of the fee schedule amounts payable for the same services when furnished by participating physicians in their offices in the same locality, less deductible and coinsurance as defined in section 1866(a)(2)(A)(ii) of the Act.

(c) Payment for other diagnostic procedures. (1) The aggregate payment for other diagnostic procedures performed by a hospital on an outpatient basis on or after October 1, 1989, is equal to the sum of—

(i) 65 percent of the hospital-specific amount (the hospital’s reasonable cost or customary charges, whichever is less, reduced by the applicable Part B annual deductible and coinsurance amounts); and

(ii) 35 percent of a prevailing charge or fee schedule amount that is calculated as 80 percent of the amount determined by subtracting the applicable Part B annual deductible from 42 percent of the prevailing charges for the same services furnished by participating physicians in their offices in the same locality.

(3) For other diagnostic procedures performed by a hospital on or after October 1, 1990, the blended payment is equal to 50 percent of the hospital-specific amount and 50 percent of the prevailing charge amount.

(4) For other diagnostic services furnished on or after October 1, 1997, the blended payment amount is equal to the sum of—

(i) 50 percent of the hospital-specific amount; and

(ii) 50 percent of the fee schedule amount calculated as 42 percent of the sum of the fee schedule amounts payable for the same services when furnished by participating physicians in their offices in the same locality, less deductible and coinsurance as defined in section 1866(a)(2)(A)(ii) of the Act.

§413.123 Payment for screening mammography performed by hospitals on an outpatient basis.

(a) Basis and scope. This section implements section 1834(c)(1)(C) of the Act and establishes the method for determining Medicare payment for screening mammographies performed by hospitals.

(b) Payment to hospitals for outpatient services. Payment to hospitals for
screening mammography services performed on an outpatient basis is determined in accordance with the technical component billing requirements in § 405.534(d) of this chapter.


§ 413.124 Reduction to hospital outpatient operating costs.

(a) Except for sole community hospitals, as defined in §412.92 of this chapter, and critical access hospitals, the reasonable costs of outpatient hospital services (other than capital-related costs of these services) are reduced by 5.8 percent for services furnished during portions of cost reporting periods occurring on or after October 1, 1990 and until the first date that the prospective payment system under part 419 of this chapter is implemented.

(b) For purposes of determining the blended payment amounts of ambulatory surgical center approved surgical procedures performed in the hospital outpatient setting under §413.118 and hospital outpatient radiology services and other diagnostic procedures under §413.122, the reduction is applicable only to the hospital-specific portion of the blended payment amounts.


§ 413.125 Payment for home health agency services.

(a) For additional rules on the allowability of certain costs incurred by home health agencies, see §§409.46 and 409.49(b) of this chapter.

(b) The reasonable cost of outpatient rehabilitation services furnished by a home health agency to homebound patients who are not entitled to home health benefits may not exceed the amounts payable under the physician fee schedule for comparable services effective January 1, 1999.

and rentals, including licenses and royalty fees, are includable in capital-related costs if they relate to the use of assets that would be depreciable if the provider owned them outright or they relate to land, which is neither depreciable nor amortizable if owned outright. The terms "leases" and "rentals of assets" signify that a provider has possession, use, and enjoyment of the assets.

(2) For sale and leaseback agreements for hospitals and SNFs entered into before October 23, 1992 and for sale and leaseback agreements for other providers entered into at any time, a provider may include incurred rental charges in its capital-related costs, as specified in a sale and leaseback agreement with a nonrelated purchaser (including shared service organizations not related within the meaning of §413.17) involving plant facilities or equipment only if the following conditions are met:

(i) The rental charges are reasonable based on the following—

(A) Consideration of rental charges of comparable facilities and market conditions in the area;

(B) The type, expected life, condition, and value of the facilities or equipment rented; and

(C) Other provisions of the rental agreements.

(ii) Adequate alternative facilities or equipment that would serve the purpose are not or were not available at lower cost.

(iii) The leasing was based on economic and technical considerations.

(3) If the conditions of paragraph (b)(2) of this section are not met, the amount a provider may include in its capital-related costs as rental or lease expense may not exceed the amount that the provider would have included in its capital-related costs had the provider retained legal title to the facilities or equipment, such as interest expense on mortgages, taxes, depreciation, and insurance costs (the costs of ownership). This limitation applies both on an annual basis and over the useful life of the asset.

(i) If in the early years of the lease, the annual rental or lease costs are less than the annual costs of ownership, but in the later years of the lease the annual rental or lease costs are more than the annual costs of ownership, in the years that the annual rental or lease costs are more than the annual costs of ownership, the provider may include in capital-related costs annually the actual amount of rental or lease costs. The aggregate rental or lease costs included in capital-related costs over the useful life of the asset had the provider retained legal title to the assets.

(ii) If in the early years of the lease, the annual rental or lease costs exceed the annual costs of ownership, but in the later years of the lease the annual rental or lease costs are less than the annual costs of ownership, the provider may carry forward amounts of rental or lease costs that were not included in capital-related costs in the early years of the lease due to the costs of ownership limitation, and include these amounts in capital-related costs in the years of the lease when the annual rental or lease costs are less than the annual costs of ownership.

(iii) In any given year the amount of actual annual rental or lease costs plus the amount carried forward to that year may not exceed the amount of the costs of ownership for that year.

(iv) In the aggregate, the amount of rental or lease costs included in capital-related costs may not exceed the amount of the costs of ownership that the provider could have included in capital-related costs had the provider retained legal title to the asset.

(5) For sale purchase transactions entered into before October 23, 1992, a lease that meets the following conditions establishes a virtual purchase:
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(i) The rental charge exceeds rental charges of comparable facilities or equipment in the area.

(ii) The term of the lease is less than the useful life of the facilities or equipment.

(iii) The provider has the option to renew the lease at a significantly reduced rental, or the provider has the right to purchase the facilities or equipment at a price that appears to be significantly less than what the fair market value of the facilities or equipment would be at the time the acquisition by the provider is permitted.

(b)(i) If a lease is a virtual purchase under paragraph (b)(5) of this section, the rental charge is includable in capital-related costs only to the extent that it does not exceed the amount that the provider would have included in capital-related costs if it had legal title to the asset (the cost of ownership), such as straight-line depreciation, insurance, and interest. A provider may not include in its capital-related costs accelerated depreciation in this situation.

(ii) The difference between the amount of rent paid and the amount of rent allowed as capital-related costs is considered a deferred charge and is capitalized as part of the historical cost of the asset when the asset is purchased.

(iii) If an asset is returned to the owner, instead of being purchased, the deferred charge may be included in capital-related costs in the year the asset is purchased.

(iv) If the term of the lease is extended for an additional period of time at a reduced lease cost and the option to purchase no longer exists, the deferred charge may be included in capital-related costs to the extent of increasing the reduced rental to an amount not in excess of the cost of ownership.

(v) If the term of the lease is extended for an additional period of time at a reduced lease cost and the option to purchase no longer exists, the deferred charge may be included in the capital-related costs to the extent of increasing the reduced rental to a fair rental value.

(7) Amounts included in lease or rental payments for repair or maintenance agreements are excluded from capital-related costs. If no amount is identified in the lease or rental agreement for maintenance, the entire lease payment is considered a capital-related cost subject to the provisions of paragraph (b)(1) of this section.

(8) For lease purchase transactions entered into on or after October 23, 1992, a lease that meets any one of the following conditions establishes a virtual purchase:

(i) The lease transfers title of the facilities or equipment to the lessee during the lease term.

(ii) The lease contains a bargain purchase option.

(iii) The lease term is at least 75 percent of the useful life of the facilities or equipment. This provision is not applicable if the lease begins in the last 25 percent of the useful life of the facilities or equipment.

(iv) The present value of the minimum lease payments (payments to be made during the lease term including bargain purchase option, guaranteed residual value, and penalties for failure to renew) equals at least 90 percent of the fair market value of the leased property. This provision is not applicable if the lease begins in the last 25 percent of the useful life of the facilities or equipment. Present value is computed using the lessee’s incremental borrowing rate, unless the interest rate implicit in the lease is known and is less than the lessee’s incremental borrowing rate, in which case the interest rate implicit in the lease is used.

(9)(i) If a lease establishes a virtual purchase under paragraph (b)(8) of this section, the rental charge is includable in capital-related costs to the extent that it does not exceed the amount that the provider would have included in capital-related costs if it had legal title to the asset (the cost of ownership). The cost of ownership includes straight-line depreciation, insurance, and interest. For purposes of computing the limitation on allowable rental cost in this paragraph, a provider may not include accelerated depreciation.

(ii) The difference between the amount of rent paid and the amount of rent allowed as capital-related costs is considered a deferred charge and is capitalized as part of the historical cost of the asset.
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cost of the asset when the asset is purchased.

(iii) If an asset is returned to the owner instead of being purchased, the deferred charge may be included in capital-related costs in the year the asset is returned.

(iv) If the term of the lease is extended for an additional period of time at a reduced lease cost and the option to purchase still exists, the deferred charge may be included in capital-related costs to the extent of increasing the reduced rental to an amount not in excess of the cost of ownership.

(v) If the term of the lease is extended for an additional period of time at a reduced lease cost and the option to purchase no longer exists, the deferred charge may be included in capital-related costs to the extent of increasing the reduced rental to a fair rental value.

(vi) If the lessee becomes the owner of the leased asset (either by operation of the lease or by other means), the amount considered as depreciation, for the purpose of having computed the limitation on rental charges in paragraph (b)(9)(i) of this section, must be used in calculating the limitation on adjustments for the purpose of determining any gain or loss under § 413.134(f) upon disposal of an asset.

(c) Betterments and improvements. (1) Betterments and improvements are changes which extend the estimated useful life of an asset at least two years beyond its original estimated useful life, or increase the productivity of an asset significantly over its original productivity.

(2) A provider must capitalize and prorate the costs of betterments and improvements over the remaining estimated useful life of the asset, as modified by the betterment or improvement.

(d) Minor equipment. A provider must include in its capital-related costs the costs of minor equipment that are capitalized rather than charged off to expense if—

(1) The net book value of minor equipment at the time the provider enters the program is prorated over three years (that is, one-third of the net book value is written off each year), and new purchases are also prorated over a 3-year period; or

(2) The cost of minor equipment is prorated over their actual useful lives.

(e) Insurance. (1) A provider must include in its capital-related costs the costs of insurance on depreciable assets used for patient care or insurance that provides for the payment of capital-related costs during business interruption.

(2) If an insurance policy also provides protection for other than the replacement of depreciable assets or to pay capital-related costs in the case of business interruption insurance, only that portion of the premium related to the replacement of depreciable assets or to pay capital-related costs in the case of business interruption insurance is includable in capital-related costs.

(f) Debt premiums and debt discounts. Debt premiums or debt discount are applied as adjustments to capital-related costs if the associated debt is incurred for acquiring land or depreciable assets used for patient care or for refinancing existing debt for which the original purpose was to acquire land or depreciable assets used for patient care.

(g) Interest expense. (1) A provider must include in its capital-related costs interest expense, as described in § 413.153, if such expense is incurred in—

(i) Acquiring land or depreciable assets (either through purchase or lease) used for patient care; or

(ii) Refinancing existing debt, if the original purpose of the refinanced debt was to acquire land or depreciable assets used for patient care.

(2) If investment income offset is required under § 413.153(b)(2)(iii), only that portion of investment income that bears the same relationship to total investment income, as the portion of capital-related interest expense bears to total interest expense, is offset against capital-related costs.

(h) Costs of supplying organizations—

(1) Supplying organizations related to the provider. (i) If the supplying organization is related to the provider within the meaning of § 413.17, except as provided in paragraph (g)(1)(iii) of this section, a provider’s capital-related costs include the capital-related costs of the supplying organization.
(ii) If the costs of the services, facilities or supplies being furnished exceed the open market price, or if the provisions of §413.17(d) apply, no part of the cost to the provider of the services, facilities, or supplies are considered capital-related costs, unless the services, facilities, or supplies would otherwise be considered capital-related.

(2) Supplying organizations not related to the provider. If the supplying organization is not related to the provider within the meaning of §413.17, no part of the charge to the provider may be considered a capital-related cost (unless the services, facilities, or supplies are capital-related in nature) unless—

(i) The capital-related equipment is leased or rented (as described in paragraph (b) of this section) by the provider;

(ii) The capital-related equipment is located on the provider’s premises, or is located offsite and is on real estate owned, leased or rented by the provider; and

(iii) The capital-related portion of the charge is separately specified in the charge to the provider.

(i) Costs excluded from capital-related costs. The following costs are not capital-related costs. The extent to which they are allowable, they must be included in determining each provider’s operating costs:

(1) Costs incurred for the repair or maintenance of equipment or facilities.

(2) Amounts included in rentals or lease payments for repair or maintenance agreements.

(3) Interest expense incurred to borrow working capital (for operating expenses).

(4) General liability insurance or any other form of insurance to provide protection other than for the replacement of depreciable assets or to pay capital-related costs in the case of business interruption.

(5) Taxes other than those assessed on the basis of some valuation of land or depreciable assets used for patient care. (Taxes not related to patient care, such as income taxes, are not allowable, and are therefore not included among either capital-related or operating costs.)

(6) The costs of minor equipment that are charged off to expense rather than capitalized as described in paragraph (d) of this section.

(7) The costs incurred for maintenance and repair insurance agreements (commonly referred to as maintenance agreements).

(j) Reduction to capital-related costs. (1) Except for sole community hospitals and critical access hospitals, the amount of capital-related costs of all hospital outpatient services is reduced by—

(i) 15 percent for portions of cost reporting periods occurring on or after October 1, 1989, through September 30, 1991; and

(ii) 10 percent for portions of cost reporting periods occurring on or after October 1, 1991, and until the first date that the prospective payment system under part 419 of this chapter is implemented.

(2) For purposes of determining the blended payment amounts for hospital outpatient services under §413.118 and §413.122, the reduction is applicable only to the hospital-specific portion of the blended amounts.

§413.134 Depreciation: Allowance for depreciation based on asset costs.

(a) Principle. An appropriate allowance for depreciation on buildings and equipment used in the provision of patient care is an allowable cost. The depreciation must be—

(1) Identifiable and recorded in the provider’s accounting records;

(2) Based on the historical cost of the asset, except as specified in paragraph (j) of this section regarding donated assets; and

(3) Prorated over the estimated useful life of the asset using—

(i) The straight-line method; or

(ii) Accelerated depreciation under a declining balance method (not to exceed double the straight-line rate) or the sum-of-the-years’ digits method in the following situations:

(A) Depreciable assets for which accelerated depreciation was used for Medicare purposes before August 1,
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1970, including those assets for which a timely request to change from straight-line depreciation to accelerated depreciation was received by an intermediary before August 1, 1970;

(B) Depreciable assets acquired before August 1, 1970, if no election to use straight-line or accelerated depreciation was in effect on August 1, 1970, and the provider was participating in the program on August 1, 1970;

(C) Depreciable assets of a provider if construction of such depreciable asset began before February 5, 1970, and the provider was participating in the program on February 5, 1970; or

(D) Depreciable assets of a provider if a valid written contract was entered into by a provider participating in the program before February 5, 1970, for construction, acquisition, or for the permanent financing thereof, and such contract was binding on a provider on February 5, 1970, and at all times thereafter; or

(iii) A declining balance method, not to exceed 150 percent of the straight-line rate, for a depreciable asset acquired after July 31, 1970; however, this declining balance method may be used only if the cash flow from depreciation on the total assets of the institution during the reporting period, including straight-line depreciation on the assets in question, is insufficient (assuming funding of available capital not required currently for amortization and assuming reasonable interest income on such funds) to supply the funds required to meet the reasonable principal amortization schedules on the capital debts related to the provider’s total depreciable assets. For each depreciable asset for which a provider requests authorization to use a declining balance method for Medicare reimbursement purposes, but not to exceed 150 percent of the straight-line rate, the provider must demonstrate to the intermediary’s satisfaction that the required cash flow need exists. For each depreciable asset in which a provider justifies the use of accelerated depreciation, the intermediary must give written approval for the use of a depreciation method other than straight-line before basing any interim payment on this accelerated depreciation or making its reasonable cost determination which includes an allowance for such depreciation.

(b) General rules—(1) Historical cost. Historical cost is the cost incurred by the present owner in acquiring the asset.

(i) All providers—(A) Depreciable assets acquired after July 31, 1970 and before December 1, 1997. For depreciable assets acquired after July 31, 1970 and before December 1, 1997, and for a hospital or an SNF, acquired before July 18, 1984, the historical cost may not exceed the lower of current reproduction cost adjusted for straight-line depreciation over the life of the asset to the time of the purchase or the fair market value of the asset at the time of its purchase.

(B) Depreciable assets acquired on or after December 1, 1997. For depreciable assets acquired on or after December 1, 1997, the historical cost of the asset that will be recognized under this program must not exceed the historical cost less depreciation allowed to the owner of record as of August 5, 1997 (or if an asset did not exist as of August 5, 1997, the first owner of record after August 5, 1997). For this paragraph (b)(1)(i)(B), the following apply:

(1) An asset that was not in existence as of August 5, 1997 includes an asset that physically existed but was not owned by a provider participating in the Medicare program as of that date.

(2) The acquisition cost to the owner of record is subject to the limitation on historical costs described in paragraphs (g) (1), (2), and (3) of this section, and is reduced by any depreciation taken by the owner of record. The limitation on historical cost is also applied to the purchase of land, which is a capital asset that is neither depreciable nor amortizable under any circumstances.

(3) Acquisition cost to the owner of record includes the costs of betterment or improvements that extend the estimated useful life of an asset at least 2 years beyond its original estimated useful life or that increase the productivity of an asset significantly over its original productivity.
(4) For assets acquired prior to a provider’s entrance into the Medicare program, the acquisition cost to the owner of record is the historical cost when acquired, rather than when the provider entered the program.

(5) For assets subject to the optional depreciation allowance as described in §413.139, the acquisition cost to the owner of record is the historical cost established for those assets when the provider changed to actual depreciation as described in §413.139(e). If the provider did not change to actual depreciation, as described in §413.139(e), for optional allowance assets, the acquisition cost to the owner of record is based on the provider’s recorded historical cost of the asset when acquired. If the provider has no historical cost records for optional allowance assets, the acquisition cost to the owner of record is established by appraisal.

(6) The historical cost of an asset acquired on or after July 18, 1984 may not include costs attributable to the negotiation or settlement of the sale or purchase (by acquisition, merger, or consolidation) of any capital asset for which any payment was previously made under the Medicare program. The costs to be excluded include, but are not limited to, appraisal costs (except those incurred at the request of the intermediary under paragraph (f)(2)(iv) of this section), legal fees, accounting and administrative costs, travel costs, and the costs of feasibility studies.

(ii) Hospitals and SNFs only. (A) For assets acquired on or after July 18, 1984 and before December 1, 1997 and not subject to an enforceable agreement entered into before July 18, 1984, historical cost may not exceed the lowest of the following:

1. The allowable acquisition cost of the asset to the owner of record as of July 18, 1984 (or, in the case of an asset not in existence as of July 18, 1984, the first owner of record of the asset after that date);

2. The acquisition cost of the asset to the new owner;

3. The fair market value of the asset on the date of acquisition.

(B) For purposes of applying paragraph (b)(2)(ii)(A) of this section, an asset not in existence as of July 18, 1984 includes any asset that physically existed, but was not owned by a hospital or SNF participating in the Medicare program as of July 18, 1984.

(C) The acquisition cost to the owner of record is subject to any limitation on historical costs described in paragraphs (b)(1)(i) or (g)(1) and (2) of this section, and is not reduced by any depreciation taken by the owner of record. This limitation on historical cost is also applied to the purchase of land, a capital asset that is neither depreciable nor amortizable under any circumstances. (See §§413.153(d) and 413.157(b) for application of the limitation to the cost of land for purposes of determining allowable interest expense and return on equity capital or proprietary providers.)

(D) Acquisition cost to the owner of record includes the costs of betterments or improvements that extend the estimated useful life of an asset at least two years beyond its original estimated useful life or increase the productivity of an asset significantly over its original productivity.

(E) For assets acquired prior to a provider’s or SNF’s entrance into the Medicare program, the acquisition cost to the owner of record is the historical cost of the asset when acquired, rather than when the hospital or SNF entered the program.

(F) For assets subject to the optional depreciation allowance as described in §413.139, the acquisition cost to the owner of record is the historical cost established for those assets when the hospital or SNF changed to actual depreciation as described in §413.139(e). If the hospital or SNF did not change to actual depreciation, as described in §413.139(e), for optional allowance assets, the acquisition cost to the owner of record is established by reference to the hospital’s or SNF’s recorded historical cost of the asset when acquired. If the hospital or SNF has no historical cost records for optional allowance assets, the acquisition cost to the owner of record is established by appraisal.

(G) The historical cost of an asset acquired on or after July 18, 1984 may not include costs attributable to the negotiation or settlement of the sale or purchase (by acquisition, merger, or consolidation) of any capital asset for which any payment was previously made under the Medicare program.
made under the Medicare program. The costs to be excluded include, but are not limited to, appraisal costs (except those incurred at the request of the intermediary under paragraph (f)(2)(iv) of this section), legal fees, accounting and administrative costs, travel costs, and the costs of feasibility studies.

(iii) Hospital-based providers other than SNFs and SNF-based providers. For changes of ownership that involve assets of a hospital-based provider other than a SNF, or assets of a SNF-based provider, the provisions of paragraph (b)(1)(ii) of this section are not applicable. A reasonable allocation of the purchase price must be made, so that the hospital-based provider other than a SNF, or a SNF-based provider, is not affected by the limitations described in paragraph (b)(1)(ii) of this section. The historical cost of assets of providers other than hospitals and SNFs is governed by paragraph (b)(1)(i) of this section.

(2) Fair market value. Fair market value is the price that the asset would bring by bona fide bargaining between well-informed buyers and sellers at the date of acquisition. Usually the fair market price is the price that bona fide sales have been consummated for assets of like type, quality, and quantity in a particular market at the time of acquisition.

(3) The straight-line method. Under the straight-line method of depreciation, the cost or other basis (for example, fair market value in the case of donated assets) of the asset, less its estimated salvage value, if any, is determined first. Then this amount is distributed in equal amounts over the period of the estimated useful life of the asset.

(4) Declining balance method. Under the declining balance method, the annual depreciation allowance is computed by multiplying the undepreciated cost of the asset each year by a uniform rate up to double the straight-line rate or 150 percent, as the case may be (see paragraph (a)(3) of this section for limitations on use of accelerated methods of depreciation).

(5) Sum-of-the-years’ digits method. Under the sum-of-the-years’ digits method, the annual depreciation allowance is computed by multiplying the depreciable cost basis (cost less salvage value) by a constantly decreasing fraction. The numerator of the fraction is represented by the remaining years of useful life of the asset at the beginning of each year, and the denominator is always represented by the sum of the years’ digits of useful life at the time of acquisition.

(6) Current reproduction cost. Current reproduction cost is the cost at current prices, in a particular locality or market area, of reproducing an item of property or a group of assets. Where depreciable assets are concerned, this means the reasonable cost to have built, reproduce in kind, or, in the case of equipment or similar assets, to purchase in the competitive market.

(7) Useful life. The estimated useful life of a depreciable asset is its normal operating or service life to the provider, subject to the provisions in paragraph (b)(7)(i) of this section. Factors to be considered in determining useful life include normal wear and tear; obsolescence due to normal economic and technological changes; climatic and other local conditions; and the provider’s policy for repairs and replacement.

(i) Initial selection of useful life. In selecting a proper useful life for computing depreciation under the Medicare program, providers must use the useful life guidelines published by HCFA. If HCFA has not published applicable useful life guidelines, providers must use—

(A) The edition of the American Hospital Association useful life guidelines, as specified in HCFA Medicare program manuals; or

(B) A different useful life specifically requested by the provider and approved by the intermediary. A different useful life may be approved by the intermediary if the provider’s request is properly supported by acceptable factors that affect the determination of useful life. However, such factors as an expected early sale, retirement, demolition or abandonment of an asset, or termination of the provider from the Medicare program may not be used.

(ii) Application of guidelines. The provisions concerning the selection of useful life guidelines described in paragraph (b)(7)(i) of this section apply to assets acquired on or after January 1,
1981. For assets acquired before January 1, 1981, providers must use the useful life guidelines published by the American Hospital Association in its 1973 edition of Chart of Accounts for Hospitals, or those published by the Internal Revenue Service, or those approved for use by intermediaries as provided in paragraph (b)(7)(i)(B) of this section.

(iii) Changing useful life. A change in the estimated useful life may be made if clear and convincing evidence justifies a redetermination of the useful life used by the provider. Such a change must be approved by the intermediary in writing, and the factors cited in paragraphs (b)(7) and (b)(7)(i) of this section are applicable in making such redeterminations of useful life. If the request is approved, the change is effective with the reporting period immediately following the period in which the provider’s request is submitted for approval.

(8) Donated asset. An asset is considered donated when the provider acquires the asset without making payment in the form of cash, new debt, assumed debt, property or services. Except as provided in paragraph (j)(3) of this section, if a provider makes payment in any form to acquire an asset, the payment is considered the purchase price for the purpose of determining allowable historical cost.

(9) Net book value. The net book value of an asset is the depreciable basis used for the Medicare program by the asset’s last participating owner less depreciation recognized under the Medicare program.

(c) Recording of depreciation. Appropriate recording of depreciation includes the identification of the depreciable assets in use, the assets’ historical costs, the assets’ dates of acquisition, the method of depreciation, estimated useful lives, and the assets’ accumulated depreciation.

(1) Depreciation methods—(1) General. Proration of the cost of an asset over its useful life is allowed on the straight-line method, or, when permitted under paragraph (a)(3) of this section, the declining balance or the sum-of-the-years’ digits methods. One method may be used on a single asset or group of assets and another method on others. In applying the declining balance or sum-of-the-years’ digits method to an asset that is not new, the undepreciated cost of the asset is treated as the cost of a new asset in computing depreciation.

(2) Change in method. Prior to August 1, 1970, a provider may change from the straight-line method to an accelerated method or vice versa, upon advance approval from the intermediary on a prospective basis with the request being made before the end of the first month of the prospective reporting period. Only one such change with respect to a particular asset may be made by a provider. Effective with August 1, 1970, a provider may only change from an accelerated method or optional method (see §413.139) to the straight-line method. Such a change may be made without intermediary approval and the basis for depreciation is the undepreciated cost reduced by the salvage value. Thereafter, once straight-line depreciation is selected for a particular asset, an accelerated method may not be established for that asset.

(3) Recovery of accelerated depreciation—(i) General. If a provider who has used an accelerated method of depreciation for any of its assets terminates participation in the program, or if the Medicare proportion of its allowable costs decreases so that cumulatively substantially more depreciation was paid than would have been paid using the straight-line method of depreciation, the excess of reimbursable cost determined by using accelerated depreciation methods and paid under the program over the reimbursable cost that would have been determined and paid under the program by using the straight-line method of depreciation, will be recovered as an offset to current reimbursement due or, if the provider has terminated participation in the program, as an overpayment. In this determination of excess payment, recognition will be given to the effects the adjustment to straight-line depreciation would have on the return on equity capital and on the allowance in lieu of specific recognition of other costs in the respective years.

(ii) Transaction between related organizations—(A) General. If the termination of the provider agreement is due
to a change in provider ownership, as defined in §489.18 of this chapter, resulting from a transaction between related organizations, as defined in §413.17, and the criteria in paragraph (b) of this section are met, the excess of reimbursable cost, as determined in paragraph (d)(3)(i) of this section may not be recovered if there is a continuation of participation by the facility in the Medicare program.

(B) Criteria. The following criteria must be met if the recovery of excess reimbursable cost is not to be made:

(1) The termination of the provider agreement is due to a change in ownership of the provider resulting from a transaction between related organizations.

(2) The successor provider continues to participate in the Medicare program.

(3) Control and the extent of the financial interest of the owners of the provider before and after the termination remain the same; that is, the successor owners acquire the same percentage of control or financial investment as the transferors had.

(4) All assets and liabilities of the terminated provider are transferred to the related successor participating provider.

(C) Effect of transaction. In transactions meeting the criteria specified in paragraph (d)(3)(ii)(B) of this section, the provision concerning recovery of excess reimbursable cost (§413.134(d)(3)(i)) is not applied, and the transaction is treated as follows:

(1) The successor provider must record the historical cost and accumulated depreciation and the method of depreciation recognized under the Medicare program, and these are considered as incurred by the successor provider for Medicare purposes.

(2) The Medicare program’s utilization of the terminated provider is considered as having been incurred by the successor provider for Medicare purposes.

(3) The equity capital of the terminated provider as of the closing of its final cost reporting period must be wholly contained in the equity capital of the successor provider as of the beginning of its first cost reporting period.

(e) Funding of depreciation. Although funding of depreciation is not required, it is strongly recommended that providers use this mechanism as a means of conserving funds for replacement of depreciable assets. Funded depreciation account funds must be placed in readily marketable investments of the type that assures the availability and conservation of the funds. Additions to the funded depreciation account must remain in the account for at least 6 months to be considered valid funding transactions.

(1) Incentive. As an incentive for funding, investment income on funded depreciation is not treated as a reduction of allowable interest expense provided such investment income is deposited in, and becomes part of, the funded depreciation account at the time of receipt by the provider. Investment income earned on deposits before the 6-month period elapses are not offset unless the deposits are withdrawn for an improper purpose during this period. If a provider transfers assets of the funded depreciation account to a related organization (for example, pooling of several chain organization providers’ funded depreciation accounts at the chain home office for investment purposes), these assets shall be treated as the provider’s funds and are subject to all the requirements specified in paragraph (e) of this section.

(2) Availability of funded depreciation.

(i) HCFA considers funded depreciation available for use in the acquisition or replacement of depreciable assets related to patient care unless the funded depreciation funds have been committed by contract for the acquisition of depreciable assets related to the furnishing of patient care or for other capital purposes related to patient care.

(ii) Borrowing for a purpose for which funded depreciation account funds should have been used makes the borrowing unnecessary to the extent that funded depreciation account funds were available at the time of the borrowing. Available funds in the funded depreciation account, to the extent of the unnecessary borrowing, are called “tainted” funds. Interest expense incurred on borrowing for a capital purpose is not an allowable cost to the extent that
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funded depreciation account funds were available at the time of the borrowing.

(iii) A provider can remove the “unnecessary” characterization of borrowing, and thereby cure tainted funded depreciation, by using the tainted funds for a proper purpose described in paragraph (e)(3)(i) of this section. However, any funded depreciation that existed at the time of the unnecessary borrowing and is not classified as tainted must be used before any of the tainted funds.

(iv) When only a portion of the borrowing is considered unnecessary under paragraph (e)(2)(ii) of this section, subsequent repayments of such borrowing from general funds are applied first to the allowable portion of the borrowing and then, when all of the allowable borrowing is repaid, to the unallowable portion of the borrowing. When funds from the funded depreciation account are used for the repayment of the unnecessary borrowing, an equivalent amount of tainted funds is cured without regard to the provisions of paragraphs (e)(2)(ii) and (e)(3)(i)(C) of this section. Similarly, where general funds are used to pay for the unallowable borrowing after the necessary borrowing has been repaid, an equivalent amount of tainted funded depreciation is cured without regard to the provisions of paragraphs (e)(2)(ii) and (e)(3)(i)(C) of this section.

(3) Withdrawals of funded depreciation—(i) Proper withdrawals. (A) Withdrawals from funded depreciation are considered proper if made either for the acquisition or replacement of depreciable assets related to the furnishing of patient care or for other capital purposes related to patient care.

(B) First-in, first-out basis. Proper withdrawals from funded depreciation are made on a first-in, first-out basis.

(C) Exception. If HCFA determines that a borrowing is unnecessary because of the existence of available funded depreciation, and additional deposits have been made to funded depreciation after the occurrence of the unnecessary borrowing, withdrawals made after the date of the additional deposits are deemed to be made on a last-in, first-out basis.

(ii) Improper withdrawals. (A) Withdrawals from funded depreciation that do not meet the requirements for proper withdrawals under the provisions in paragraph (e)(3)(i)(A) of this section are considered improper withdrawals.

(B) Improper withdrawals from funded depreciation are made on a last-in, first-out basis. If improper withdrawals are made, interest expense is reduced in accordance with section §413.153(c)(3).

(C) Improper withdrawals will result in the offset of otherwise allowable interest expense under the offset provisions in §413.153(c)(3).

(4) Loans from funded depreciation. (i) When the general fund of the provider borrows from the funded depreciation to obtain working capital for normal operating expenses to furnish patient care, interest incurred by the general fund is an allowable operating cost only if the interest expense is supported by documents that evidence that the funds were borrowed and that payment of interest and repayment of the funds are required, is separately identified in the provider’s accounting records, and meets the necessary and proper tests described in §413.153(b)(2) and (b)(3). However, if the general fund of the provider borrows from the funded depreciation account to acquire depreciable assets used in furnishing patient care, or for other capital purposes related to patient care, interest expense paid by the general fund to the funded depreciation account is not an allowable cost. Providers are expected to use the funded depreciation for these purposes.

(ii) Loans from funded depreciation to the general fund are considered investments of funded depreciation, but do not have to meet the readily marketable test described in paragraph (e) of this section. Loans made from funded depreciation are subject to the requirement that funded depreciation must be available for the acquisition of depreciable assets used to furnish patient care, or for other capital purposes related to patient care. Costs incurred to secure lines of credit from lending institutions to ensure such availability are not allowable costs.

(iii) Funding of depreciation from general funds will not be recognized to the extent of any outstanding loans from the funded depreciation account...
to the general fund. Deposits from the general fund into the funded depreciation account must be first applied to reduce any loans outstanding from the funded depreciation to the general fund. When the loans are repaid in full, general funds deposited in the funded depreciation account are considered as repayments of the general fund. Therefore, any subsequent interest expense of the general fund paid to the funded depreciation fund is not an allowable cost.

(iv) A provider may loan its funded depreciation to a related organization for any purpose subject to the following conditions:

(A) Authorization for such a loan by the provider's appropriate managing body of the provider, such as Board of Trustees or Board of Directors, must be on file.

(B) The funded depreciation loaned must remain available, as specified in paragraph (e)(2) of this section, to the provider making the loan. Costs incurred for lines of credit to assure such availability are not allowable costs. During the period of time that the loan is outstanding, if the provider making the loan resorts to outside borrowing for a purpose for which its funded depreciation should have been used, interest expense on an amount of the outside borrowing up to the amount of the funded depreciation that should have been available would be disallowed as unnecessary.

(C) Such loans shall be considered investments of the provider's funded depreciation, but the requirement that funded depreciation be invested in readily marketable investments as required in paragraph (e) of this section is relaxed for such loans.

(D) The funded depreciation account must earn interest on such loans at a rate that does not exceed the rate that would be charged for a comparable loan from an independent lending institution. This investment income will not be used to reduce the provider's interest expense if all the other conditions in paragraph (e) of this section are met. If the entity borrowing the funds is another provider participating in the Medicare program, the interest expense incurred on such loans would be allowable if the loan meets all of the interest expense requirements specified in §413.153. (For purposes of §413.153(b)(3)(ii), such loans are not considered to be with a related lender.)

(f) Gains and losses on disposal of assets—

(1) General. Depreciable assets may be disposed of through sale, scrapping, trade-in, exchange, demolition, abandonment, condemnation, fire, theft, or other casualty. If disposal of a depreciable asset, including the sale or scrapping of an asset before December 1, 1997, results in a gain or loss, an adjustment is necessary in the provider's allowable cost. (No gain or loss is recognized on either the sale or the scrapping of an asset that occurs on or after December 1, 1997.) The amount of a gain included in the determination of allowable cost is limited to the amount of depreciation previously included in Medicare allowable costs. The amount of a loss to be included is limited to the undepreciated basis of the asset permitted under the program. The treatment of the gain or loss depends upon the manner of disposition of the asset, as specified in paragraphs (f)(2) through (6) of this section. The gain or loss on the disposition of depreciable assets has no retroactive effect on a proprietary provider's equity capital for years prior to the year of disposition.

(2) Bona fide sale or scrapping before December 1, 1997. For the bona fide sale or scrapping of depreciable assets before December 1, 1997, the following apply:

(i) Except as specified in paragraph (f)(3) of this section, gains and losses realized from the bona fide sale or scrapping of depreciable assets are included in the determination of allowable cost only if the sale or scrapping occurs while the provider is participating in Medicare. The extent to which such gains and losses are included is calculated by prorating the basis for depreciation of the asset in accordance with the proportion of the asset's useful life for which the provider participated in Medicare. For purposes of this paragraph (f)(2)(i), scrapping refers to the physical removal from the provider's premises of tangible personal properties that are no longer useful for their intended purpose and are only salable for their scrap or junk value.
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(ii) If the total amount of gains or losses realized from bona fide sales or scrapping does not exceed $5,000 within the cost reporting period or if the provider's cumulative utilization under the Medicare program is less than 5 percent, the net amount of gains or losses realized from sale or scrapping will be allowed as a depreciation adjustment in the period of disposal. For purposes of this paragraph (f)(2)(ii), the provider's cumulative Medicare utilization percentage is determined by comparing the cumulative total of the Medicare inpatient days for all reporting periods in which depreciation on the asset disposed of was claimed under the Medicare program to the cumulative total of inpatient days of the participating provider for the same reporting periods.

(iii) If the conditions specified in paragraph (f)(2)(ii) of this section are not met, the adjustment to reimbursable cost in the reporting period of asset disposition is calculated as follows:

(A) The total amount of gains or losses shall be allocated to all reporting periods under the Medicare program, based on the ratio of the depreciation allowed on the assets in each reporting period to the total depreciation allowed under the Medicare program.

(B) The results of this allocation are multiplied by the ratio of Medicare reimbursable cost to total allowable cost for each reporting period.

(C) The results of this multiplication are then added.

(D) Effective for cost reporting periods beginning on or after October 1, 1991, no adjustment will be made for the portion of gains or losses allocated to inpatient hospital services for which the hospital was paid under the fully prospective payment methodology as described in § 412.340 of this chapter or under the hold-harmless methodology based on the Federal rate as described in § 412.344(a)(1) of this chapter for new capital costs or in § 412.344(a)(2) of this chapter.

(iv) If a provider sells more than one asset for a lump sum sales price, the gain or loss on the sale of each depreciable asset must be determined by allocating the lump sum sales price among all the assets sold, in accordance with the fair market value of each asset as it was used by the provider at the time of sale. If the buyer and seller cannot agree on an allocation of the sales price, or if they do agree but there is insufficient documentation of the current fair market value of each asset, the intermediary for the selling provider will require an appraisal by an independent appraisal expert to establish the fair market value of each asset and will make an allocation of the sales price in accordance with the appraisal.

(3) Sale within 1 year after termination. Gains and losses realized from a bona fide sale of depreciable assets within 1 year immediately following the date on which the provider terminates participation in the Medicare program are also included in the determination of allowable cost, in accordance with the procedure specified in paragraph (f)(2) of this section. However, if several assets are sold for a lump sum sales price, the determination of fair market value must be based on the appraised value of the assets as they were last used by the provider while participating in the Medicare program.

(4) Exchange, trade-in or donation. Gains or losses realized from the exchange, trade-in, or donation of depreciable assets are not included in the determination of allowable cost. When the disposition of an asset is by means of exchange or trade-in, the historical cost of the new asset is the sum of the undepreciated cost of the asset disposed of and the additional cash or other assets transferred (or to be transferred) to acquire the new asset. However, if the asset disposed of was acquired by the provider before its participation in the Medicare program and the sum of the undepreciated cost and the cash or other assets transferred (or to be transferred) exceed the list price or fair market value of the new asset, the historical cost of the new asset is limited to the lower of its list price or fair market value.

(5) Demolition or abandonment. (i) For purposes of this section, the term "abandonment" means the permanent retirement of an asset for any future purpose, not merely the provider's ceasing to use the asset for patient
care purposes. To claim an abandonment under the Medicare program, the provider must have relinquished all rights, title, claim, and possession of the asset with the intention of never reclaiming it or resuming its ownership, possession, or enjoyment.

(ii) If losses resulting from the demolition or abandonment of depreciable assets do not exceed $5,000 within the cost-reporting period, the losses are to be allowed in the period of disposal.

(iii) If losses exceed $5,000 and, at the date of disposition, the demolished or abandoned assets are at least 80 percent depreciated as computed under the straight-line method, such losses are includable in the determination of allowable cost under the Medicare program in the period of disposal and the procedure provided in paragraph (f)(2)(iii) of this section must be used in determining the adjustment to reimbursable cost.

(iv) Losses in excess of $5,000 resulting from the demolition or abandonment of assets, which at the date of disposition are not 80 percent depreciated as computed under the straight-line method, must be capitalized as a deferred charge and amortized as follows:

(A) If the State Health Planning and Development Agency (SHPDA) designated under section 1521 of the Public Health Service Act approves the demolition or abandonment of a depreciable asset as being consistent with the health systems plan of the health service area in which the provider is located, the net loss realized shall be capitalized as a deferred charge and amortized over the remaining life of the demolished or abandoned asset, or at the rate of $5,000 per year, whichever is greater. If no SHPDA exists or if such agency is unable or unwilling to perform this function, the provider must submit a request for approval to the intermediary. The intermediary, after reviewing this request and before issuing the approval, will submit the request along with its recommendation to the appropriate Regional Office for its approval.

(B) If a provider fails to obtain approval as specified in paragraph (f)(5)(iv)(A) of this section, a loss is not allowable unless the demolished or abandoned asset is replaced. If the asset is replaced, the loss resulting from the unapproved demolition or abandonment must be capitalized as a deferred charge and amortized over the estimated useful life of the replacement asset or at the rate of $5,000 per year, whichever is greater.

(v) If a loss resulting from the demolition or abandonment is deferred and amortized and the provider terminates its participation in the Medicare program or ceases to use a replacement asset in the provision of patient care services, the unamortized deferred charge remaining at that time must not be included in determining allowable cost under the Medicare program.

(vi) Losses on demolition must include the demolition cost incurred by the provider for razing and removal of the asset, less any salvage value recovered by the provider. However, if a provider demolishes a depreciable asset for the purpose of preparing land for future sale, the net demolition cost incurred by the provider (razing and removal costs less salvage recovered) is considered a capital expenditure and added to the historical basis of the land.

(vii) If a provider purchases land on which there is a building, no depreciation will be allowed under the Medicare program unless the building is used in providing patient care. If the building is demolished, the entire purchase price and demolition cost shall be considered the historical cost of the land. If the building is used for patient care, but demolished within 5 years of purchase, the entire purchase price, less allowed depreciation, plus demolition cost will be considered the historical cost of the land.

(B) Involuntary conversion. (i) Losses resulting from the involuntary conversion of depreciable assets, such as condemnation, fire, theft, or other casualty, are generally included in the determination of allowable cost on a deferred basis if the asset is restored or replaced. However, losses resulting from a provider’s imprudent management of its depreciable assets, such as the failure to obtain proper insurance coverage, are not included in the determination of allowable cost.

(ii) The net allowable loss from involuntary conversion must consist of the...
undepreciated cost of unrecovered book value of the asset, less amounts received from insurance proceeds gifts, and grants received from local, State, or Federal government, or any other source as a result of the involuntary conversion.

(iii) If the asset is replaced and the net allowable loss in any cost-reporting period does not exceed $5,000, the entire amount must be included in allowable cost in the period in which the loss is incurred. If the asset is replaced and the net allowable loss in any cost-reporting period exceeds $5,000, the loss must be capitalized as a deferred charge and amortized over the useful life of the replacement or restored asset. If a replaced or restored asset ceases to be used in the provision of patient care services or the provider terminates its participation in the Medicare program, the unamortized deferred charge remaining at that time will not be included in determining allowable cost under the Medicare program.

(iv) If the provider fails to replace or restore an involuntarily converted asset, the loss is not included in determining allowable cost. However, if the provider intends to replace or restore the asset but is unable to do so because the designated SHPDA finds such replacement or restoration to be inconsistent with the health systems plan of the provider’s health service area, the loss is allowable so long as the provider continues to participate in Medicare. In this case, the loss must be capitalized as a deferred charge and amortized over the remaining life of the involuntarily converted asset, or at the rate of $5,000 per year, whichever is greater.

(v) If a gain is realized from an involuntary conversion of depreciable assets, the net amount realized reduces the basis of the restored or replacement asset. If the asset is not restored or replaced, the gain is to be treated in accordance with paragraph (f)(2) of this section.

(7) Effect on equity capital. The unrecovered loss entered on the books of the provider as a deferred charge, in accordance with paragraphs (f)(5) and (6) of this section, is not includable in the computation of equity capital under §413.157.

(8) Sale of replacement or restored assets. If a provider sells a replacement or restored asset while participating in the Medicare program or within 1 year immediately following the date on which it terminates its participation in the Medicare program, the unrecovered loss entered on the books of the provider as a deferred charge in accordance with paragraphs (f)(5) and (6) of this section will not be included in determining the gain or loss realized from the sale of the replacement or restored asset. However, if the sale of such asset is made to a related organization, as defined in §413.17, and the purchasing organization continues as a provider in the Medicare program, the remaining deferred charge representing the unrecovered depreciable basis of the demolished, abandoned or destroyed asset must continue to be amortized over the remaining expected useful life of the replacement or restored asset. If the sale is made to an unrelated organization, further amortization of the deferred charge is not allowed.

(g) Establishment of cost basis on purchase of facility as an ongoing operation—(1) Assets acquired after July 1, 1966 and before August 1, 1970. The cost basis for the assets of a facility purchased as an ongoing operation after July 1, 1966, and before August 1, 1970, is the lowest of the—

(i) Total price paid for the facility by the purchaser, as allocated to the individual assets of the facility;

(ii) Total fair market value of the facility at the time of the sale, as allocated to the individual assets; or

(iii) Combined fair market value of the individually identified assets at the time of the sale.

(2) Assets acquired after July 31, 1970 and, for hospitals and SNFs, before July 18, 1984. For depreciable assets acquired after July 31, 1970 and, for hospitals and SNFs, before July 18, 1984, in addition to the limitations specified in paragraph (g)(1) of this section, the cost basis of the depreciable assets may not exceed the current reproduction cost depreciated on a straight-line basis over the life of the asset to the time of the sale.

(3) Assets acquired by hospitals and SNFs on or after July 18, 1984 and not
subject to an enforceable agreement entered into before that date. Subject to paragraphs (b)(1)(ii) (B) through (G) and (b)(1)(iii) of this section, historical cost may not exceed the lowest of the following:

(i) The allowable acquisition cost of the asset to the owner of record as of July 18, 1984 (or, in the case of an asset not in existence as of July 18, 1984, the first owner of record of the asset);

(ii) The acquisition cost to the new owner; or

(iii) The fair market value of the asset on the date of acquisition.

(2) Assets acquired by all providers on or after December 1, 1997. Subject to the provisions of paragraph (b)(1)(i)(A) of this section, the historical cost may not exceed the historical cost of the asset, as recognized under the Medicare program, less depreciation allowed, to the owner of record as of August 5, 1997 (or for an asset not in existence as of August 5, 1997, the first owner of record after August 5, 1997).

(5) Transactions other than bona fide. If the purchaser cannot demonstrate that the sale was bona fide, in addition to the limitations specified in paragraph (g)(1), (2), and (3) of this section, the purchaser’s cost basis may not exceed the seller’s cost basis, less accumulated depreciation.

(h) Sale and leaseback agreements and other lease transactions. (1) For sale and leaseback agreements for all providers, and for sale and leaseback agreements for hospitals and SNFs entered into before October 23, 1992, a provider may include in its allowable costs incurred rental charges, as specified in a sale and leaseback agreement with a non-related purchaser involving plant facilities or equipment, only if—

(i) The rental charges are reasonable based on consideration of rental charges of comparable facilities and market conditions in the area; the type, expected life, condition, and value of the facilities or equipment rented; and other provisions of the rental agreement;

(ii) Adequate alternate facilities or equipment that would serve the purpose are not or were not available at lower cost; and

(iii) The leasing was based on economic and technical considerations.

(2) If the conditions of paragraph (h)(1) of this section are not met, the amount a provider may include in its allowable costs as rental or lease expense under a sale and leaseback agreement may not exceed the amount that the provider would have included in its allowable costs had the provider retained legal title to the facilities or equipment such as interest expense on mortgages, taxes, depreciation, and insurance costs.

(3) For hospitals and SNFs entering into sale and leaseback agreements on or after October 23, 1992, the amount a provider may include in its allowable costs as rental or lease expense may not exceed the amount that the provider would have included in its allowable costs had the provider retained legal title to the facilities or equipment, such as interest expense on mortgages, taxes, depreciation, and insurance costs (the costs of ownership). This limitation applies both on an annual basis and over the useful life of the asset.

(i) If in the early years of the lease, the annual rental or lease costs are less than the annual costs of ownership, but in the later years of the lease the annual rental or lease costs are more than the annual costs of ownership, in the years that the annual rental or lease costs are more than the costs of ownership the provider may include in allowable costs annually the actual amount of rental or lease costs. The aggregate rental or lease costs included in allowable costs may not exceed the aggregate costs of ownership that would have been included in allowable costs over the useful life of the asset had the provider retained legal title to the asset.

(ii) If in the early years of the lease, the annual rental or lease costs exceed the annual costs of ownership, but in the later years of the lease the annual rental or lease costs are less than the annual costs of ownership, the provider may carry forward amounts of rental or lease costs that were not included in allowable costs in the early years of the lease due to the costs of ownership limitation, and include these amounts in allowable costs in the years of the lease when the annual rental or lease costs are less than the annual costs of
ownership. In any given year the amount of actual annual rental or lease costs plus the amount carried forward to that year may not exceed the amount of the costs of ownership for that year.

(iii) In the aggregate, the amount of rental or lease costs included in allowable costs may not exceed the amount of the costs of ownership that the provider could have included in allowable costs had the provider retained legal title to the asset.

(4) For lease transactions of all providers entered into before October 23, 1992, a lease that meets the following conditions establishes a virtual purchase:

(i) The rental charge exceeds rental charges of comparable facilities or equipment in the area.

(ii) The term of the lease is less than the useful life of the facilities or equipment.

(iii) The provider has the option to renew the lease at a significantly reduced rental, or the provider has the right to purchase the facilities or equipment at a price that appears to be significantly less than what the fair market value of the facilities or equipment would be at the time acquisition by the provider is permitted.

(5)(i) If a lease is a virtual purchase under paragraph (h)(4) of this section, the rental charge is includable in allowable costs if it had legal title to the asset (the cost of ownership), such as straight-line depreciation, insurance, and interest. For purposes of computing the limitation on allowable rental cost in this paragraph, a provider may not include accelerated depreciation.

(ii) The difference between the amount of rent paid and the amount of rent allowed as rental expense is considered a deferred charge and must be capitalized as part of the historical cost of the asset when the asset is purchased.

(iii) If an asset is returned to the owner instead of being purchased, the deferred charge may be expensed in the year the asset is returned.

(iv) If the term of the lease is extended for an additional period of time at a reduced lease cost and the option to purchase still exists, the deferred charge may be expensed to the extent of increasing the reduced rental to an amount not in excess of the cost of ownership.

(v) If the term of the lease is extended for an additional period of time at a reduced lease cost and the option to purchase no longer exists, the deferred charge may be expensed to the extent of increasing the reduced rental to a fair rental value.

(6) For lease transactions entered into on or after October 23, 1992, a lease that meets any one of the following conditions establishes a virtual purchase:

(i) The lease transfers title of the facilities or equipment to the lessee during the lease term.

(ii) The lease contains a bargain purchase option.

(iii) The lease term is 75 percent or more of the useful life of the facilities or equipment. This provision is not applicable if the lease begins in the last 25 percent of the useful life of the facilities or equipment.

(iv) The present value of the minimum lease payments (that is, payments to be made during the lease term, including bargain purchase option, guaranteed residual value, or penalties for failure to renew) equals 90 percent or more of the fair market value of the leased property. This provision is not applicable if the lease begins in the last 25 percent of the useful life of the facilities or equipment. The present value is computed using the lessee's incremental borrowing rate, unless the interest rate implicit in the lease is known and is less than the lessee's incremental borrowing rate, in which case, the interest rate implicit in the lease is used.

(7)(i) If a lease is a virtual purchase under paragraph (h)(6) of this section, the rental charge is includable in allowable costs only to the extent that it does not exceed the amount that the provider would have included in allowable costs if it had legal title to the asset (the costs of ownership), such as straight-line depreciation, insurance,
and interest. For purposes of computing the limitation on allowable rental cost as described in this paragraph, a provider may not include accelerated depreciation in its allowable costs.

(ii) The difference between the amount of rent paid and the amount of rent allowed as rental expense is considered a deferred charge and is capitalized as part of the historical cost of the asset when the asset is purchased.

(iii) If an asset is returned to the owner instead of being purchased, the deferred charge may be expensed in the year the asset is returned.

(iv) If the term of the lease is extended for an additional period of time at a reduced lease cost and the option to purchase still exists, the deferred charge may be expensed to the extent of increasing the reduced rental to an amount not in excess of the cost of ownership.

(v) If the term of the lease is extended for an additional period of time at a reduced lease cost and the option to purchase no longer exists, the deferred charge may be expensed to the extent of increasing the reduced rental to a fair rental value.

(vi) If the lessee becomes the owner of the leased asset (either by operation of the lease or by other means), the amount considered as depreciation, for the purpose of determining any gain or loss upon disposal of an asset under paragraph (f) of this section.

(j) Basis of assets donated to a provider—(1) Assets not used or depreciated under the Medicare program. If an asset has never been used or depreciated under the Medicare program and is donated to a provider, the basis for the purpose of calculating depreciation and equity capital (if applicable) is the fair market value of the asset at the time of donation.

(ii) Assets used or depreciated under the Medicare program. If an asset has been used or depreciated under the Medicare program and is donated to a provider, the basis for the purpose of calculating depreciation and equity capital (if applicable) is the lesser of—

(i) The fair market value at the time of donation; or
(ii) The net book value in the hands of the owner last participating in the Medicare program.

(3) Transfers of State hospitals to nonprofit corporations without monetary consideration. If a State transfers a hospital to a nonprofit corporation without monetary consideration on or after July 18, 1984, the depreciable basis of the assets to the new owner is the net book value of the assets as recorded on the State's books at the time of the transfer. For purposes of this section, monetary consideration includes cash, new debt, and assumed debt.

(k) Transactions involving a provider’s capital stock—(1) Acquisition of capital stock of a provider. If the capital stock of a provider is acquired, the provider’s assets may not be revalued. For example, if Corporation A purchases the capital stock of Corporation B, the provider, Corporation B continues to be the provider after the purchase and Corporation A is merely the stockholder. Corporation B’s assets may not be revalued.

(2) Statutory merger. A statutory merger is a combination of two or more corporations under the corporation laws of the State, with one of the corporations surviving. The surviving corporation acquires the assets and liabilities of the merged corporation(s) by operation of State law. The effect of a statutory merger upon Medicare reimbursement is as follows:

(i) Statutory merger between unrelated parties. If the statutory merger is between two or more corporations that are unrelated (as specified in §413.17), the assets of the merged corporation(s) acquired by the surviving corporation may be revalued in accordance with paragraph (g) of this section. If the merged corporation was a provider before the merger, then it is subject to the provisions of paragraphs (d)(3) and (f) of this section concerning recovery of accelerated depreciation and the revaluation of gains and losses. The basis of the assets owned by the surviving corporation are unaffected by the transaction. An example of this type of transaction is one in which Corporation A purchase the capital stock of Corporation B, the provider. Immediately after the acquisition of the capital stock of Corporation B, there is a statutory merger of Corporation B and Corporation A, with Corporation A being the surviving corporation. Under these circumstances, at the time of the merger the transaction is one between related parties and is not a basis for revaluation of the provider’s assets.

(3) Consolidation. A consolidation is the combination of two or more corporations resulting in the creation of a new corporate entity. If at least one of the original corporations is a provider, the effect of a consolidation upon Medicare reimbursement for the provider is as follows:

(i) Consolidation between unrelated parties. If the consolidation is between two or more corporations that are unrelated (as specified in §413.17), the assets of the provider corporation(s) may be revalued in accordance with paragraph (g) of this section.

(ii) Consolidation between related parties. If the consolidation is between two or more related corporations (as specified in §413.17), no revaluation of provider assets is permitted.

§413.139 Depreciation: Optional allowance for depreciation based on a percentage of operating costs.

(a) Principle. With respect to all assets acquired before 1966, the provider, at its option, may choose an allowance for depreciation based on a percentage of operating costs. The operating costs
to be used are the provider's 1965 operating costs or the provider's current year's allowable costs, whichever are the lower. The percentage to be applied is 5 percent starting with the year 1966-67, with such percentage being uniformly reduced by one-half percent each succeeding year. The allowance based on operating costs is in addition to regular depreciation on assets acquired after 1965; however, if the optional allowance is selected, the combined amount of such allowance on pre-1966 assets and the straight-line depreciation on assets acquired after 1965 (including the estimated depreciation on assets held on a rental basis during the current year) may not exceed 6 percent of the provider's allowable cost for the current year.

(b) Definitions—(1) Operating costs. Operating costs are the total costs incurred by the provider in operating the institution or facility.

(2) Allowable costs. Allowable costs are the costs of a provider that are includable under the principles for cost reimbursement. Through application of apportionment methods to the total amount of such allowable costs, the share of a provider's total cost that is attributable to covered services for beneficiaries is determined.

(c) Application. If a provider has inadequate historical cost records for pre-1966 depreciable assets, the provider may elect to receive an allowance for depreciation on such assets based on a percentage of operating costs. The optional allowance for depreciation for such assets may be used, however, whether or not a provider has records of the cost of pre-1966 depreciable assets currently in use.

(d) Allowance based on a percentage of operating costs. The allowance for depreciation based on a percentage of operating costs is to be computed by applying a specified percentage to a base amount equal to the provider's 1965 total operating costs, without adjustments to these principles or the current year's allowable operating costs, whichever is lower. The percentage to be applied is five for the reporting period that begins during 1967-68, and continues to decline annually by equal amounts to become zero in 1976-77.

(2) If used as a base for determining the optional allowance for depreciation, neither the 1965 operating costs nor the current year's allowable costs are to include any actual depreciation, estimated depreciation on rented depreciable-type assets, allowance in lieu of specific recognition of other costs, or return on equity capital. Such exclusions are to be made only for the purpose of computing the allowance for depreciation based on operating costs. For other purposes, the excluded amounts are recognized in determining allowable costs and for computing the costs of services furnished to Medicare beneficiaries during the reporting period.

(e) Change to actual depreciation. (1) A provider that elects this allowance may at any time before 1976 change to actual depreciation on all pre-1966 depreciable assets. In such case, this option is eliminated and the provider can no longer elect to receive an allowance for depreciation based on a percentage of operating costs.

(2) If the provider desires to change to actual depreciation but either has no historical cost records or has incomplete records, the determination of historical cost may be made through appropriate means involving expert consultation with the determination being subject to review and approval by the intermediary.

(f) Determination of optional allowance based on percentage of operating costs illustrated. The following illustrates how the provider would determine the optional allowance for depreciation based on operating costs.

Example No. 1. The provider keeps its records on a calendar year basis. The current year's actual allowable cost and the actual operating cost for 1965 do not include any actual depreciation or rentals on depreciable-type assets. The current year's allowable cost also does not include any allowance in lieu of specific recognition of other costs or return on equity capital.

| Year 1966 |
|-------------------|-------------------|
| Current year's allowable cost | $1,100,000 |
| Operating cost for 1965 | $1,000,000 |
| Percent for determining the allowance | 5 |
§ 413.139

Calculation of allowance for depreciation based on a percentage of operating costs and the estimated straight-line depreciation on depreciable-type assets rented after 1965.

(g) Limitation on depreciation if optional allowance is used. This optional allowance only is subject to a limitation based on the provider's total allowable operating cost for the current year. To determine this limitation, compute the sum of the actual depreciation claimed, the allowance based on a percentage of operating costs, and the estimated straight-line depreciation on depreciable-type assets rented after 1965. If this sum exceeds six percent of the provider's current year's allowable cost (exclusive of any actual depreciation claimed, estimated depreciation on rented depreciable-type assets, allowance in lieu of specific recognition of other costs, and return on equity capital), the allowance for depreciation based on a percentage of operating costs is reduced by the amount of excess. In applying this limitation, if the actual depreciation claimed is on an accelerated basis, it must be converted to a straight-line basis only for use in calculating this limitation. It is presumed that pre-1966 assets will not be retired at a greater than normal rate, and the limitation of six percent, as it affects the availability of the allowance, is designed as a safeguard if the presumption is not borne out. If the provider does not elect to use the optional allowance, the combined allowance for depreciation based on costs of pre-1966 assets and those subsequently acquired is not subject to the six percent limitation.

Example No. 1. The following illustration demonstrates how this limitation would be determined.

YEARS 1966—Continued

<table>
<thead>
<tr>
<th>Allowance</th>
<th>$50,000</th>
</tr>
</thead>
</table>

1 1965 Operating cost was used in computing the allowance for depreciation based on a percentage of operating costs because it was lower than 1966 allowable cost.

YEAR 1967

<table>
<thead>
<tr>
<th>Current year's allowable cost</th>
<th>$1,200,000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operating cost for 1965</td>
<td>$1,000,000</td>
</tr>
<tr>
<td>Percent for determining the allowance</td>
<td>5</td>
</tr>
<tr>
<td>Allowance</td>
<td>$50,000</td>
</tr>
</tbody>
</table>

1 1965 Operating cost was used in computing the allowance for depreciation based on a percentage of operating costs because it was lower than 1967 allowable cost.

YEAR 1968

<table>
<thead>
<tr>
<th>Current year's allowable cost</th>
<th>$1,000,000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operating cost for 1965</td>
<td>$900,000</td>
</tr>
<tr>
<td>Percent for determining the allowance</td>
<td>4 1/2</td>
</tr>
<tr>
<td>Allowance</td>
<td>$40,500</td>
</tr>
</tbody>
</table>

1 The current year's allowable cost was used in computing the allowance for depreciation based on percentage of operating costs because it was lower than 1965 allowable cost.
2 Since the reporting period began during the year 1966-1967 (July 1, 1966-June 30, 1967), 5 percent is the percentage to be used.

Example No. 2. When the provider pays rent for depreciable-type assets rented prior to 1966, the estimated depreciation on such assets must be deducted from the allowance.

The following illustration demonstrates how the allowance is determined.

The provider keeps its records on a calendar year basis. The current year's actual allowable cost and the actual operating cost for 1965 did not include any actual depreciation, allowance in lieu of specific recognition of other costs, or return on equity capital. However, such costs have been adjusted to exclude estimated depreciation on rented depreciable-type assets.

YEAR 1966

<table>
<thead>
<tr>
<th>Adjusted current year's allowable cost</th>
<th>$1,100,000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adjusted operating cost for 1965</td>
<td>$1,000,000</td>
</tr>
<tr>
<td>Percent for determining the allowance</td>
<td>5</td>
</tr>
<tr>
<td>Allowance</td>
<td>$50,000</td>
</tr>
</tbody>
</table>

Less estimated depreciation for depreciable-type assets rented prior to 1966 on which rental is paid in 1966

| Amount                              | $3,000    |

Adjusted allowance

| Amount                              | $47,000   |

1 1965 operating cost was used in computing the allowance for depreciation based on a percentage of operating costs because it was lower than 1966 allowable cost.

(g) Limitation on depreciation if optional allowance is used. This optional allowance only is subject to a limitation based on the provider's total allowable operating cost for the current year. To determine this limitation, compute the sum of the actual depreciation claimed, the allowance based on a percentage of operating costs, and the estimated straight-line depreciation on depreciable-type assets rented after 1965. If this sum exceeds six percent of the provider's current year's allowable cost (exclusive of any actual depreciation claimed, estimated depreciation on rented depreciable-type assets, allowance in lieu of specific recognition of other costs, and return on equity capital), the allowance for depreciation based on a percentage of operating costs is reduced by the amount of excess. In applying this limitation, if the actual depreciation claimed is on an accelerated basis, it must be converted to a straight-line basis only for use in calculating this limitation. It is presumed that pre-1966 assets will not be retired at a greater than normal rate, and the limitation of six percent, as it affects the availability of the allowance, is designed as a safeguard if the presumption is not borne out. If the provider does not elect to use the optional allowance, the combined allowance for depreciation based on costs of pre-1966 assets and those subsequently acquired is not subject to the six percent limitation.

Example No. 1. The following illustration demonstrates how this limitation would be determined.

YEAR 1966

[The provider keeps its records on a calendar year basis. The current year's actual allowable cost and the actual operating cost for 1965 have been adjusted to exclude actual depreciation, the estimated depreciation on rented depreciable-type assets, allowance in lieu of specific recognition of other costs, and return on equity capital.]

| Adjusted operating cost for 1965     | $1,000,000 |
| Percent for determining the allowance | 5          |

In 1966 assets were acquired which produce a straight-line depreciation of

| Estimated depreciation on assets rented in 1966 | $18,000    |

Adjustable allowable operating cost for 1966

| Amount                              | $1,100,000 |

CALCULATION OF ALLOWANCE FOR DEPRECIATION BASED ON A PERCENTAGE OF OPERATING COSTS

Gross allowance

| 5 percent times adjusted 1965 operating costs ($1,000,000) | $50,000    |

Estimated depreciation on assets rented in 1966

| Amount                              | 2,000      |

Straight-line depreciation on post-1965 assets

| Amount                              | 18,000     |

Total

| Amount                              | 70,000     |

6 percent of adjusted 1966 allowable operating cost

| Amount                              | 66,000     |

Reduction in allowance

| Amount                              | 4,000      |

Allowance

| Amount                              | 50,000     |

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§ 413.144 Depreciation: Allowance for depreciation on fully depreciated or partially depreciated assets.

(a) Principle. Depreciation on assets being used by a provider at the time it enters into the Medicare program is allowed. This principle applies even though such assets may be fully or partially depreciated on the provider's books.

(b) Application. Depreciation is allowable on assets being used at the time the provider enters into the program. This applies even though such assets may be fully depreciated on the provider's books or fully depreciated with respect to other third-party payers. So long as an asset is being used, its useful life is considered not to have ended, and consequently the asset is subject to depreciation based upon a revised estimate of the asset's useful life as determined by the provider and approved by the intermediary. Correction of prior years' depreciation to reflect revision of estimated useful life should be made in the first year of participation in the program unless the provider has used the optional method (§ 413.139), in which case the correction should be made at the time of discontinuing the use of that method. If an asset has become fully depreciated under Medicare, further depreciation is not appropriate or allowable, even though the asset may continue in use.

(c) Example of an allowance for a fully-depreciated asset. For example, if a 50-year-old building is in use at the time the provider enters into the program, depreciation is allowable on the building even though it has been fully depreciated on the provider's books. Assuming that a reasonable estimate of the asset's continued life is 20 years (70 years from the date of acquisition), the provider may claim depreciation over the next 20 years—if the asset is in use that long—or a total depreciation of as much as twenty-seventieths of the asset's historical cost.

(d) Corrections to depreciation. If the asset is disposed of before the expiration of its estimated useful life, the depreciation would be adjusted to the actual useful life. Likewise, a provider may not have fully depreciated other assets it is using and finds that it has incorrectly estimated the useful lives of those assets. In such cases, the provider may use the corrected useful lives in determining the amount of depreciation, provided such corrections have been approved by the intermediary.

§ 413.149 Depreciation: Allowance for depreciation on assets financed with Federal or public funds.

(a) Principle. Depreciation is allowed on assets financed with Hill-Burton or other Federal or public funds.

(b) Application. Like other assets (including other donated depreciable assets), assets financed with Hill-Burton or other Federal or public funds become a part of the provider institution's plant and equipment to be used in furnishing services. It is the function of payment of depreciation to provide funds that make it possible to maintain the assets and preserve the capital employed in the production of services. Therefore, irrespective of the source of financing of an asset, if it is used in the providing of services for beneficiaries of the program, payment for depreciation of the asset is, in fact, a cost of the production of those services. Moreover, recognition of this cost is necessary to maintain productive capacity for the future. An incentive for funding of depreciation is provided in

[The provider keeps its records on a calendar year basis. The current year's actual allowable cost and the actual operating cost for 1965 have been adjusted to exclude actual depreciation, the estimated depreciation on rented depreciable-type assets, allowance in lieu of specific recognition of other costs, and return on equity capital.)

<table>
<thead>
<tr>
<th>Description</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reduction</td>
<td>4,000</td>
</tr>
<tr>
<td>Adjusted allowance</td>
<td>46,000</td>
</tr>
<tr>
<td>Total depreciation allowance for 1966</td>
<td>64,000</td>
</tr>
</tbody>
</table>

Assume in this illustration that the provider had elected to use the declining balance method in computing its allowable depreciation and the rental expense for depreciable-type assets was $3,500. In that case, it would include in its 1966 allowable cost not only the $46,000 allowance based on operating costs but also $36,000 (in this instance 2×straight-line rate is used) in actual depreciation and the rental expense of $3,500—or a total of $85,500 covering all its depreciable assets.
§ 413.153 Interest expense.

(a)(1) Principle. Necessary and proper interest on both current and capital indebtedness is an allowable cost. However, interest costs are not allowable if incurred as a result of—

(i) Judicial review by a Federal court (as described in § 413.64(j));

(ii) An interest assessment on a determined overpayment (as described in § 405.377 of this chapter); or

(iii) Interest on funds borrowed to repay an overpayment (as described in § 413.64(j) or § 405.378 of this chapter), up to the amount of the overpayment, unless the provider had made a prior commitment to borrow funds for other purposes (for example, capital improvements).

(2) Exception. In those cases of administrative or judicial reversal, interest paid on funds borrowed to repay an overpayment is an allowable cost, in accordance with this section.

(b) Definitions—(1) Interest. Interest is the cost incurred for the use of borrowed funds. Interest on current indebtedness is the cost incurred for funds borrowed for a relatively short term. This is usually for such purposes as working capital for normal operating expenses. Interest on capital indebtedness is the cost incurred for funds borrowed for capital purposes, such as acquisition of facilities and equipment, and capital improvements. Generally, loans for capital purposes are long-term loans.

(2) Necessary. Necessary interest is interest that meets the following requirements:

(i) It is incurred on a loan made to satisfy a financial need of the provider. Loans that result in excess funds or investments are not considered necessary.

(ii) It is incurred on a loan made for a purpose reasonably related to patient care.

(iii) It is reduced by investment income except income from—

(A) Gifts, grants, and endowments, whether held separately or pooled with other funds;

(B) Funded depreciation that meets the program's qualifying criteria;

(C) The provider's qualified pension funds;

(D) The provider's deferred compensation funds that meet the program's qualifying criteria; and

(E) The provider's self-insurance trust funds that meet the program's qualifying criteria.

(iv) It is not reduced by interest received as a result of judicial review by a Federal court (as described in § 413.64(j)).

(3) Proper. Proper requires that interest be—

(i) Incurred at a rate not in excess of what a prudent borrower would have had to pay in the money market existing at the time the loan was made; and

(ii) Paid to a lender not related through control or ownership, or personal relationship to the borrowing organization. However, interest is allowable if paid on loans from the provider's donor-restricted funds, the funded depreciation account, the provider's qualified pension fund.

(4) Zero coupon bonds. Zero coupon bonds are issued by government agencies, corporations, and banks at a price substantially below the face value. The difference between the purchase price and the face value reflects the actual amount of interest and is neither a discount nor an adjustment to the interest rate as with other bonds. Interest is paid at maturity when the bond is redeemed at face value.

(c) Borrower-lender relationship. (1) Except as described in paragraph (c)(2) of this section, to be allowable, interest expense must be incurred on indebtedness established with lenders or lending organizations not related through control, ownership, or personal relationship to the borrower. Presence of any of these factors could affect the "bargaining" process that usually accompanies the making of a loan, and could thus be suggestive of an agreement on higher rates of interest or of unnecessary loans. Loans should be made under terms and conditions that a prudent borrower would make in arms-length transactions with lending organizations.
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institutions. The intent of this provision is to assure that loans are legitimate and needed, and that the interest rate is reasonable. Thus, interest paid by the provider to partners, stockholders, or related organizations of the provider would not be allowable. If the owner uses his own funds in a business, it is reasonable to treat the funds as invested funds or capital, rather than borrowed funds. Therefore, if interest on loans by partners, stockholders, or related organizations is disallowed as a cost solely because of the relationship factor, the principal of such loans is treated as invested funds in the computation of the provider’s equity capital under §413.157.

(2) Exceptions to the general rule regarding interest on loans from controlled sources of funds are made in the following circumstances. Interest on loans to providers by partners, stockholders, or related organizations made prior to July 1, 1966, is allowable as cost, provided that the terms and conditions of payment of such loans have been maintained in effect without modification subsequent to July 1, 1966. If the general fund of a provider “borrows” from a donor-restricted fund and pays interest to the restricted fund, this interest expense is an allowable cost. The same treatment is accorded interest paid by the general fund on money “borrowed” from the funded depreciation account of the provider or from the provider’s qualified pension fund. In addition, if a provider operated by members of a religious order borrows from the order, interest paid to the order is an allowable cost.

(3) When a provider borrows funds, but only some of the funds are necessary, repayments of the loan (principal and interest portions) are applied first to pay for the necessary portion of the loan. Only after all of the necessary portion of the loan (principal and interest) has been repaid are any repayments applied to the unnecessary portion of the loan. Repayments toward non-allowable borrowing pertaining to assets or activities not related to patient care are considered investments, and the provisions of paragraph (b)(2)(iii) of this section are applied.

(e) Zero coupon bonds—(1) Interest on bonds issued on or after August 15, 1996. For zero coupon bonds issued on or after August 15, 1996, interest income earned for investment purposes is an allowable expense, and interest income earned for investment purposes is an allowable offset, in the cost reporting period in which the interest accrues.
(2) Interest income offset. Interest income from zero coupon bonds must be offset against allowable interest expense as prescribed in paragraph (b)(2) of this section and in §413.130(g)(2). If zero coupon bonds are purchased with the proceeds of an advanced refunding of debt, offset of the investment income is required under §413.153(b)(2)(iii), but the investment income is not prorated under §413.130(g)(2).

(3) Use of effective interest method. (i) Interest expense and interest income from zero coupon bonds that are reported as they accrue must be amortized using the effective interest method. This method recognizes the actual accrual of interest expense or income for each interest computation period (as specified by the bond instrument) throughout the life of the bond.

(ii) A constant effective yield rate is determined and applied to the book value (outstanding loan balance including prior accrued interest) of the bond at the beginning of each period to determine the total interest for the period.

(iii) If the interest computation period involves portions of more than one cost reporting period, the amount of interest for that computation period shall be apportioned to each cost reporting period.

(iv) An example of the computation of interest using the effective interest method follows:

Facts
Life of zero coupon bond: 15 years.
Value at maturity: $50,000.
Bondholder pays $6,996 for the bond.
Annual interest rate is 13.5506% compounded semi-annually.

From the table below, interest for the first year would be $980.11 ($474.00 plus $506.11).

<table>
<thead>
<tr>
<th>Col 1 Six-month periods</th>
<th>Col 2 Book value beginning of period</th>
<th>Col. 3 Effective interest*</th>
<th>Col 4 Book value end of period (colums 2 + 3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$6,996.00</td>
<td>$474.00</td>
<td>$7,470.00</td>
</tr>
<tr>
<td>2</td>
<td>7,470.00</td>
<td>506.11</td>
<td>7,976.11</td>
</tr>
<tr>
<td>3</td>
<td>7,976.11</td>
<td>540.40</td>
<td>8,516.51</td>
</tr>
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<td>4</td>
<td>8,516.51</td>
<td>577.02</td>
<td>9,093.53</td>
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<tr>
<td>29</td>
<td>43,855.94</td>
<td>2,971.37</td>
<td>46,827.31</td>
</tr>
</tbody>
</table>

*Computed by multiplying the book value at the beginning of each period (Column 2) by 6.7753% (the annual interest rate of 13.5506% / 2 = 6.7753%).

§413.157 Return on equity capital of proprietary providers.

(a) Definitions. For purposes of this section—
Proprietary provider means a provider that is organized and operated with the expectation of earning a profit for its owners (as distinguished from a provider that is organized and operated on a nonprofit basis). Proprietary providers may be sole proprietorships, partnerships, or corporations. Effective for cost reporting periods beginning on or after July 6, 1987, the term applies only to proprietary hospitals and SNFs.

(b) General rule. A reasonable return on equity capital invested and used in the provision of patient care is paid as an allowance in addition to the reasonable cost of covered services furnished to beneficiaries by proprietary providers.

(1) Rate of return applicable to proprietary providers for cost reporting periods beginning before July 6, 1987. Except as provided in paragraphs (b)(2), (b)(3), and (b)(4) of this section, the amount allowable on an annual basis, for cost reporting periods beginning before July 6, 1987, is determined by multiplying the provider’s equity capital by a percentage equal to one and one-half times the average of the rates of interest on special issues of public debt obligations issued for purchase by the Medicare Part A Trust Fund for each of the months during the provider’s reporting period or portion thereof covered under the program.

(2) Rate of return for inpatient hospital services furnished by proprietary hospitals. The rate used in determining the return for inpatient hospital services is a percentage of the average of the rates of interest described in paragraph (b)(1)
of this section. The percentages applicable to inpatient hospital services are as follows:

(i) 150 percent for cost reporting periods beginning before April 20, 1983.

(ii) 100 percent for cost reporting periods beginning on or after April 20, 1983 and before October 1, 1986.

(iii) 75 percent for cost reporting periods beginning on or after October 1, 1986 and before October 1, 1987.

(iv) 50 percent for cost reporting periods beginning on or after October 1, 1987 and before October 1, 1988.

(v) 25 percent for cost reporting periods beginning on or after October 1, 1988 and before October 1, 1989.

(vi) Zero percent for cost reporting periods beginning on or after October 1, 1989.

(3) Rate of return related to proprietary SNFs. (i) For cost reporting periods beginning on or after October 1, 1985, the rate used in determining the return for SNF services furnished before October 1, 1993, is a percentage equal to the average of the rates of interest described in paragraph (b)(1) of this section.

(ii) There is no allowance for return for SNF services furnished on or after October 1, 1993.

(4) Rate of return related to outpatient hospital services. (i) For cost reporting periods beginning on or after October 1, 1985, the rate used in determining the return for outpatient hospital services furnished before January 1, 1988, is a percentage equal to the average of the rates of interest described in paragraph (b)(1) of this section.

(ii) There is no allowance for return for outpatient hospital services furnished on or after January 1, 1988.

(5) Rate of return for proprietary services of all nonhospital and non-SNF providers. (i) For cost reporting periods beginning on or after October 1, 1985, but before July 6, 1987, the rate used in determining the return for services of all nonhospital and non-SNF providers is a percentage equal to the average of the rates of interest described in paragraph (b)(1) of this section.

(ii) For cost reporting periods beginning on or after July 6, 1987, there is no allowance for return on equity capital for nonhospital and non-SNF providers.

(c) Application—(1) Computation of equity capital. For purposes of computing the allowable return, the provider’s equity capital means—

(i) The provider’s investment in plant, property, and equipment related to patient care (net of depreciation) and funds deposited by a provider who leases plant, property, or equipment related to patient care and is required by the terms of the lease to deposit such funds (net of noncurrent debt related to such investment or deposited funds); and

(ii) Net working capital maintained for necessary and proper operation of patient care activities. However, debt representing loans from partners, stockholders, or related organizations on which interest payments would be allowable as costs but for the provisions of §413.153(b)(3)(ii), is not subtracted in computing the amount of equity capital in order that the proceeds from such loans be treated as part of the provider’s equity capital. In computing the amount of equity capital upon which a return is allowable, investment in facilities is recognized on the basis of the historical cost, or other basis, used for depreciation and other purposes under Part A of Medicare.

(2) Acquisitions after July 1970. With respect to a facility or any tangible assets of a facility acquired on or after August 1, 1970, the excess of the price paid for such facility or such tangible assets over the historical cost, as defined in §413.134(b), or the cost basis, as determined under §413.134(g) (whichever is appropriate), is not includable in equity capital, and loans made to finance such excess portion of the cost of such acquisitions (see §413.153(d)) are excluded in computing equity capital.

(3) Acquisitions prior to August 1970. With respect to a facility or any tangible assets of a facility acquired before August 1970, the excess of the price paid for such facility or assets over the fair market value of tangible assets at the time of purchase is includable in equity capital to the extent that it is reasonable except that the cumulative allowable return for such excess may not exceed 100 percent of such excess. For purposes of this section, the cumulative allowable return means the sum of the allowable rate of return on equity capital for all months starting...
from August 1, 1970. For example, if the allowable rates of return on equity capital for a provider are 9 percent for the first year (and such year started August 1, 1970), 8.5 percent for the second year, and 10.5 percent for the third year, the cumulative allowable return at the end of the third year would be 28 percent. After the cumulative allowable return equals 100 percent, the inclusion in equity capital of the excess is no longer allowable.

(4) Computation of return on equity capital. For purposes of computing the allowable return, the amount of equity capital is the average investment during the reporting period. The rate of return allowed, as derived from time to time based upon interest rates in accordance with this principle, is determined by HCFA and communicated through intermediaries. Return on investment as an element of allowable costs is subject to apportionment in the same manner as other elements of allowable costs.

Example of calculation of cumulative allowable return. X purchased a provider on July 1, 1969, paying $100,000 in excess of the fair market value of the assets acquired. Provider X files its cost report on a calendar-year basis. The allowable rate of return on equity capital for August 1, 1970-December 31, 1970 (4.538 percent), is obtained by multiplying the allowable rate of return for the period ending December 31, 1970 (10.891) by 4/12 (a fraction of which the numerator is the number of months from August 1, 1970, to the end of the cost-reporting period and the denominator is the number of months in the cost-reporting period). The cumulative allowable return for Provider X for the period August 1, 1970-December 31, 1973, (32.367 percent) is computed as follows:

<table>
<thead>
<tr>
<th>Cost reporting year ending</th>
<th>Rate of return on equity capital (percent)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dec. 31, 1970</td>
<td>4.538</td>
</tr>
<tr>
<td>Dec. 31, 1971</td>
<td>8.969</td>
</tr>
<tr>
<td>Dec. 31, 1972</td>
<td>8.891</td>
</tr>
<tr>
<td>Dec. 31, 1973</td>
<td>9.969</td>
</tr>
<tr>
<td>Total</td>
<td>32.367</td>
</tr>
</tbody>
</table>

(The $100,000 paid in excess of the fair market value of the assets acquired is included in equity capital until the sum of the allowable rate of return on equity capital equals 100 percent. Of course, no portion of the $100,000 may be amortized as an allowable cost or is otherwise allowable for any program reimbursement purposes other than for determining the provider’s equity capital.

Subpart H—Payment for End-Stage Renal Disease (ESRD) Services and Organ Procurement Costs

§ 413.170 Scope.
This subpart implements sections 1881(b)(2) and (b)(7) of the Act by—
(a) Setting forth the principles and authorities under which HCFA is authorized to establish a prospective payment system for outpatient maintenance dialysis furnished in or under the supervision of an ESRD facility approved under subpart U of part 405 of this chapter (referred to as “facility” in this section). For purposes of this section and §413.172 through §413.198, “outpatient maintenance dialysis” means outpatient dialysis, home dialysis, self-dialysis, and home dialysis training, as defined in §405.2102(f)(2)(ii), (f)(2)(iii), and (f)(3) of this chapter, and includes all items and services specified in §§410.50 and 410.52 of this chapter.
(b) Providing procedures and criteria under which a facility may receive an exception to the prospective payment rates; and
(c) Establishing procedures that a facility must follow to appeal its payment amount under the prospective payment system.

§ 413.172 Principles of prospective payment.
(a) Payments for outpatient maintenance dialysis are based on rates set prospectively by HCFA.
(b) All approved ESRD facilities must accept the prospective payment rates established by HCFA as payment in full for covered outpatient maintenance dialysis.
(c) HCFA publishes the methodology used to establish payment rates and
§ 413.174 Prospective rates for hospital-based and independent ESRD facilities.

(a) Establishment of rates. HCFA establishes prospective payment rates for ESRD facilities using a methodology that—

(1) Differentiates between hospital-based facilities and independent ESRD facilities;

(2) Effectively encourages efficient delivery of dialysis services; and

(3) Provides incentives for increasing the use of home dialysis.

(b) Determination of independent facility. For purposes of rate-setting and payment under this section, HCFA considers any facility that does not meet all of the criteria of a hospital-based facility to be an independent facility. A determination under this paragraph (b) is an initial determination under § 498.3 of this chapter.

(c) Determination of hospital-based facility. A determination under this paragraph (c) is an initial determination under § 498.3 of this chapter. For purposes of rate-setting and payment under this section, HCFA determines that a facility is hospital-based if the—

(1) Facility and hospital are subject to the bylaws and operating decisions of a common governing board. This governing board, which has final administrative responsibility, approves all personnel actions, appoints medical staff, and carries out similar management functions;

(2) Facility’s director or administrator is under the supervision of the hospital’s chief executive officer and reports through him or her to the governing board;

(3) Facility personnel policies and practices conform to those of the hospital;

(4) Administrative functions of the facility (for example, records, billing, laundry, housekeeping, and purchasing) are integrated with those of the hospital; and

(5) Facility and hospital are financially integrated, as evidenced by the cost report, which reflects allocation of overhead to the facility through the required step-down methodology.

(d) Nondetermination of hospital-based facility. In determining whether a facility is hospital-based, HCFA does not consider—

(1) An agreement between a facility and a hospital concerning patient referral;

(2) A shared service arrangement between a facility and a hospital; or

(3) The physical location of a facility on the premises of a hospital.

(e) Add-on amounts. If all the physicians furnishing services to patients in an ESRD facility elect the initial method of payment (as described in § 414.313(c) of this chapter), the prospective rate (as described in paragraph (a) of this section) paid to that facility is increased by an add-on amount as described in § 414.313.

(f) Erythropoietin/Epoietin (EPO). (1) When EPO is furnished to an ESRD patient by a Medicare-approved ESRD facility or a supplier of home dialysis equipment and supplies, payment is based on the amount specified in paragraph (f)(3) of this section.

(2) The payment is made only on an assignment basis, that is, directly to the facility or supplier, which must accept, as payment in full, the amount that HCFA determines.

(3) HCFA determines the payment amount in accordance with the following rules:

(i) The amount is prospectively determined, as specified in section 1881(b)(11)(B)(ii) of the Act, reviewed and adjusted by HCFA, as necessary, and paid to hospital-based and independent dialysis facilities and to suppliers of home dialysis equipment and supplies, regardless of the location of the facility, supplier, or patient.

(ii) If HCFA determines that an adjustment to the payment amount is necessary, HCFA publishes a Federal Register notice proposing a revision to the EPO payment amount and requesting public comment.

(iii) Any increase in this amount for a year does not exceed the percentage increase (if any) in the implicit price deflator for gross national product (as published by the Department of Commerce) for the second quarter of the preceding year over the implicit price deflator for the second quarter of the second preceding year.
The Medicare payment amount is subject to the Part B deductible and coinsurance.

Additional payment for certain drugs. In addition to the prospective payment described in this section, HCFA makes an additional payment for certain drugs furnished to ESRD patients by a Medicare-approved ESRD facility. HCFA makes this payment directly to the ESRD facility. The facility must accept the allowance determined by HCFA as payment in full. Payment for these drugs is made as follows:

1. Hospital-based facilities. HCFA makes payments in accordance with the cost reimbursement rules set forth in this part.
2. Independent facilities. HCFA makes payment in accordance with the methodology set forth in §405.517 of this chapter for paying for drugs that are not paid on a cost or prospective payment basis.

§413.176 Amount of payments.

(a) If the beneficiary has incurred the full deductible applicable under Part B of Medicare before the dialysis treatment, the intermediary pays the facility 80 percent of its prospective payment rate.

(b) If the beneficiary has not incurred the full deductible applicable under Part B of Medicare before the dialysis treatment, the intermediary subtracts the amount applicable to the deductible from the facility's prospective rate and pays the facility 80 percent of the remainder, if any.

§413.178 Bad debts.

(a) HCFA will reimburse each facility its allowable Medicare bad debts, as defined in §413.80(b), up to the facility's costs, as determined under Medicare principles, in a single lump sum payment at the end of the facility's cost reporting period.

(b) A facility must attempt to collect deductible and coinsurance amounts owed by beneficiaries before requesting reimbursement from HCFA for uncollectible amounts. Section 413.80 specifies the collection efforts facilities must make.

(c) A facility must request payment for uncollectible deductible and coinsurance amounts owed by beneficiaries by submitting an itemized list that specifically enumerates all uncollectable amounts related to covered services under the composite rate.

§413.180 Procedures for requesting exceptions to payment rates.

(a) Outpatient maintenance dialysis payments. All payments for outpatient maintenance dialysis furnished at or by facilities are made on the basis of prospective payment rates.

(b) Criteria for requesting an exception. If a facility projects on the basis of prior year costs and utilization trends that it will have an allowable cost per treatment higher than its prospective rate set under §413.174, and if these excess costs are attributable to one or more of the factors in §413.182, the facility may request, in accordance with paragraph (d) of this section, that HCFA approve an exception to that rate and set a higher prospective payment rate. However, a facility may only request an exception or seek to retain its previously approved exception rate when authorized under the conditions specified in paragraphs (d) and (e) of this section.

(c) Application of deductible and coinsurance. The higher payment rate is subject to the application of deductible and coinsurance in accordance with §413.176.

(d) Payment rate exception request. A facility must request an exception to its payment rate within 180 days of—

1. The effective date of its new composite payment rate(s);
2. The effective date that HCFA opens the exceptions process; or
3. The date on which an extraordinary cost-increasing event occurs, as specified (or provided for) in §§413.182(c) and 413.188.

(e) Criteria for retaining a previously approved exception rate. A facility may elect to retain its previously approved exception rate in lieu of any composite rate increase or any other exception amount if—

1. The conditions under which the exception was granted have not changed;
§ 413.182 Criteria for approval of exception requests.

HCFA may approve exceptions to an ESRD facility’s prospective payment rate if the facility demonstrates, by convincing objective evidence, that its total per treatment costs are reasonable and allowable under the relevant regulation.

(ii) Determination of an exception request. In determining the facility’s payment rate under the exception process, HCFA excludes all costs that are not reasonable or allowable under the reasonable cost principles set forth in this part.

(ij) Approval of an exception request. A facility that has been denied an exception request during the 180 days may file another exception request if all required documentation is filed with the intermediary by the 180th day.

§ 413.182 Criteria for approval of exception requests.

HCFA may approve exceptions to an ESRD facility’s prospective payment rate if the facility demonstrates, by convincing objective evidence, that its total per treatment costs are reasonable and allowable under the relevant regulation.

(ii) Determination of an exception request. In determining the facility’s payment rate under the exception process, HCFA excludes all costs that are not reasonable or allowable under the reasonable cost principles set forth in this part.

(j) Period of approval: Payment exception request. Except for exceptions approved under §§413.180(e), 413.180(k), 413.182(c), and 413.188, a prospective exception payment rate approved by HCFA applies for the period from the date the complete exception request was filed with its intermediary until the earlier of the—

(1) Date the circumstances justifying the exception rate no longer exist; or

(2) End of the period during which the announced rate was to apply.

(k) Period of approval: Payment exception request under §§413.182(c) and 413.188. A prospective exception payment rate approved by HCFA under §§413.182(c) and 413.188 applies from the date of the extraordinary event until the end of the period during which the prospective announced rate was to apply, unless HCFA determines that another date is more appropriate. If HCFA does not extend the exception period and the facility believes that it continues to require an exception to its rate, the facility must reapply in accordance with the procedures in this section.

(l) Denial of an exception request. HCFA denies exception requests submitted without the documentation specified in §413.182 and the applicable regulations cited therein.

(m) Criteria for refiling a denied exception request. A facility that has been denied an exception request during the 180 days may file another exception request if all required documentation is filed with the intermediary by the 180th day.
cost reimbursement principles of part 413 and that its per treatment costs in excess of its payment rate are directly attributable to any of the following criteria:
(a) Atypical service intensity (patient mix), as specified in §413.184.
(b) Isolated essential facility, as specified in §413.186.
(c) Extraordinary circumstances, as specified in §413.188.
(d) Self-dialysis training costs, as specified in §413.190.
(e) Frequency of dialysis, as specified in §413.192.

§ 413.184 Payment exception: Atypical service intensity (patient mix).

(a) To qualify for an exception to the prospective payment rate based on atypical service intensity (patient mix)—

(1) A facility must demonstrate that a substantial proportion of the facility’s outpatient maintenance dialysis treatments involve atypically intense dialysis services, special dialysis procedures, or supplies that are medically necessary to meet special medical needs of the facility’s patients. Examples that may qualify under this criterion are more intense dialysis services that are medically necessary for patients such as—

(i) Patients who have been referred from other facilities on a temporary basis for more intense care during a period of medical instability and who return to the original facility after stabilization;

(ii) Pediatric patients who require a significantly higher staff-to-patient ratio than typical adult patients; or

(iii) Patients with medical conditions that are not commonly treated by ESRD facilities and that complicate the dialysis procedure.

(2) The facility must demonstrate clearly that these services, procedures, or supplies and its per treatment costs are prudent and reasonable when compared to those of facilities with a similar patient mix.

(3) A facility must demonstrate that—

(i) Its nursing personnel costs have been allocated properly between each mode of care; and

(ii) The additional nursing hours per treatment are not the result of an excess number of employees.

(b) Documentation.

(1) A facility must submit a listing of all outpatient dialysis patients (including all home patients) treated during the most recently completed fiscal or calendar year showing—

(i) Patients who received transplants, including the date of transplant;

(ii) Patients awaiting a transplant who are medically able, have given consent, and are on an active transplant list, and projected transplants;

(iii) Home patients;

(iv) In-facility patients, staff-assisted, or self-dialysis;

(v) Individual patient diagnosis;

(vi) Diabetic patients;

(vii) Patients isolated because of contagious disease;

(viii) Age of patients;

(ix) Mortality rate, by age and diagnosis;

(x) Number of patient transfers, reasons for transfers, and any related information; and

(xi) Total number of hospital admissions for the facility’s patients, reason for, and length of stay of each session.

(2) The facility also must—

(i) Submit documentation on costs of nursing personnel (registered nurses, licensed practical nurses, technicians, and aides) incurred during the most recently completed fiscal year cost report showing—

(A) Amount each employee was paid;

(B) Number of personnel;

(C) Amount of time spent in the dialysis unit; and

(D) Staff-to-patient ratio based on total hours, with an analysis of productive and nonproductive hours.

(ii) Submit documentation on supply costs incurred during the most recently completed fiscal or calendar year cost report showing—

(A) By modality, a complete list of supplies used routinely in a dialysis treatment;

(B) The make and model number of each dialyzer and its component cost; and

(C) That supplies are prudently purchased (for example, that bulk discounts are used when available).
§ 413.186 Payment exception: Isolated essential facility.

(a) Qualifications. To qualify for an exception to the prospective payment rate based on being an isolated essential facility—

(1) The facility must be the only supplier of dialysis in its geographical area;

(2) The facility’s patients must be unable to obtain dialysis services elsewhere without substantial additional hardship; and

(3) The facility’s excess costs must be justifiable.

(b) Criteria for determining qualifications. In determining whether a facility qualifies for an exception based on its being an isolated essential facility, HCFA considers—

(1) Local, permanent residential population density;

(2) Typical local commuting distances from medical services;

(3) Volume of treatments; and

(4) The extent that other dialysis facilities are used by area residents (other than the applying facility’s patients).

(c) Documentation. (1) Isolated. Generally, to be considered isolated, the facility must document that it is located outside an established Metropolitan Statistical Area and provides dialysis to a permanent patient population, as opposed to a transient patient population.

(2) Essential. To be considered essential, the facility must document—

(i) That a substantial number of its patients cannot obtain dialysis services elsewhere without additional hardship; and

(ii) The additional hardship the patients will incur in travel time and cost.

(d) Cost per treatment. The facility must—

(i) Document that its cost per treatment is reasonable; and

(ii) Explain how the facility’s cost per treatment in excess of its composite rate relates to the isolated essential facility criteria specified in paragraph (b) of this section.

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§ 413.188 Payment exception: Extraordinary circumstances.

(a) To qualify for an exception to the prospective payment rate based on extraordinary circumstances, the facility must substantiate that it incurs excess costs beyond its control due to a fire,
§ 413.190 Payment exception: Self-dialysis training costs.

(a) Qualifications. To qualify for an exception to the prospective payment rate based on self-dialysis training costs, the facility must establish that it incurs per treatment costs for furnishing self-dialysis and home dialysis training that exceed the facility's payment rate for such training sessions.

(b) Justification. To justify its exception request, a facility must—

(1) Separately identify those elements contributing to its costs in excess of the composite training rate; and

(2) Demonstrate that its per treatment costs are reasonable and allowable.

(c) Criteria for determining proper cost reporting. HCFA considers the facility's total costs, cost finding and apportionment, including its allocation of costs, to determine if costs are properly reported by treatment modality.

(d) Limitation of exception requests. Exception requests for a higher training rate are limited to those cost components relating to training such as technical staff, medical supplies, and the special costs of education (manuals and education materials). These requests may include overhead and other indirect costs to the extent that these costs are directly attributable to the additional training costs.

(e) Documentation. The facility must provide the following information to support its exception request:

(1) A copy of the facility's training program.

(2) Computation of the facility's cost per treatment for maintenance sessions and training sessions including an explanation of the cost difference between the two modalities.

(3) Class size and patients' training schedules.

(4) Number of training sessions required, by treatment modality, to train patients.

(5) Number of patients trained for the current year and the prior 2 years on a monthly basis.

(6) Projection for the next 12 months of future training candidates.

(7) The number and qualifications of staff at training sessions.

(f) Accelerated training exception. (1) An ESRD facility may bill Medicare for a dialysis training session only when a patient receives a dialysis treatment (normally three times a week for hemodialysis). Continuous cycling peritoneal dialysis (CCPD) and continuous ambulatory peritoneal dialysis (CAPD) are daily treatment modalities; ESRD facilities are paid the equivalent of three hemodialysis treatments for each week that CCPD and CAPD treatments are provided.

(2) If an ESRD facility elects to train all its patients using a particular treatment modality more often than during each dialysis treatment and, as a result, the number of billable training dialysis sessions is less than the number of actual training sessions, the facility may request a composite rate exception, limited to the lesser of the—

(i) Facility's projected training cost per treatment; or

(ii) Cost per treatment the facility would have received in training a patient if it had trained patients only during a dialysis treatment, that is, three times per week.

(3) An ESRD facility may bill a maximum of 25 training sessions per patient for hemodialysis training and 15 sessions for CCPD and CAPD training.

(4) In computing the payment amount under an accelerated training exception, HCFA uses a minimum number of training sessions per patient (15 for hemodialysis and 5 for CAPD and CCPD) when the facility actually provides fewer than the minimum number of training sessions.

(5) To justify an accelerated training exception request, an ESRD facility must document that a significant number of training sessions for a particular modality are provided during a shorter but more condensed period.
§ 413.192 Payment exception: Frequency of dialysis.

(a) Qualification. To qualify for an exception to the prospective payment rate based on frequency of dialysis, the facility must establish that it has a substantial portion of outpatient maintenance dialysis treatments furnished to patients who dialyze less frequently than three times per week.

(b) Definition. For purposes of this section, “substantial” means the number of treatments furnished by the facility is at least 15 percent lower than the number would be if all patients dialyzed three times a week.

(c) Limitation for per treatment payment rates. Per treatment payment rates granted under this exception may not exceed the amount that produces weekly payments per patient equal to three times the facility’s prospective composite rate, exclusive of any exception amounts.

(d) Documentation. To document that an ESRD facility furnishes a substantial number of dialysis treatments at a frequency less than three times per week per patient, the facility must submit the following information:

(1) A list of patients receiving outpatient dialysis treatments for the cost report that is filed with the request. The list must indicate—

(i) Whether the patients are permanent, transient, or temporary;

(ii) The medically prescribed frequency of dialysis; and

(iii) The number of dialysis treatments that each patient received on a weekly and yearly basis and an explanation of any discrepancy between that calculation and the number of treatments reported on the facility’s cost report.

(2) A list of patients used to project treatments. The list must indicate—

(i) Whether the patients are permanent, transient, or temporary;

(ii) The medically prescribed frequency of dialysis;

(iii) The number of dialysis treatments that each patient is projected to receive on a weekly and yearly basis, an explanation of any discrepancy between that calculation and the number of treatments reported on the facility’s projected cost report, and an explanation for any change among prior, actual, and projected data.

(3) A schedule showing the number of treatments to be furnished twice a week and the number of treatments that would have been furnished if each patient were dialyzed three times a week.

(4) A computation of the facility’s projected costs per treatment using the—

(i) Projected number of treatments furnished twice a week; and

(ii) Number of treatments if patients dialyze three times a week.

(5) A schedule showing the computation of the percentage decrease in the number of treatments.

§ 413.194 Appeals.

(a) Appeals under section 1878 of the Act. (1) A facility that disputes the amount of its allowable Medicare bad debts reimbursed by HCFA under § 413.178 may request review by the intermediary or the Provider Reimbursement Review Board (PRRB) in accordance with subpart R of part 405 of this chapter.

(2) A facility must request and obtain a final agency decision prior to seeking judicial review of a dispute regarding the amount of allowable Medicare bad debts.

(b) Other appeals. (1) A facility that has requested higher payment per treatment in accordance with § 413.180 may request review from the intermediary or the PRRB if HCFA has denied the request in whole or in part. In such a case, the procedure in subpart R of part 405 of this chapter is followed to the extent that it is applicable.
§ 413.198 Recordkeeping and cost reporting requirements for outpatient maintenance dialysis.

(a) Purpose and Scope. This section implements section 1881(b)(2)(B)(i) of the Act by specifying recordkeeping and cost reporting requirements for ESRD facilities approved under subpart U of part 405 of this chapter. The records and reports will enable HCFA to determine the costs incurred in furnishing outpatient maintenance dialysis as defined in § 413.170(a).

(b) Recordkeeping and reporting requirements.

(1) Each facility must keep adequate records and submit the appropriate HCFA-approved cost report in accordance with §§ 413.20 and 413.24, which provide rules on financial data and reports, and adequate cost data and cost finding, respectively.

(2) The cost reimbursement principles set forth in this part (beginning with § 413.134, Depreciation, and excluding the principles listed in paragraph (b)(4) of this section), apply in the determination and reporting of the allowable cost incurred in furnishing outpatient maintenance dialysis treatments to patients dialyzing in the facility, or incurred by the facility in furnishing home dialysis service, supplies, and equipment.

(3) Allowable cost is the reasonable cost related to dialysis treatments. Reasonable cost includes all necessary and proper expenses incurred by the facility in furnishing the dialysis treatments, such as administrative costs, maintenance costs, and premium payments for employee health and pension plans. It includes both direct and indirect costs and normal standby costs. Reasonable cost does not include costs that—

(i) Are not related to patient care for outpatient maintenance dialysis;

(ii) Are for services or items specifically not reimbursable under the program;

(iii) Flow from the provision of luxury items or services (items or services substantially in excess of or more

§ 413.196 Notification of changes in rate-setting methodologies and payment rates.

(a) HCFA or the facility's intermediary notifies each facility of changes in its payment rate. This notice includes changes in individual facility payment rates resulting from corrections or revisions of particular geographic labor cost adjustment factors.

(b) Changes in payment rates resulting from incorporation of updated cost data or general revisions of geographic labor cost adjustment factors are announced by notice published in the Federal Register without opportunity for prior comment. Revisions of the rate-setting methodology are published in the Federal Register in accordance with the Department's established rulemaking procedures.
§413.200 Payment of independent organ procurement organizations and histocompatibility laboratories.

(a) Principle. Covered services furnished after September 30, 1978 by organ procurement organizations (OPOs) and histocompatibility laboratories in connection with kidney acquisition and transplantation will be reimbursed under the principles for determining reasonable cost contained in this part. Services furnished by freestanding OPOs and histocompatibility laboratories, that have an agreement with the Secretary in accordance with paragraph (c) of this section, will be reimbursed by making an interim payment to the transplant hospitals using these services and by making a retroactive adjustment, directly with the OPO or laboratory, based upon a cost report filed by the OPO or laboratory. (The reasonable costs of services furnished by hospital based OPOs or laboratories will be reimbursed in accordance with the principles contained in §§413.60 and 413.64.)

(b) Definitions. For purposes of this section:

Freestanding refers to an OPO or a histocompatibility laboratory that is not—

1. Subject to the control of the hospital with respect to the hiring, firing, training, and paying of employees; and
2. Considered as a department of the hospital for insurance purposes (including malpractice insurance, general liability insurance, worker’s compensation insurance, and employee retirement insurance).

Histocompatibility laboratory means a laboratory meeting the standards and providing the services for kidneys or other organs set forth in §413.2171(d) of this chapter.

OPO means an organization defined in §486.302 of this chapter.

(c) Agreements with independent OPOs and laboratories. (1) Any freestanding OPO or histocompatibility laboratory that wishes to have the cost of its pretransplant services reimbursed under the Medicare program must file an agreement with HCFA under which the OPO or laboratory agrees—

(i) To file a cost report in accordance with §413.24(f) within three months after the end of each fiscal year;

(ii) To permit HCFA to designate an intermediary to determine the interim reimbursement rate payable to the transplant hospitals for services provided by the OPO or laboratory and to make a determination of reasonable cost based upon the cost report filed by the OPO or laboratory;

(iii) To provide such budget or cost projection information as may be required to establish an initial interim reimbursement rate;

(iv) To pay to HCFA amounts that have been paid by HCFA to transplant hospitals that are determined to be in excess of the reasonable cost of the services provided by the OPO or laboratory; and

(v) Not to charge any individual for items or services for which that individual is entitled to have payment made under section 1861 of the Act.

(2) The initial cost report due from an OPO or laboratory is for its first fiscal year during any portion of which it had an agreement with the Secretary under paragraphs (c) (1) and (2) of this section. The initial cost report covers...
only the period covered by the agreement.

(d) Interim reimbursement. (1) Hospitals eligible to receive Medicare reimbursement for renal transplantation will be paid for the pretransplantation services of a freestanding OPO or histocompatibility laboratory that has an agreement with the Secretary under paragraph (c) of this section, on the basis of an interim rate established by an intermediary for that OPO or laboratory.

(2) The interim rate will be based on the average cost per service incurred by an OPO or laboratory, during its previous fiscal year, associated with procuring a kidney for transplantation. This interim rate may be adjusted if necessary for anticipated cost changes. If there is not adequate cost data to determine the initial interim rate, it will be determined according to the OPO’s or laboratory’s estimate of its projected costs for the fiscal year.

(3) Payments made on the basis of the interim rate will be reconciled directly with the OPO or laboratory after the close of its fiscal year, in accordance with paragraph (e) of this section.

(4) Information on the interim rate for all freestanding OPOs and histocompatibility laboratories shall be disseminated to all transplant hospitals and intermediaries.

(e) Retroactive adjustment. (1) Cost reports. Information provided in cost reports by freestanding OPOs and histocompatibility laboratories must meet the requirements for cost data and cost finding specified in paragraphs (a) through (e) of §413.24. These cost reports must provide a complete accounting of the cost incurred by the agency or laboratory in providing covered services, the total number of Medicare beneficiaries who received those services, and any other data necessary to enable the intermediary to make a determination of the reasonable cost of covered services provided to Medicare beneficiaries.

(2) Audit and adjustment. A cost report submitted by a freestanding OPO or histocompatibility laboratory will be reviewed by the intermediary and a new interim reimbursement rate for the succeeding fiscal year will be established based upon this review. A retroactive adjustment in the amount paid under the interim rate will be made in accordance with §413.64(f). If the determination of reasonable cost reveals an overpayment or underpayment resulting from the interim reimbursement rate paid to transplant hospitals, a lump sum adjustment will be made directly between that intermediary and the OPO or laboratory.

(f) For services furnished on or after April 1, 1988, no payment may be made for services furnished by an OPO that does not meet the requirements of part 405, subpart D of this chapter.

(g) Appeals. Any OPO or histocompatibility laboratory that disagrees with an intermediary’s cost determination under this section is entitled to an intermediary hearing, in accordance with the procedures contained in §§405.1811 through 405.1833, if the amount in controversy is $1,000 or more.

§ 413.202 Organ procurement organization (OPO) cost for kidneys sent to foreign countries or transplanted in patients other than Medicare beneficiaries.

An OPO’s total costs for all kidneys is reduced by the costs associated with procuring kidneys sent to foreign transplant centers or transplanted in patients other than Medicare beneficiaries. OPOs, as defined in §435.302 of this chapter, must separate costs for procuring kidneys that are sent to foreign transplant centers and kidneys transplanted in patients other than Medicare beneficiaries from Medicare allowable costs prior to final settlement by the Medicare fiscal intermediary. Medicare costs are based on the ratio of the number of usable kidneys transplanted into Medicare beneficiaries to the total number of usable kidneys applied to reasonable costs. Certain long-standing arrangements that existed before March 3, 1988 (for example, an OPO that procures kidneys at a military transplant hospital for transplant at that hospital), will be deemed to be Medicare kidneys for cost reporting statistical purposes. The OPO must submit a request to the fiscal intermediary for review and approval of these arrangements.
§ 413.203 Transplant center costs for organs sent to foreign countries or transplanted in patients other than Medicare beneficiaries.

(a) A transplant center’s total costs for all organs is reduced by the costs associated with procuring organs sent to foreign transplant centers or transplanted in patients other than Medicare beneficiaries. Organs are defined in §486.302 (only covered organs will be paid for on a reasonable cost basis).

(b) Transplant center hospitals must separate costs for procuring organs that are sent to foreign transplant centers and organs transplanted in patients other than Medicare beneficiaries from Medicare allowable costs prior to final cost settlement by the Medicare fiscal intermediaries.

(c) Medicare costs are based on the ratio of the number of usable organs transplanted into Medicare beneficiaries to the total number of usable organs applied to reasonable costs.

Subpart I—Prospectively Determined Payment Rates for Low-Volume Skilled Nursing Facilities, for Cost Reporting Periods Beginning Prior to July 1, 1998

SOURCE: 60 FR 37594, July 21, 1995, unless otherwise noted.

§ 413.300 Basis and scope.

(a) Basis. This subpart implements section 1888(d) of the Act, which provides for optional prospectively determined payment rates for qualified SNFs.

(b) Scope. This subpart sets forth the eligibility criteria an SNF must meet to qualify, the process governing election of prospectively determined payment rates, and the basis and methodology for determining prospectively determined payment rates.

§ 413.302 Definitions.

For purposes of this subpart—

Area wage level means the average wage per hour for all classifications of employees as reported by health care facilities within a specified area.

Census region means one of the 9 census divisions, comprising the 50 States and the District of Columbia, established by the Bureau of the Census for statistical and reporting purposes.

Routine capital-related costs means the capital-related costs, allowable for Medicare purposes (as described in Subpart G of this Part), that are allocated to the SNF participating inpatient routine service cost center as reported on the Medicare cost report.

Routine operating costs means the cost of regular room, dietary, and nursing services, and minor medical and surgical supplies for which a separate charge is not customarily made. It does not include the costs of ancillary services, capital-related costs, or, where appropriate, return on equity.

Rural area means any area outside an urban area in a census region.

Urban area means a Metropolitan Statistical Area (MSA) or New England County Metropolitan Area (NECMA), as defined by the Office of Management and Budget, or a New England county deemed to be an urban area, as listed in §412.62(f)(1)(ii)(B) of this chapter.

§ 413.304 Eligibility for prospectively determined payment rates.

(a) General rule. An SNF is eligible to receive a prospectively determined payment rate for a cost reporting period if it had fewer than 1,500 Medicare covered inpatient days as reported on a Medicare cost report in its immediately preceding cost reporting period. This criterion applies even if the SNF received a prospectively determined payment rate during the preceding cost reporting period.

(b) Less than a full cost reporting period. If the cost reporting period that precedes an SNF’s request for prospectively determined payment is not a full cost reporting period, the SNF is eligible to receive prospectively determined payment rates only if the average daily Medicare census for the period (Medicare inpatient days divided by the total number of days in the cost reporting period) is not greater than 4.1.

(c) Newly-participating SNFs. An SNF is eligible to receive prospectively determined payment rates for its first cost reporting period for which it is approved to participate in Medicare.
§ 413.308 Rules governing election of prospectively determined payment rates.

(a) Requirements. An SNF must notify its intermediary at least 30 calendar days before the beginning of the cost reporting period for which it requests to receive such payment that it elects prospectively determined payment rates. A separate request must be made for each cost reporting period for which an SNF seeks prospectively determined payment. A newly participating SNF with no preceding cost reporting period must make its election within 30 days of its notification of approval to participate in Medicare.

(b) Intermediary notice. After evaluating an SNF’s request for prospectively determined payment rates, the intermediary notifies the SNF in writing as to whether the SNF meets any of the eligibility criteria described in § 413.304 and the timely election requirements under § 413.308(a). The intermediary must notify the SNF of its initial and final determinations within 10 working days after it receives all the data necessary to make each determination. The intermediary’s determination is limited to one cost reporting period.

(c) Prohibition against revocation. An SNF may not revoke its request after it has received the initial determination of eligibility from the intermediary and the cost reporting period has begun.

(d) Revocation by intermediary. If an SNF is given tentative approval to receive a prospectively determined payment rate, and, after the start of the applicable cost reporting period, the intermediary determines that the SNF does not meet the eligibility criteria, the intermediary must revoke the prospectively determined payment option.

§ 413.310 Basis of payment.

(a) Method of payment. Under the prospectively determined payment rate system, a qualified SNF receives a per diem payment of a predetermined rate for inpatient services furnished to Medicare beneficiaries. Each SNF’s routine per diem payment rate is determined according to the methodology described in § 413.312 and is based on various components of SNF costs.

(b) Payment in full. The payment rate represents payment in full for routine services as described in § 413.314 (subject to applicable coinsurance as described in Subpart G of Part 409 of this title), and for routine capital costs. Payment is made in lieu of payment on a reasonable cost basis for routine services and for routine capital costs.

§ 413.312 Methodology for calculating rates.

(a) Data used. (1) To calculate the prospectively determined payment rates, HCFA uses:

(i) The SNF cost data that were used to develop the applicable routine service cost limits;

(ii) A wage index to adjust for area wage differences; and

(iii) The most recent projections of increases in the costs from the SNF market basket index.

(2) In the annual schedule of rates published in the FEDERAL REGISTER under the authority of § 413.320, HCFA announces the wage index and the annual percentage increases in the market basket used in the calculation of the rates.

(b) Calculation of per diem rate. (1) Routine operating component of rate—(i) Adjusting cost report data. The SNF market basket index is used to adjust the routine operating cost from the SNF cost report to reflect cost increases occurring between cost reporting periods represented in the data collected and the midpoint of the initial cost reporting period to which the payment rates apply.

(ii) Calculating a per diem cost. For each SNF, an adjusted routine operating per diem cost is computed by dividing the adjusted routine operating cost calculated under paragraph (b)(1)(i) of this section by the SNF’s total patient days.

(iii) Adjusting for wage levels. (A) The SNF’s adjusted per diem routine operating cost calculated under paragraph (b)(1)(i) of this section is then divided into labor-related and nonlabor-related portions.

(B) The labor-related portion is obtained by multiplying the SNF’s adjusted per diem routine operating cost by a percentage that represents the labor-related portion of cost from the
§ 413.314 Determining payment amounts: Routine per diem rate.

(a) General rule. An SNF that elects to be paid under the prospectively determined payment rate system, and qualifies for such payment, is paid a per diem rate for inpatient routine services. This rate is adjusted to reflect area wage differences and the cost reporting period beginning date (if necessary) and is subject to the limitation specified in paragraph (d) of this section.

(b) Per diem rate. The prospectively determined payment rate for each urban and rural area in each census region is comprised of the following:

(1) A routine operating component, which is divided into:
   (i) A labor-related portion adjusted by the appropriate wage index; and
   (ii) A nonlabor-related portion.

(2) A routine capital-related cost portion.

(3) For proprietary SNFs only, a portion that is based on the return on owner’s equity related to routine cost, applicable only for services furnished before October 1, 1993.

(c) Adjustment for cost reporting period.

(1) If a facility has a cost reporting period beginning after the beginning of the Federal fiscal year, the intermediary increases the labor-related and nonlabor-related portions of the prospective payment rate that would otherwise apply to the SNF by a adjustment factor. Each factor represents the projected increase in the market basket index for a specific 12-month period. The factors are used to account for inflation in costs for cost reporting periods beginning after October 1. Adjustment factors are published in the annual notice of prospectively determined payment rates described in § 413.320.

(2) If a facility uses a cost reporting period that is not 12 months in duration, the intermediary must obtain a special adjustment factor from HCFA for the specific period.

(d) Limitation of prospectively determined payment rate. The per diem prospectively determined payment rate for an SNF, excluding capital-related costs and excluding return on equity for services furnished prior to October 1, 1993, may not exceed the individual
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§ 413.330 Basis and scope.
(a) Basis. This subpart implements section 1888(e) of the Act, which provides for the implementation of a prospective payment system for SNFs for cost reporting periods beginning on or after July 1, 1998.
(b) Scope. This subpart sets forth the framework for the prospective payment system for SNFs, including the methodology used for the development of payment rates and associated adjustments, the application of a transition phase, and related rules.

§ 413.333 Definitions.
As used in this subpart—
Case-mix index means a scale that measures the relative difference in resource intensity among different groups in the resident classification system.
Market basket index means an index that reflects changes over time in the prices of an appropriate mix of goods and services included in covered skilled nursing services.
Resident classification system means a system for classifying SNF residents into mutually exclusive groups based on clinical, functional, and resource-based criteria. For purposes of this subpart, this term refers to the current version of the Resource Utilization Groups, as set forth in the annual publication of Federal prospective payment rates described in §413.345.
Rural area means any area outside of an urban area.
Urban area means a metropolitan statistical area (MSA) or New England County Metropolitan Area (NECMA), as defined by the Office of Management and Budget, or a New England county deemed to be an urban area, as listed in §412.62(f)(1)(ii)(B) of this chapter.

§ 413.335 Basis of payment.
(a) Method of payment. Under the prospective payment system, SNFs receive a per diem payment of a predetermined rate for inpatient services furnished to Medicare beneficiaries. The per diem payments are made on the basis of the Federal payment rate described in §413.337 and, during a transition period,
§ 413.337 Methodology for calculating the prospective payment rates.

(a) Data used. (1) To calculate the prospective payment rates, HCFA uses—

(i) Medicare data on allowable costs from freestanding and hospital-based SNFs for cost reporting periods beginning in fiscal year 1995. SNFs that received “new provider” exemptions under § 413.30(e)(2) are excluded from the data base used to compute the Federal payment rates. In addition, allowable costs related to exceptions payments under § 413.30(f) are excluded from the data base used to compute the Federal payment rates;

(ii) An appropriate wage index to adjust for area wage differences;

(iii) The most recent projections of increases in the costs from the SNF market basket index;

(iv) Resident assessment and other data that account for the relative resource utilization of different resident types; and

(v) Medicare Part B SNF claims data reflecting amounts payable under Part B for covered SNF services (other than those services described in § 411.15(p)(2) of this chapter) furnished during SNF cost reporting periods beginning in fiscal year 1995 to individuals who were residents of SNFs and receiving Part A covered services.

(b) Methodology for calculating the per diem Federal payment rates—(1) Determining SNF costs. In calculating the initial unadjusted Federal rates applicable for services provided during the period beginning July 1, 1998 through September 30, 1999, HCFA determines each SNF’s costs by summing its allowable costs for the cost reporting period beginning in fiscal year 1995 and its estimate of Part B payments (described in paragraphs (a)(1)(i) and (a)(1)(v) of this section).

(2) Use of market basket index. The SNF market basket index is used to adjust the SNF cost data to reflect cost increases occurring between cost reporting periods represented in the data and the initial period (beginning July 1, 1998 and ending September 30, 1999) to which the payment rates apply. For each year, the cost data are updated by a factor equivalent to the annual market basket index percentage minus 1 percentage point.

(3) Calculation of the per diem cost. For each SNF, the per diem cost is computed by dividing the cost data for each SNF by the corresponding number of Medicare days.

(4) Standardization of data for variation in area wage levels and case-mix. The cost data described in paragraph (b)(2) of this section are standardized to remove the effects of geographic variation in wage levels and facility variation in case-mix. The cost data are standardized for geographic variation in wage levels using the wage index. The cost data are standardized for facility variation in case-mix using the case-mix indices and other data that indicate facility case-mix.

(5) Calculation of unadjusted Federal payment rates. HCFA calculates the national per diem unadjusted payment rates by urban and rural classification in the following manner:

(i) By computing the average per diem standardized cost of freestanding SNFs weighted by Medicare days.

(ii) By computing the average per diem standardized cost of freestanding and hospital-based SNFs combined weighted by Medicare days.

(iii) By computing the average of the amounts determined under paragraphs (b)(5)(i) and (b)(5)(ii) of this section.

(c) Calculation of adjusted Federal payment rates for case-mix and area wage levels. The Federal rate is adjusted to account for facility case-mix using a resident classification system and associated case-mix indices that account for the relative resource utilization of
different patient types. This classification system utilizes the resident assessment instrument completed by SNFs as described at §483.20 of this chapter, according to the assessment schedule described in §413.343(b). The Federal rate is also adjusted to account for geographic differences in area wage levels using an appropriate wage index.

(d) Annual updates of Federal unadjusted payment rates. HCFA updates the unadjusted Federal payment rates on a fiscal year basis.

(1) For fiscal years 2000 through 2002, the unadjusted Federal rate is equal to the rate for the previous period or fiscal year increased by a factor equal to the SNF market basket index percentage minus one percentage point.

(2) For subsequent fiscal years, the unadjusted Federal rate is equal to the rate for the previous fiscal year increased by the applicable SNF market basket index amount.

§ 413.340 Transition period.

(a) Duration of transition period and proportions for the blended transition rate. Beginning with an SNF’s first cost reporting period beginning on or after July 1, 1998, there is a transition period covering three cost reporting periods. During this transition phase, SNFs receive a payment rate comprising a blend of the adjusted Federal rate and a facility-specific rate. For the first cost reporting period beginning on or after July 1, 1998, payment is based on 75 percent of the facility-specific rate and 25 percent of the Federal rate. For the subsequent cost reporting period, the rate is comprised of 50 percent of the facility-specific rate and 50 percent of the Federal rate. In the final cost reporting period of the transition, the rate is comprised of 25 percent of the facility-specific rate and 75 percent of the Federal rate. For all subsequent cost reporting periods, payment is based entirely on the Federal rate.

(b) Calculation of facility-specific rate for the first cost reporting period. The facility-specific rate is computed based on the SNF’s Medicare allowable costs from its fiscal year 1995 cost report plus an estimate of the amounts payable under Part B for covered SNF services (other than those services described in §411.15(p)(2) of this chapter) furnished during fiscal year 1995 to individuals who were residents of SNFs and receiving Part A covered services. Allowable costs associated with exceptions, as described in §413.30(f), are included in the calculation of the facility-specific rate. Allowable costs associated with exemptions, as described in §413.30(e)(2), are included in the calculation of the facility-specific rate but only to the extent that they do not exceed 150 percent of the routine cost limit. Low Medicare volume SNFs that were paid a prospectively determined rate under §413.300 for their cost reporting period beginning in fiscal year 1995 will utilize that rate as the basis for the allowable costs of routine (operating and capital-related) expenses in determining the facility-specific rate. Each SNF’s allowable costs are updated to the first cost reporting period to which the payment rates apply using annual factors equal to the SNF market basket percentage minus one percentage point.

(c) SNFs participating in the Multistate Nursing Home Case-Mix and Quality Demonstration. SNFs that participated in the Multistate Nursing Home Case-Mix and Quality Demonstration in a cost reporting period that began in calendar year 1997 will utilize their allowable costs from that cost reporting period, including prospective payment amounts determined under the demonstration payment methodology.

(d) Update of facility-specific rates for subsequent cost reporting periods. The facility-specific rate for a cost reporting period that is subsequent to the first cost reporting period is equal to the facility-specific rate for the first cost reporting period (described in paragraph (a) of this section) updated by the market basket index.

(1) For a subsequent cost reporting period beginning in fiscal years 1998 and 1999, the facility-specific rate is equal to the facility-specific rate for the previous cost reporting period updated by the applicable market basket index percentage minus one percentage point.

(2) For a subsequent cost reporting period beginning in fiscal year 2000, the facility-specific rate is equal to the facility-specific rate for the previous cost reporting period updated by the
§ 413.343 Resident assessment data.

(a) Submission of resident assessment data. SNFs are required to submit the resident assessment data described at §483.20 of this chapter in the manner necessary to administer the payment rate methodology described in §413.337. This provision includes the frequency, scope, and number of assessments required.

(b) Assessment schedule. In accordance with the methodology described in §413.337(c) related to the adjustment of the Federal rates for case-mix, SNFs must submit assessments according to an assessment schedule. This schedule must include performance of patient assessments on the 5th, 14th, 30th, 60th, and 90th days of posthospital SNF care and such other assessments that are necessary to account for changes in patient care needs.

(c) Noncompliance with assessment schedule. HCFA pays a default rate for the Federal rate when a SNF fails to comply with the assessment schedule in paragraph (b) of this section. The default rate is paid for the days of a patient’s care for which the SNF is not in compliance with the assessment schedule.

§ 413.345 Publication of Federal prospective payment rates.

HCFA publishes information pertaining to each update of the Federal payment rates in the Federal Register. This information includes the standardized Federal rates, the resident classification system that provides the basis for case-mix adjustment (including the designation of those specific Resource Utilization Groups under the resident classification system that represent the required SNF level of care, as provided in §409.30 of this chapter), and the wage index. This information is published before May 1 for the fiscal year 1998 and before August 1 for the fiscal years 1999 and after.

§ 413.348 Limitation on review.

Judicial or administrative review under sections 1869 or 1878 of the Act or otherwise is prohibited with regard to the establishment of the Federal rates. This prohibition includes the methodology used in the computation of the Federal standardized payment rates, the case-mix methodology, and the development and application of the wage index. This prohibition on judicial and administrative review also extends to the methodology used to establish the facility-specific rates but not to determinations related to reasonable cost in the fiscal year 1995 cost reporting period used as the basis for these rates.

§ 413.350 Periodic interim payments for skilled nursing facilities receiving payment under the skilled nursing facility prospective payment system for Part A services.

(a) General rule. Subject to the exceptions in paragraphs (b) and (c) of this section, SNFs receiving payment under the PPS for Part A services do not receive interim payments during the cost reporting year, and receive payment only following submission of a bill. Paragraph (d) of this section provides for accelerated payments in certain circumstances.

(b) Periodic interim payments. (1) An SNF receiving payment under the prospective payment system may receive periodic interim payments (PIP) for Part A SNF services under the PIP method subject to the provisions of §413.64(h). To be approved for PIP, the SNF must meet the qualifying requirements in §413.64(h)(3). Moreover, as provided in §413.64(h)(5), intermediary approval is conditioned upon the intermediary’s best judgment as to whether payment can be made under the PIP method without undue risk of its resulting in an overpayment to the provider.

(2) Frequency of payment. The intermediary estimates an SNF’s prospective payments net of estimated beneficiary coinsurance and makes biweekly payments equal to 1/26 of the total estimated amount of payment for
the year. If an SNF has payment experience under the prospective payment system, the intermediary estimates PIP based on that payment experience, adjusted for projected changes supported by substantiated information for the current year. Each payment is made 2 weeks after the end of a biweekly period of service as described in §413.64(h)(6). The interim payments are reviewed at least twice during the reporting period and adjusted if necessary. Fewer reviews may be necessary if an SNF receives interim payments for less than a full reporting period. These payments are subject to final settlement.

(3) Termination of PIP—(i) Request by the SNF. An SNF receiving PIP may convert to receiving prospective payments on a non-PIP basis at any time.

(ii) Removal by the intermediary. An intermediary terminates PIP if the SNF no longer meets the requirements of §413.64(h).

(c) Interim payments for Medicare bad debts and for Part A costs not paid under the prospective payment system. For Medicare bad debts and for costs of an approved education program and other costs paid outside the prospective payment system, the intermediary determines the interim payments by estimating the reimbursable amount for the year based on the previous year’s experience, adjusted for projected changes supported by substantiated information for the current year, and makes biweekly payments equal to 1/26 of the total estimated amount. Each payment is made 2 weeks after the end of a biweekly period of service as described in §413.64(h)(6). The interim payments are reviewed at least twice during the reporting period and adjusted if necessary. Fewer reviews may be necessary if an SNF receives interim payments for less than a full reporting period. These payments are subject to final cost settlement.

(d) Accelerated payments—(1) General rule. Upon request, an accelerated payment may be made to an SNF that is receiving payment under the prospective payment system and is not receiving PIP under paragraph (b) of this section if the SNF is experiencing financial difficulties because of the following:

(i) There is a delay by the intermediary in making payment to the SNF.

(ii) Due to an exceptional situation, there is a temporary delay in the SNF’s preparation and submittal of bills to the intermediary beyond its normal billing cycle.

(2) Approval of payment. An SNF’s request for an accelerated payment must be approved by the intermediary and HCFA.

(3) Amount of payment. The amount of the accelerated payment is computed as a percentage of the net payment for unbilled or unpaid covered services.

(4) Recovery of payment. Recovery of the accelerated payment is made by recoupment as SNF bills are processed or by direct payment by the SNF.

[64 FR 41682, July 30, 1999]
§ 414.48 Limits on actual charges of non-participating suppliers.
§ 414.50 Physician billing for purchased diagnostic tests.
§ 414.52 Payment for physician assistants' services.
§ 414.54 Payment for certified nurse-midwives' services.
§ 414.56 Payment for nurse practitioners' and clinical nurse specialists' services.
§ 414.58 Payment of charges for physician services to patients in providers.
§ 414.60 Payment for the services of CRNAs.
§ 414.62 Fee schedule for clinical psychologist services.
§ 414.65 Payment for consultations via interactive telecommunications systems.

Subpart C [Reserved]

Subpart D—Payment for Durable Medical Equipment and Prosthetic and Orthotic Devices
§ 414.200 Purpose.
§ 414.202 Definitions.
§ 414.210 General payment rules.
§ 414.220 Inexpensive or routinely purchased items.
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§ 414.230 Determining a period of continuous use.
§ 414.232 Special payment rules for transcutaneous electrical nerve stimulators (TENS).

Subpart E—Determination of Reasonable Charges Under the ESRD Program
§ 414.300 Scope of subpart.
§ 414.310 Determination of reasonable charges for physician services furnished to renal dialysis patients.
§ 414.313 Initial method of payment.
§ 414.314 Monthly capitation payment method.
§ 414.316 Payment for physician services to patients in training for self-dialysis and home dialysis.
§ 414.320 Determination of reasonable charges for physician renal transplantation services.
§ 414.330 Payment for home dialysis equipment, supplies, and support services.
§ 414.335 Payment for EPO furnished to a home dialysis patient for use in the home.

Subparts F–H—[Reserved]
(2) Supplies and services covered “incident to” physician services (excluding drugs as specified in §414.36).

(3) Outpatient physical and occupational therapy services if furnished by a person or an entity that is not a Medicare provider of services as defined in §400.202 of this chapter.

(4) Diagnostic x-ray tests and other diagnostic tests (excluding diagnostic laboratory tests paid under the fee schedule established under section 1833(h) of the Act).

(5) X-ray, radium, and radioactive isotope therapy, including materials and services of technicians.

(6) Antigens, as described in section 1861(s)(2)(G) of the Act.

(7) Bone mass measurement.

§414.21 Medicare payment basis.

Medicare payment is based on the lesser of the actual charge or the applicable fee schedule amount.

§414.22 Relative value units (RVUs).

HCFA establishes RVUs for physicians' work, practice expense, and malpractice insurance.

(a) Physician work RVUs—(1) General rule. Physician work RVUs are established using a relative value scale in which the value of physician work for a particular service is rated relative to the value of work for other physician services.

(ii) Radiology services. HCFA bases the RVUs for all radiology services on the relative value scale developed under section 1834(b)(1)(A) of the Act, with appropriate modifications to ensure that the RVUs established for radiology services that are similar or related to other physician services are consistent with the RVUs established for those similar or related services.

(b) Practice expense RVUs. (1) Practice expense RVUs are computed for each service or class of service by applying average historical practice cost percentages to the estimated average allowed charge during the 1991 base period.

(2) The average practice expense percentage for a service or class of services is computed as follows:

(i) Multiply the average practice expense percentage for each specialty by the proportion of a particular service or class of service performed by that specialty.

(ii) Add the products for all specialties.

(3) For services furnished beginning calendar year (CY) 1994, for which 1994 practice expense RVUs exceed 1994 work RVUs and that are performed in
(4) For services furnished beginning January 1, 1998, practice expense RVUs for certain services are reduced to 110 percent of the work RVUs for those services. The following two categories of services are excluded from this limitation:

(i) The service is provided more than 75 percent of the time in an office setting; or

(ii) The service is one described in section 1848(c)(2)(G)(v) of the Act, codified at 42 U.S.C. 1395w-4(c)(2)(G). Section 1848(c)(2)(G)(v) of the Act refers to the 1998 proposed resource-based practice expense RVUs (as specified in the June 18, 1997 physician fee schedule proposed rule (62 FR 33158)) for the specific site, either in-office or out-of-office, increased from its 1997 practice expense RVUs.

(5) For services furnished beginning January 1, 1999, the practice expense RVUs are based on 75 percent of the practice expense RVUs applicable to services furnished in 1998 and 25 percent of the relative practice expense resources involved in furnishing the service. For services furnished in 2000, the practice expense RVUs are based on 50 percent of the practice expense RVUs applicable to services furnished in 1998 and 50 percent of the relative practice expense resources involved in furnishing the service. For services furnished in 2001, the practice expense RVUs are based on 25 percent of the practice expense RVUs applicable to services furnished in 1998 and 75 percent of the relative practice expense resources involved in furnishing the service. For services furnished in 2002 and subsequent years, the practice expense RVUs are based entirely on relative practice expense resources.

(i) Usually one of two levels of practice expense RVUs can be applied to each code. The lower facility practice expense RVUs apply to services furnished to patients in the hospital, skilled nursing facility, or ambulatory surgical center when the physician performs procedures on the ASC approved procedures list. The higher non-facility practice expense RVUs apply to services performed in a physician’s office or in an ASC. If the physician is performing a procedure not on the ASC approved procedures list, services furnished to patients in a nursing facility, in a facility or institution other than a hospital, skilled nursing facility, or in the home. The facility practice expense RVUs for a particular code may not be greater than the non-facility RVUs for that code.

(ii) Only one practice expense RVU per code can be applied for each of the following services: services that have only technical component practice expense RVUs or only professional component practice expense RVUs; evaluation and management services, such as hospital or nursing facility visits, that are furnished exclusively in one setting; and major surgical services.

(6)(i) HCFA establishes criteria for supplemental surveys regarding specialty practice expenses submitted to HCFA by August 1, 2000 that may be used in determining practice expense RVUs for the 2001 physician fee schedule.

(ii) Any HCFA-designated specialty group may submit a supplemental survey.

(iii) Survey data and related materials submitted to HCFA between August 2, 2000 and August 1, 2001 will be considered for use in determining practice expense RVUs for the 2002 physician fee schedule.

(c) Malpractice insurance RVUs. (1) Malpractice insurance RVUs are computed for each service or class of services by applying average malpractice insurance historical practice cost percentages to the estimated average allowed charge during the 1991 base period.

(2) The average historical malpractice insurance percentage for a service or class of services is computed as follows:

(i) Multiply the average malpractice insurance percentage for each specialty by the proportion of a particular service or class of services performed by that specialty.
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§ 414.26 Determining the GAF.

HCFA establishes a GAF for each service in each fee schedule area.

(a) Geographic indices. HCFA uses the following indices to establish the GAF:

(1) An index that reflects one-fourth of the difference between the relative value of physicians’ work effort in each of the different fee schedule areas as determined under §414.22(a) and the national average of that work effort.

(2) An index that reflects the relative costs of the mix of goods and services comprising practice expenses (other than malpractice expenses) in each of the different fee schedule areas as determined under §414.22(b) compared to the national average of those costs.

(3) An index that reflects the relative costs of malpractice expenses in each of the different fee schedule areas as determined under §414.22(c) compared to the national average of those costs.

(b) Class-specific practice cost indices. If the application of a single index to different classes of services would be substantially inequitable because of differences in the mix of goods and services comprising practice expenses for the different classes of services, more than one index may be established under paragraph (a)(2) of this section.

(c) Computation of GAF. The GAF for each fee schedule area is the sum of the physicians’ work adjustment factor, the practice expense adjustment factor, and the malpractice cost adjustment factor, as defined in this section:

(1) The geographic physicians’ work adjustment factor for a service is the product of the proportion of the total relative value for the service that reflects the RVUs for the work component and the geographic physicians’ work index value established under paragraph (a)(1) of this section.

(2) The geographic practice expense adjustment factor for a service is the product of the proportion of the total relative value for the service that reflects the RVUs for the practice expense component, multiplied by the geographic practice cost index (GPCI).
§ 414.28 Conversion factors.

HCFA establishes CFs in accordance with section 1848(d) of the Act.

(a) Base-year CFs. HCFA established the CF for 1992 so that had section 1848 of the Act applied during 1991, it would have resulted in the same aggregate amount of payments for physician services as the estimated aggregate amount of these payments in 1991, adjusted by the update for 1992 computed as specified in § 414.30.

(b) Subsequent CFs. For calendar years 1993 through 1995, the CF for each year is equal to the CF for the previous year, adjusted in accordance with § 414.30. Beginning January 1, 1996, the CF for each calendar year may be further adjusted so that adjustments to the fee schedule in accordance with section 1848(c)(2)(B)(ii) of the Act do not cause total expenditures under the fee schedule to differ by more than $20 million from the amount that would have been spent if these adjustments had not been made.

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(b) Subsequent CFs. For calendar years 1993 through 1995, the CF for each year is equal to the CF for the previous year, adjusted in accordance with § 414.30. Beginning January 1, 1996, the CF for each calendar year may be further adjusted so that adjustments to the fee schedule in accordance with section 1848(c)(2)(B)(ii) of the Act do not cause total expenditures under the fee schedule to differ by more than $20 million from the amount that would have been spent if these adjustments had not been made.

§ 414.30 Conversion factor update.

Unless Congress acts in accordance with section 1848(d)(3) of the Act—

(a) General rule. The CF update for a CY equals the Medicare Economic Index increased or decreased by the number of percentage points by which the percentage increase in expenditures for physician services (or for a particular category of physician services, such as surgical services) in the second preceding FY over the third preceding FY exceeds the performance standard rate of increase established for the second preceding FY.

(b) Downward adjustment. The downward adjustment may not exceed the following:

(1) For CY's 1992 and 1993, 2 percentage points.
(2) For CY 1994, 2.5 percentage points.
(3) For CY's 1995 and thereafter, 5 percentage points.

§ 414.32 Determining payments for certain physicians' services furnished in facility settings.

(a) Definition. As used in this section, facility settings include the following facilities:

(1) Hospital outpatient departments, including clinics and emergency rooms.
(2) Hospital inpatient departments.
(3) Comprehensive outpatient rehabilitation facilities.
(4) Comprehensive inpatient rehabilitation facilities.
(5) Inpatient psychiatric facilities.
(6) Skilled nursing facilities.

(b) General rule. If physicians' services of the type routinely furnished in physicians' offices are furnished in facility settings before January 1, 1999, the physician fee schedule amount for those services is determined by reducing the practice expense RVUs for the services by 50 percent. For services furnished on or after January 1, 1999, the practice expense RVUs are determined in accordance with § 414.30.

(c) Services covered by the reduction. HCFA establishes a list of services routinely furnished in physicians' offices nationally. Services furnished at least 50 percent of the time in physicians' offices are subject to this reduction.

(d) Services excluded from the reduction. The reduction established under this section does not apply to the following:

(1) Rural health clinic services.
(2) Surgical services not on the ambulatory surgical center covered list of procedures published under § 416.65(c) of this chapter when furnished in an ambulatory surgical center.
§ 414.38 Special rules for payment of low osmolar contrast media.

(a) General. Payment for low osmolar contrast media is included in the technical component payment for diagnostic procedures except as specified in paragraph (b) of this section.

(b) Conditions for separate payment. For diagnostic procedures furnished to beneficiaries who are neither inpatients nor outpatients of any hospital, separate payment is made for low osmolar contrast media used in all intrathecal injections and in intravenous, and intra-arterial injections, if it is used for patients with one or more of the following characteristics:

(1) A history of a previous adverse reaction to contrast material, with the exception of a sensation of heat, flushing, or a single episode of nausea or vomiting.

(2) A history of asthma or allergy.

(3) Significant cardiac dysfunction including recent or imminent cardiac decompensation, severe arrhythmias, unstable angina pectoris, recent myocardial infarction, and pulmonary hypertension.

(4) Generalized severe debilitation.

(5) Sickle cell disease.

(c) Method of payment. If one of the conditions of paragraph (b) of this section is met, payment is made for low osmolar contrast media as set forth in §414.36 as a drug furnished incident to a physician's service, subject to paragraph (d) of this section.

(d) Drug payment reduction. If separate payment is made for low osmolar contrast media, the payment amount calculated in accordance with §414.36 is reduced by 8 percent to account for the allowance for contrast media already paid as if the physician had personally furnished the service.
§414.39 Special rules for payment of care plan oversight.

(a) General. Except as specified in paragraph (b) of this section, payment for care plan oversight is included in the payment for visits and other services under the physician fee schedule.

(b) Exception. Separate payment is made under the following conditions for physician care plan oversight services furnished to beneficiaries who receive HHA and hospice services that are covered by Medicare:

- (1) The care plan oversight services require recurrent physician supervision of therapy involving 30 or more minutes of the physician’s time per month.
- (2) Payment is made to only one physician per patient for services furnished during a calendar month period. The physician must have furnished a service requiring a face-to-face encounter with the patient at least once during the 6-month period before the month for which care plan oversight payment is first billed. The physician may not have a significant ownership interest in, or financial or contractual relationship with, the HHA in accordance with §424.22(d) of this chapter. The physician may not be the medical director or employee of the hospice and may not furnish services under an arrangement with the hospice.
- (3) If a physician furnishes care plan oversight services during a post-operative period, payment for care plan oversight services is made if the services are documented in the patient’s medical record as unrelated to the surgery.

[59 FR 63463, Dec. 8, 1994; 60 FR 49, Jan. 3, 1995; 60 FR 36733, July 18, 1995]

§414.40 Coding and ancillary policies.

(a) General rule. HCFA establishes uniform national definitions of services, codes to represent services, and payment modifiers to the codes.

(b) Specific types of policies. HCFA establishes uniform national ancillary policies necessary to implement the fee schedule for physician services. These include, but are not limited to, the following policies:

- (1) Global surgery policy (for example, post- and pre-operative periods and services, and intra-operative services).
- (2) Professional and technical components (for example, payment for services, such as an EEG, which typically comprise a technical component (the taking of the test) and a professional component (the interpretation)).
- (3) Payment modifiers (for example, assistant-at-surgery, multiple surgery, bilateral surgery, split surgical global services, team surgery, and unusual services).

§414.42 Adjustment for first 4 years of practice.

(a) General rule. For services furnished during CYs 1992 and 1993, except as specified in paragraph (b) of this section, the fee schedule payment amount or prevailing charge must be phased in as specified in paragraph (d) of this section for physicians, physical therapists (PTs), occupational therapists (OTs), and all other health care practitioners who are in their first through fourth years of practice.

(b) Exception. The reduction required in paragraph (d) of this section does not apply to primary care services or to services furnished in a rural area as defined in section 1886(d)(2)(D) of the Act that is designated under section 332(a)(1)(A) of the Public Health Service Act as a Health Professional Shortage Area.

(c) Definition of years of practice. (1) The “first year of practice” is the first full CY during the first 6 months of which the physician, PT, OT, or other health care practitioner furnishes professional services for which payment may be made under Medicare Part B, plus any portion of the prior CY if that prior year does not meet the first 6 months test.

- (2) The “second, third, and fourth years of practice” are the first, second, and third CYs following the first year of practice, respectively.

(d) Amounts of adjustment. The fee schedule payment for the service of a new physician, PT, OT, or other health care practitioner is limited to the following percentages for each of the indicated years:
§ 414.44 Transition rules.

(a) Adjusted historical payment basis—

(1) All services other than radiology and nuclear medicine services. For all physician services other than radiology services, furnished in a fee schedule area, the adjusted historical payment basis (AHPB) is the estimated weighted average prevailing charge applied in the fee schedule area for the service in CY 1991, as determined by HCFA without regard to physician specialty and as adjusted to reflect payments for services below the prevailing charge, adjusted by the update established for CY 1992.

(2) Radiology services. For radiology services, the AHPB is the amount paid for the service in the fee schedule area in CY 1991 under the fee schedule established under section 1834(b), adjusted by the update established for CY 1992.

(3) Nuclear medicine services. For nuclear medicine services, the AHPB is the amount paid for the service in the fee schedule area in CY 1991 under the fee schedule established under section 6105(b) of Public Law 101-239 and section 4102(g) of Public Law 101-508, adjusted by the update established for CY 1992.

(4) Transition adjustment. HCFA adjusts the AHPB for all services by 5.5 percent to produce budget-neutral payments for 1992.

(b) Adjustment of 1992 payments for physician services other than radiology services. For physician services furnished during CY 1992 the following rules apply:

(1) If the AHPB determined under paragraph (a) of this section is from 85 percent to 115 percent of the fee schedule amount for the area for services furnished in 1992, payment is at the fee schedule amount.

(2) If the AHPB determined under paragraph (a) of this section is less than 85 percent of the fee schedule amount for the area for services furnished in 1992, an amount equal to the AHPB minus 15 percent of the fee schedule amount is substituted for the fee schedule amount.

(3) If the AHPB determined under paragraph (a) of this section is greater than 115 percent of the fee schedule amount for the area for services furnished in 1992, an amount equal to the AHPB minus 15 percent of the fee schedule amount is substituted for the fee schedule amount.

(c) Adjustment of 1992 payments for radiology services. For radiology services furnished during CY 1992 the following rules apply:

(1) If the AHPB determined under paragraph (a) of this section is from 85 percent to 109 percent of the fee schedule amount for the area for services furnished in 1992, payment is at the fee schedule amount.

(2) If the AHPB determined under paragraph (a) of this section is less than 85 percent of the fee schedule amount for the area for services furnished in 1992, an amount equal to the AHPB plus 15 percent of the fee schedule amount is substituted for the fee schedule amount.

(3) If the AHPB determined under paragraph (a) of this section is greater than 109 percent of the fee schedule amount for the area for services furnished in 1992, an amount equal to the AHPB minus 9 percent of the fee schedule amount is substituted for the fee schedule amount.

(d) Computation of payments for CY 1993. For physician services subject to the transition rules in CY 1992 and furnished during CY 1993, the fee schedule is equal to 75 percent of the amount that would have been paid in the fee schedule area under the 1992 transition rules, adjusted by the amount of the 1993 update, plus 25 percent of the 1993 fee schedule amount.

(e) Computation of payments for CY 1994. For physician services subject to the transition rules in CY 1993, and furnished during CY 1994, the fee schedule is equal to 67 percent of the amount that would have been paid in the fee schedule area under the 1992 transition rules, adjusted by the amount of the 1994 update, plus 33 percent of the 1994 fee schedule amount.

(f) Computation of payments for CY 1995. For physician services subject to
§ 414.46 Additional rules for payment of anesthesia services.

(a) Definitions. For purposes of this section, the following definitions apply:

(1) Base unit means the value for each anesthesia code that reflects all activities other than anesthesia time. These activities include usual preoperative and postoperative visits, the administration of fluids and blood incident to anesthesia care, and monitoring services.

(2) Anesthesia practitioner, for the purpose of anesthesia time, means a physician who performs the anesthesia service alone, a CRNA who is not medically directed who performs the anesthesia service alone, or a medically directed CRNA.

(3) Anesthesia time means the time during which an anesthesia practitioner is present with the patient. It starts when the anesthesia practitioner begins to prepare the patient for anesthesia services and ends when the anesthesia practitioner is no longer furnishing anesthesia services to the beneficiary, that is, when the beneficiary may be placed safely under postoperative care. Anesthesia time is a continuous time period from the start of anesthesia to the end of an anesthesia service. In counting anesthesia time, the anesthesia practitioner can add blocks of anesthesia time around an interruption in anesthesia time as long as the anesthesia practitioner is furnishing continuous anesthesia care within the time periods around the interruption.

(b) Determinations of payment amount—Basic rule. For anesthesia services performed, medically directed, or medically supervised by a physician, HCFA pays the lesser of the actual charge or the anesthesia fee schedule amount.

(1) The carrier bases the fee schedule amount for an anesthesia service on the product of the sum of allowable base and time units and an anesthesia-specific CF. The carrier calculates the time units from the anesthesia time reported by the anesthesia practitioner for the anesthesia procedure. The physician who fulfills the conditions for medical direction in § 415.110 (Conditions for payment: Anesthesiology services) reports the same anesthesia time as the medically-directed CRNA.

(2) HCFA furnishes the carrier with the base units for each anesthesia procedure code. The base units are derived from the 1988 American Society of Anesthesiologists' Relative Value Guide except that the number of base units recognized for anesthesia services furnished during cataract or iridectomy surgery is four units.

(3) Modifier units are not allowed. Modifier units include additional units charged by a physician or a CRNA for patient health status, risk, age, or unusual circumstances.

(c) Physician personally performs the anesthesia procedure.

(1) HCFA considers an anesthesia service to be personally performed under any of the following circumstances:

(i) The physician performs the entire anesthesia service alone.

(ii) The physician establishes an attending physician relationship in one or two concurrent cases involving an intern or resident and the service was furnished before January 1, 1994.

(iii) The physician establishes an attending physician relationship in one case involving an intern or resident and the service was furnished on or after January 1, 1994 but prior to January 1, 1996. For services on or after January 1, 1996, the physician must be the teaching physician as defined in §§ 415.170 through 415.184 of this chapter.

(iv) The physician and the CRNA or AA are involved in a single case and the services of each are found to be medically necessary.

(v) The physician is continuously involved in a single case involving a student nurse anesthetist.

(vi) The physician is continuously involved in a single case involving a CRNA or AA and the service was furnished prior to January 1, 1996.
(2) HCFA determines the fee schedule amount for an anesthesia service personally performed by a physician on the basis of an anesthesia-specific fee schedule CF and unreduced base units and anesthesia time units. One anesthesia time unit is equivalent to 15 minutes of anesthesia time, and fractions of a 15-minute period are recognized as fractions of an anesthesia time unit.

(d) Anesthesia services medically directed by a physician. (1) HCFA considers an anesthesia service to be medically directed by a physician if:

(i) The physician performs the activities described in §415.110 of this chapter.

(ii) The physician directs qualified individuals involved in two, three, or four concurrent cases.

(iii) Medical direction can occur for a single case furnished on or after January 1, 1998 if the physician performs the activities described in §415.110 of this chapter and medically directs a single CRNA or AA.

(2) The rules for medical direction differ for certain time periods depending on the nature of the qualified individual who is directed by the physician. If more than two procedures are directed on or after January 1, 1994, the qualified individuals could be AAs, CRNAs, interns, or residents. The medical direction rules apply to student nurse anesthetists only if the physician directs one case involving a student nurse anesthetist and the other involving a CRNA, AA, intern, or resident.

(3) Payment for medical direction is based on a specific percentage of the payment allowance recognized for the anesthesia service personally performed by a physician alone. The following percentages apply for the years specified:

(i) CY 1994—60 percent of the payment allowance for personally performed procedures.

(ii) CY 1995—57.5 percent of the payment allowance for personally performed services.

(iii) CY 1996—55 percent of the payment allowance for personally performed services.

(iv) CY 1997—52.5 percent of the payment allowance for personally performed services.

(v) CY 1998 and thereafter—50 percent of the payment allowance for personally performed services.

(e) Physician medically supervises anesthesia services. If the physician medically supervises more than four concurrent anesthesia services, HCFA bases the fee schedule amount on an anesthesia-specific CF and three base units. This represents payment for the physician’s involvement in the pre-surgical anesthesia services.

(f) Payment for medical or surgical services furnished by a physician while furnishing anesthesia services. (1) HCFA allows separate payment under the fee schedule for certain reasonable and medically necessary medical or surgical services furnished by a physician while furnishing anesthesia services to the patient. HCFA makes payment for these services in accordance with the general physician fee schedule rules in §414.20. These services are described in program operating instructions.

(2) HCFA makes no separate payment for other medical or surgical services, such as the pre-anesthetic examination of the patient, pre- or post-operative visits, or usual monitoring functions, that are ordinarily included in the anesthesia service.

(g) Physician involved in multiple anesthesia services. If the physician is involved in multiple anesthesia services for the same patient during the same operative session, the carrier makes payment according to the base unit associated with the anesthesia service having the highest base unit value and anesthesia time that encompasses the multiple services.

§414.48 Limits on actual charges of nonparticipating suppliers.

(a) General rule. A supplier, as defined in §400.202 of this chapter, who is nonparticipating and does not accept assignment may charge a beneficiary an amount up to the limiting charge described in paragraph (b) of this section.
§ 414.50 Physician billing for purchased diagnostic tests.

(a) General rule. For services covered under section 1861(s)(3) of the Act and paid for under this part 414 subpart A, if a physician bills for a diagnostic test performed by an outside supplier, the payment to the physician less the applicable deductibles and coinsurance may not exceed the lowest of the following amounts:

1. The supplier's net charge to the physician.
2. The physician's actual charge.
3. The fee schedule amount for the test that would be allowed if the supplier billed directly.

(b) Restriction on payment. The physician must identify the supplier and indicate the supplier's net charge for the test. If the physician fails to provide this information, HCFA makes no payment to the physician and the physician may not bill the beneficiary.

(1) Physicians who accept Medicare assignment may bill beneficiaries for only the applicable deductibles and coinsurance.

(2) Physicians who do not accept Medicare assignment may not bill the beneficiary more than the payment amount described in paragraph (a) of this section.


§ 414.52 Payment for physician assistants' services.

Allowed amounts for the services of a physician assistant furnished beginning January 1, 1992 and ending December 31, 1997 may not exceed the limits specified in paragraphs (a) through (c) of this section. Allowed amounts for the services of a physician assistant furnished beginning January 1, 1998, may not exceed the limits specified in paragraph (d) of this section.

(a) For assistant-at-surgery services, 65 percent of the amount that would be allowed under the physician fee schedule if the assistant-at-surgery service was furnished by a physician.

(b) For services (other than assistant-at-surgery services) furnished in a hospital, 75 percent of the physician fee schedule amount for the service.

(c) For all other services, 85 percent of the physician fee schedule amount for the service.

(d) For services (other than assistant-at-surgery services) furnished beginning January 1, 1998, 85 percent of the physician fee schedule amount for the service. For assistant-at-surgery services, 85 percent of the physician fee schedule amount that would be allowed under the physician fee schedule if the assistant-at-surgery service were furnished by a physician.


§ 414.54 Payment for certified nurse-midwives' services.

For services furnished after December 31, 1991, allowed amounts under the fee schedule established under section 1833(a)(1)(K) of the Act for the payment of certified nurse-midwife services may not exceed 65 percent of the physician fee schedule amount for the service.

§ 414.56 Payment for nurse practitioners' and clinical nurse specialists' services.

(a) Rural areas. For services furnished beginning January 1, 1992 and ending December 31, 1997, allowed amounts for the services of a nurse practitioner or a clinical nurse specialist in a rural area (as described in section 1861(s)(2)(K)(iii) of the Act) may not exceed the following limits:

1. For services furnished in a hospital (including assistant-at-surgery services), 75 percent of the physician fee schedule amount for the service.

2. For all other services, 85 percent of the physician fee schedule amount for the service.
§ 414.65 Payment for consultations via interactive telecommunications systems.

(a) Limitations on payment. Medicare payment for a professional consultation conducted via interactive telecommunications systems is subject to the following limitations:

(1) The payment may not exceed the current fee schedule amount applicable to the consulting practitioner for the health care service provided.

(2) The payment may not include reimbursement for any telephone line charges or any facility fees.

(3) The payment is subject to the co-insurance and deductible requirements of sections 1833(a)(1) and (b) of the Act.
§ 414.200 Purpose.

This subpart implements sections 1834 (a) and (h) of the Act by specifying how payments are made for the purchase or rental of new and used durable medical equipment and prosthetic and orthotic devices for Medicare beneficiaries.

[57 FR 57689, Dec. 7, 1992]

§ 414.202 Definitions.

For purposes of this subpart, the following definitions apply:

Covered item update means the percentage increase in the consumer price index for all urban consumers (U.S. city average) (CPI-U) for the 12-month period ending with June of the previous year.

Durable medical equipment means equipment, furnished by a supplier or a home health agency that—

(1) Can withstand repeated use;
(2) Is primarily and customarily used to serve a medical purpose;
(3) Generally is not useful to an individual in the absence of an illness or injury; and
(4) Is appropriate for use in the home.

(See §410.38 of this chapter for a description of when an institution qualifies as a home.)

Prosthetic and orthotic devices means—

(1) Devices that replace all or part of an internal body organ, including ostomy bags and supplies directly related to ostomy care, and replacement of such devices and supplies;
(2) One pair of conventional eyeglasses or contact lenses furnished subsequent to each cataract surgery with insertion of an intraocular lens; and
(3) Leg, arm, back, and neck braces, and artificial legs, arms, and eyes, including replacements if required because of a change in the beneficiary's physical condition.

The following are neither prosthetic nor orthotic devices—

(1) Parenteral and enteral nutrients, supplies, and equipment;
(2) Intraocular lenses;
(3) Medical supplies such as catheters, catheter supplies, ostomy bags, and supplies related to ostomy care that are furnished by an HHA as part of home health services under §409.40(e) of this chapter;
(4) Dental prostheses.

Region means those carrier service areas administered by HCFA regional offices.

[57 FR 57689, Dec. 7, 1992]

§ 414.210 General payment rules.

(a) General rule. For items furnished on or after January 1, 1999, except as provided in paragraphs (c) and (d) of this section, Medicare pays for durable medical equipment, prosthetics and orthotics, including a separate payment for maintenance and servicing of the items as described in paragraph (e)
of this section, on the basis of 80 percent of the lesser of—
(1) The actual charge for the item;
(2) The fee schedule amount for the item, as determined in accordance with the provisions of §§414.220 through 414.232.

(b) Payment classification. (1) The carrier determines fee schedules for the following classes of equipment and devices:
(i) Inexpensive or routinely purchased items, as specified in §414.220.
(ii) Items requiring frequent and substantial servicing, as specified in §414.222.
(iii) Certain customized items, as specified in §414.224.
(iv) Oxygen and oxygen equipment, as specified in §414.226.
(v) Prosthetic and orthotic devices, as specified in §414.228.
(vi) Other durable medical equipment (capped rental items), as specified in §414.229.
(vii) Transcutaneous electrical nerve stimulators (TENS), as specified in §414.232.

(2) HCFA designates the items in each class of equipment or device through its program instructions.

(c) Exception for certain HHAs. Public HHAs and HHAs that furnish services or items free-of-charge or at nominal prices to a significant number of low-income patients, as defined in §413.13(a) of this chapter, are paid on the basis of 80 percent of the fee schedule amount determined in accordance with the provision of §§414.220 through 414.230.

(d) Prohibition on special limits. For items furnished on or after January 1, 1989 and before January 1, 1991, neither HCFA nor a carrier may establish a special reasonable charge for items covered under this subpart on the basis of inherent reasonableness as described in §405.502(g) of this chapter.

(e) Maintenance and servicing. (1) General rule. Except as provided in paragraph (e)(2) of this section, the carrier pays the reasonable and necessary charges for maintenance and servicing of purchased equipment. Reasonable and necessary charges are those made for parts and labor not otherwise covered under a manufacturer’s or supplier’s warranty. Payment is made, as needed, in a lump sum based on the carrier’s consideration of the item. Payment is not made for maintenance and servicing of a rented item other than the maintenance and servicing fee for other durable medical equipment, as described in §414.229(e).

(2) Exception. For items purchased on or after June 1, 1989, no payment is made under the provisions of paragraph (e)(1) of this section for the maintenance and servicing of:
(i) Items requiring frequent and substantial servicing, as defined in §414.222(a);
(ii) Capped rental items, as defined in §414.229(a), that are not purchased in accordance with §414.229(d); and
(iii) Oxygen equipment, as defined in §414.226.

(f) Replacement of equipment. Except as provided in §414.229(g), if a purchased item of DME or a prosthetic or orthotic device paid for under this subpart has been in continuous use by the patient for the equipment’s reasonable useful lifetime or if the carrier determines that the item is lost or irrepairably damaged, the patient may elect to obtain a new piece of equipment.

(1) The reasonable useful lifetime of DME or prosthetic and orthotic devices is determined through program instructions. In the absence of program instructions, carriers may determine the reasonable useful lifetime of equipment but in no case can it be less than 5 years. Computation is based on the age of the equipment.

(2) If the beneficiary elects to obtain replacement equipment, payment is made on a purchase basis.

[57 FR 57689, Dec. 7, 1992]

§ 414.220 Inexpensive or routinely purchased items.

(a) Definitions. (1) Inexpensive equipment means equipment the average purchase price of which did not exceed $150 during the period July 1, 1986 through July 1987.

(2) Routinely purchased equipment means equipment that was acquired on a national basis at least 75 percent of the time during the period July 1986 through June 1987.

(3) Accessories. Effective January 1, 1994, accessories used in conjunction
with a nebulizer, aspirator, or ventilator excluded from §414.222 meet the definitions of "inexpensive equipment" and "routinely purchased equipment" in paragraphs (a)(1) and (a)(2) of this section, respectively.

(b) Payment rules. (1) Subject to the limitation in paragraph (b)(3) of this section, payment for inexpensive and routinely purchased items is made on a rental basis or in a lump sum amount for purchase of the item based on the applicable fee schedule amount.

(2) Effective January 1, 1994, payment for ostomy supplies, tracheostomy supplies, urologicals, and surgical dressings not furnished as incident to a physician's professional service or furnished by an HHA is made using the methodology for the inexpensive and routinely purchased class.

(3) The total amount of payments made for an item may not exceed the fee schedule amount recognized for the purchase of that item.

(c) Fee schedule amount for 1989 and 1990. The fee schedule amount for payment of purchase or rental of inexpensive or routinely purchased items furnished in 1989 and 1990 is the local payment amount determined as follows:

(1) The carrier determines the average reasonable charge for inexpensive or routinely purchased items that were furnished during the period July 1, 1986 through June 30, 1987 based on the mean of the carrier's allowed charges for the item. A separate determination of an average reasonable charge is made for rental equipment, new purchased equipment, and used purchased equipment.

(2) The carrier adjusts the amount determined under paragraph (c)(1) of this section by the change in the level of the CPI-U for the 6-month period ending December 1987.

(d) Updating the local payment amounts for years after 1990. For each year subsequent to 1990, the local payment amounts of the preceding year are increased or decreased by the covered item update. For 1991 and 1992, the covered item update is reduced by 1 percentage point.

(e) Calculating the fee schedule amounts for years after 1990. For years after 1990, the fee schedule amounts are equal to the national limited payment amount.

(f) Calculating the national limited payment amount. The national limited payment amount is computed as follows:

(1) The 1991 national limited payment amount is equal to:

(i) 100 percent of the local payment amount if the local payment amount is neither greater than the weighted average nor less than 85 percent of the weighted average of all local payment amounts;

(ii) The sum of 67 percent of the local payment amount plus 33 percent of the weighted average of all local payment amounts if the local payment amount exceeds the weighted average of all local payment amounts; or

(iii) The sum of 67 percent of the local payment amount plus 33 percent of 85 percent of the weighted average of all local payment amounts if the local payment amount is less than 85 percent of the weighted average of all local payment amounts.

(2) The 1992 national limited payment amount is equal to:

(i) 100 percent of the local payment amount if the local payment amount is neither greater than the weighted average nor less than 85 percent of the weighted average of all local payment amounts;

(ii) The sum of 33 percent of the local payment amount plus 67 percent of the weighted average of all local payment amounts if the local payment amount exceeds the weighted average; or

(iii) The sum of 33 percent of the local payment amount plus 67 percent of 85 percent of the weighted average of all local payment amounts if the local payment amount is less than 85 percent of the weighted average.

(3) For 1993, the national limited payment amount is equal to one of the following:

(i) 100 percent of the local payment amount if the local payment amount is neither greater than the weighted average nor less than 85 percent of the weighted average of all local payment amounts.

(ii) 100 percent of the weighted average of all local payment amounts if the local payment amount exceeds the weighted average of all local payment amounts.
§ 414.224 Customized items.

(a) Criteria for a customized item. To be considered a customized item for payment purposes under paragraph (b) of this section, a covered item (including a wheelchair) must be uniquely constructed or substantially modified for a specific beneficiary according to the description and orders of a physician and be so different from another item used for the same purpose that the two items cannot be grouped together for pricing purposes.

§ 414.224 Customized items.

(a) Criteria for a customized item. To be considered a customized item for payment purposes under paragraph (b) of this section, a covered item (including a wheelchair) must be uniquely constructed or substantially modified for a specific beneficiary according to the description and orders of a physician and be so different from another item used for the same purpose that the two items cannot be grouped together for pricing purposes.

(iii) 85 percent of the weighted average of all local payment amounts if the local payment amount is less than 85 percent of the weighted average of all local payment amounts.

(4) For 1994 and subsequent years, the national limited payment amount is equal to one of the following:

(i) If the local payment amount is not in excess of the median nor less than 85 percent of the median of all local payment amounts—100 percent of the local payment amount.

(ii) If the local payment amount exceeds the median—100 percent of the median of all local payment amounts.

(iii) If the local payment amount is less than 85 percent of the median—85 percent of the median of all local payment amounts.

(g) Payment for surgical dressings. For surgical dressings furnished after December 31, 1993, the national limited payment amount is computed based on local payment amounts using average reasonable charges for the 12-month period ending December 31, 1992, increased by the covered item updates for 1993 and 1994.

[57 FR 57689, Dec. 7, 1992, as amended at 60 FR 35497, July 10, 1995]
§ 414.226 Oxygen and oxygen equipment.

(a) Payment rules. (1) Payment for rental of oxygen equipment and purchase of oxygen contents is made based on a monthly fee schedule amount.

(2) Monthly fee schedule payments continue until medical necessity ends.

(b) Monthly fee schedule amount. (1) Monthly fee schedule amounts are separately calculated for the following items:

(i) Stationary oxygen equipment and oxygen contents (stationary and portable oxygen contents).

(ii) Portable oxygen equipment only.

(iii) Stationary and portable oxygen contents only.

(iv) Portable oxygen contents only.

(2) For 1989 and 1990, the monthly fee schedule amounts are the local payment amounts determined as follows:

(i) The carrier determines the base local average monthly payment rate equal to the total reasonable charges for the item for the 12-month period ending December 1986 divided by the total number of months for all beneficiaries receiving the item for the same period. In determining the local average monthly payment rate, the following limitations apply:

(A) Purchase charges for oxygen systems are not included as items classified under paragraph (b)(1)(i) of this section.

(B) Purchase charges for portable equipment are not included as items classified under paragraph (b)(1)(ii) of this section.

(ii) The carrier determines the local monthly payment amount equal to 0.95 times the base local average monthly payment amount adjusted by the change in the CPI-U for the six-month period ending December 1987.

(3) For years after 1990, the fee schedule amounts are determined using the methodology contained in § 414.220 (d), (e), and (f).

(c) Application of monthly fee schedule amounts. (1) The fee schedule amount for items described in paragraph (b)(1)(i) of this section is paid when the beneficiary rents a stationary oxygen system.

(2) Subject to the limitation set forth in paragraph (d)(2) of this section, the fee schedule amount for items described in paragraph (b)(1)(ii) of this section is paid when the beneficiary owns a stationary gaseous or liquid oxygen system.

(3) The fee schedule amount for items described in paragraph (b)(1)(iii) of this section is paid when the beneficiary owns or rents a portable gaseous or portable liquid oxygen system and uses either a stationary oxygen concentrator or no stationary oxygen system.

(4) The fee schedule amount for items described in paragraph (b)(1)(iv) of this section is paid when the beneficiary owns or rents a portable gaseous or portable liquid oxygen system and uses a stationary oxygen concentrator.

(d) Volume adjustments: (1) The fee schedule amount for an item described in paragraph (b)(1)(i) of this section is adjusted as follows:

(i) If the attending physician prescribes an oxygen flow rate exceeding four liters per minute, the fee schedule amount is increased by 50 percent, subject to the limit in paragraph (d)(2) of this section.

(ii) If the attending physician prescribes an oxygen flow rate of less than one liter per minute, the fee schedule amount is decreased by 50 percent.

(2) If portable oxygen equipment is used and the prescribed oxygen flow rate exceeds four liters per minute, the total fee schedule amount recognized for payment is limited to the higher of—

(i) The sum of the monthly fee schedule amount for the items described in paragraphs (b)(1)(i) and (ii) of this section; or

(ii) The adjusted fee schedule amount described in paragraph (d)(1)(i) of this section.
§ 414.229 Other durable medical equipment—capped rental items.
(a) General payment rule. Subject to the limitation set forth in paragraph (b) of this section, payment is made on a rental or purchase option basis for other durable medical equipment that is not subject to the payment provisions set forth in §§ 414.220 through 414.228.

§ 414.228 Prosthetic and orthotic devices.
(a) Payment rule. Payment is made on a lump-sum basis for prosthetic and orthotic devices subject to this subpart.
(b) Fee schedule amounts. The fee schedule amount for prosthetic and orthotic devices is determined as follows:
(1) The carrier determines a base local purchase price equal to the average reasonable charge for items purchased during the period July 1, 1986 through June 30, 1987 based on the mean of the carrier’s allowed charges for the item.
(2) The carrier determines a local purchase price equal to the following:
(i) For 1989 and 1990, the base local purchase price is adjusted by the change in the level of the CPI-U for the 6-month period ending December 1987.
(ii) For 1991 through 1993, the local purchase price for the preceding year is adjusted by the applicable percentage increase for the year. The applicable percentage increase is equal to 0 percent for 1991. For 1992 and 1993, the applicable percentage increase is equal to the percentage increase in the CPI-U for the 12-month period ending with June of the previous year.
(iii) For 1994 and 1995, the applicable percentage increase is 0 percent.
(iv) For all subsequent years the applicable percentage increase is equal to the percentage increase in the CPI-U for the 12-month period ending with June of the previous year.
(3) HCFA determines the regional purchase price equal to the following:
(i) For 1992, the average (weighted by the relative volume of all claims among carriers) of the local purchase prices for the carriers in the region.
(ii) For 1993 and subsequent years, the regional purchase price for the preceding year adjusted by the applicable percentage increase for the year.
(4) HCFA determines a purchase price equal to the following:
(i) For 1989, 1990 and 1991, 100 percent of the local purchase price.
(ii) For 1992, 75 percent of the local purchase price plus 25 percent of the regional purchase price.
(iii) For 1993, 50 percent of the local purchase price plus 50 percent of the regional purchase price.
(iv) For 1994 and subsequent years, 100 percent of the regional purchase price.
(5) For 1992 and subsequent years, HCFA determines a national average purchase price equal to the unweighted average of the purchase prices determined under paragraph (b)(4) of this section for all carriers.
(6) HCFA determines the fee schedule amount equal to 100 percent of the purchase price determined under paragraph (b)(4) of this section, subject to the following limitations:
(i) For 1992, the amount cannot be greater than 125 percent nor less than 85 percent of the national average purchase price determined under paragraph (b)(5) of this section.
(ii) For 1993 and subsequent years, the amount cannot be greater than 120 percent of the national average nor less than 90 percent of the national average purchase price determined under paragraph (b)(5) of this section.

§ 414.229 Other durable medical equipment—capped rental items.
(a) General payment rule. Subject to the limitation set forth in paragraph (b) of this section, payment is made on a rental or purchase option basis for other durable medical equipment that is not subject to the payment provisions set forth in §§ 414.220 through 414.228.
(b) Fee schedule amounts for rental. (1) For 1989 and 1990, the monthly fee schedule amount for rental of other covered durable medical equipment equals 10 percent of the purchase price recognized as determined under paragraph (c) of this section subject to the following limitation: For 1989 and 1990, the fee schedule amount cannot be greater than 115 percent nor less than 85 percent of the prevailing charge, as determined under §405.504 of this chapter, established for rental of the item in January 1987, as adjusted by the change in the level of the CPI±U for the 6-month period ending December 1987.

(2) For 1991 and subsequent years, the monthly fee schedule amount for rental of other covered durable medical equipment equals 10 percent of the purchase price recognized as determined under paragraph (c) of this section for each of the first 3 months and 7.5 percent of the purchase price for each of the remaining months.

c) Determination of purchase price. The purchase price of other covered durable medical equipment is determined as follows:

(1) For 1989 and 1990. (i) The carrier determines a base local purchase price amount equal to the average of the purchase prices submitted on an assignment-related basis of new items supplied during the 6-month period ending December 1986.

(ii) The purchase price is equal to the base local purchase price adjusted by the change in the level of the CPI±U for the 6-month period ending December 1987.

(2) For 1991. (i) The local payment amount is the purchase price for the preceding year adjusted by the covered item update for 1991 and decreased by the percentage by which the average of the reasonable charges for claims paid for all other items described in §414.229, is lower than the average of the purchase prices submitted for such items during the final 9 months of 1988.

(ii) The purchase price for 1991 is the national limited payment amount as determined using the methodology contained in §414.220.(f).

(iii) The purchase price for years after 1991. The purchase price is determined using the methodology contained in paragraphs (d) through (f) of §414.220.

d) Purchase option. Suppliers must offer a purchase option to beneficiaries during the 10th continuous rental month and, for power-driven wheelchairs, the purchase option must also be made available at the time the equipment is initially furnished.

(1) Suppliers must offer beneficiaries the option of purchasing power-driven wheelchairs at the time the supplier first furnishes the item. Payment must be on a lump-sum fee schedule purchase basis if the beneficiary chooses the purchase option. The purchase fee is the amount established in §414.229(c).

(2) Suppliers must offer beneficiaries the option of converting capped rental items (including power-driven wheelchairs not purchased when initially furnished) to purchased equipment during their 10th continuous rental month. Beneficiaries have one month from the date the supplier makes the offer to accept the purchase option.

(i) If the beneficiary does not accept the purchase option, payment continues on a rental basis not to exceed a period of continuous use of longer than 15 months. After 15 months of rental payments have been paid, the supplier must continue to provide the item without charge, other than a charge for maintenance and servicing fees, until medical necessity ends or Medicare coverage ceases. A period of continuous use is determined under the provisions in §414.230.

(ii) If the beneficiary accepts the purchase option, payment continues on a rental basis not to exceed a period of continuous use of longer than 13 months. On the first day after 13 continuous rental months during which payment is made, the supplier must transfer title to the equipment to the beneficiary.

(e) Payment for maintenance and servicing. (1) The carrier establishes a reasonable fee for maintenance and servicing for each rented item of other durable medical equipment. The fee may not exceed 10 percent of the purchase price recognized as determined under paragraph (c) of this section.

(2) Payment of the fee for maintenance and servicing of other durable medical equipment that is rented is made only for equipment that continues to be used after 15 months of
rental payments have been made and is limited to the following:

(i) For the first 6-month period, no payments are to be made.

(ii) For each succeeding 6-month period, payment may be made during the first month of that period.

(3) Payment for maintenance and servicing DME purchased in accordance with paragraphs (d)(1) and (d)(2)(ii) of this section, is made on the basis of reasonable and necessary charges.

(f) Transition to the fee schedules. For purposes of computing the 10-month or 15-month period of continuous use for other durable medical equipment, as described in §414.230, the carrier counts the first month that the beneficiary continued renting the equipment without regard to whether that month occurred before January 1, 1989 or after. If a beneficiary’s 15-month rental period ends prior to January 1, 1989, no further purchase or rental payments are to be made except for maintenance and servicing of equipment as described in paragraph (e) of this section.

(g) Replacement of equipment. If the item of equipment has been in continuous use by the patient on either a rental or purchase basis for the equipment’s useful lifetime, or if the carrier determines that the item is lost or irreparably damaged, the patient may elect to obtain a new piece of equipment.

(1) The reasonable useful lifetime of DME or prosthetic and orthotic devices is determined through program instructions. In the absence of program instructions, carriers may determine the reasonable useful lifetime of equipment but in no case can it be less than 5 years. Computation is based on when the equipment is delivered to the beneficiary, not the age of the equipment.

(2) If the beneficiary elects to obtain replacement equipment, payment is made on a rental or purchase basis in accordance with paragraph (a) of this section or on a lump-sum purchase basis if a purchase agreement had been entered into in accordance with paragraph (d) of this section.

§414.230 Determining a period of continuous use.

(a) Scope. This section sets forth the rules that apply in determining a period of continuous use for rental of durable medical equipment.

(b) Continuous use. A period of continuous use begins with the first month of medical need and lasts until a beneficiary’s medical need for a particular item of durable medical equipment ends.

(c) Temporary interruption. (1) A period of continuous use allows for temporary interruptions in the use of equipment.

(2) An interruption of not longer than 60 consecutive days plus the days remaining in the rental month in which use ceases is temporary, regardless of the reason for the interruption.

(3) Unless there is a break in medical necessity that lasts longer than 60 consecutive days plus the days remaining in the rental month in which use ceases, medical necessity is presumed to continue.

(d) Criteria for a new rental period. If an interruption in the use of equipment continues for more than 60 consecutive days plus the days remaining in the rental month in which use ceases, a new rental period begins if the supplier submits all of the following information—

(1) A new prescription.

(2) New medical necessity documentation.

(3) A statement describing the reason for the interruption and demonstrating that medical necessity in the prior episode ended.

(e) Beneficiary moves. A permanent or temporary move made by a beneficiary does not constitute an interruption in the period of continuous use.

(f) New equipment. If a beneficiary changes equipment or requires additional equipment based on a physician’s prescription, and the new or additional equipment is found to be necessary, a new period of continuous use begins for the new or additional equipment. A new period of continuous use does not begin for base equipment that is modified by an addition.

[57 FR 57691, Dec. 7, 1992, as amended at 60 FR 35468, July 10, 1995]
§ 414.232 Special payment rules for transcutaneous electrical nerve stimulators (TENS).

(a) General payment rule. Except as provided in paragraph (b) of this section, payment for TENS is made on a purchase basis with the purchase price determined using the methodology for purchase of inexpensive or routinely purchased items as described in §414.220. The payment amount for TENS computed under §414.220(c)(2) is reduced according to the following formula:

(1) Effective April 1, 1990—the original payment amount is reduced by 15 percent.

(2) Effective January 1, 1991—the reduced payment amount in paragraph (a)(1) is reduced by 15 percent.

(3) Effective January 1, 1994—the reduced payment amount in paragraph (a)(1) is reduced by 45 percent.

(b) Exception. In order to permit an attending physician time to determine whether the purchase of the TENS is medically appropriate for a particular patient, two months of rental payments may be made in addition to the purchase price. The rental payments are equal to 10 percent of the purchase price.

[57 FR 57692, Dec. 7, 1992, as amended at 60 FR 35498, July 10, 1995]

Subpart E—Determination of Reasonable Charges Under the ESRD Program

§ 414.300 Scope of subpart.

This subpart sets forth criteria and procedures for payment of the following services furnished to ESRD patients:

(a) Physician services related to renal dialysis.

(b) Physician services related to renal transplantation.

(c) Home dialysis equipment, supplies, and support services.

(d) Epoetin (EPO) furnished by a supplier of home dialysis equipment and supplies to a home dialysis patient for use in the home.

the facility and ends when the patient departs from the facility. In the case of home dialysis, the period begins when the patient prepares for dialysis and generally ends when the patient is disconnected from the machine. In this context, a dialysis facility includes only those parts of the building used as a facility. It does not include any areas used as a physician’s office.

Medical direction, in contrast to supervision of staff, is a routine professional service that entails substantial direct involvement and the physical presence of the physician in the delivery of services directly to the patient.

Routine professional services include all physicians’ services furnished during a dialysis session and all services listed in paragraph (d) of this section that meet the following requirements:

(1) They are personally furnished by a physician to an individual patient.
(2) They contribute directly to the diagnosis or treatment of an individual patient.
(3) They ordinarily must be performed by a physician.

Supervision of staff, in contrast to medical direction, is an administrative service that does not necessarily require the physician to be present at the dialysis session. It is a general activity primarily concerned with monitoring performance of and giving guidance to other health care personnel (such as nurses and dialysis technicians) who deliver services to patients.

(d) Types of routine professional services. Routine professional services include at least all of the following services when medically appropriate:

(1) Visits to the patient during dialysis, and review of laboratory test results, nurses’ notes and any other medical documentation, as a basis for—
   (i) Adjustment of the patient’s medication or diet, or the dialysis procedure;
   (ii) Prescription of medical supplies; and
   (iii) Evaluation of the patient’s psychosocial status and the appropriateness of the treatment modality.

(2) Medical direction of staff in delivering services to a patient during a dialysis session.

(3) Pre-dialysis and post-dialysis examinations, or examinations that could have been furnished on a pre-dialysis or post-dialysis basis.

(4) Insertion of catheters for patients who are on peritoneal dialysis and do not have indwelling catheters.

(e) Payment for routine professional services. Beginning August 7, 1990, routine professional services furnished by physicians may be paid under either the “initial method” of payment described in §414.313, (if all of the physicians at the facility elect the initial method) or under the “physician MCP method” described in §414.314. Physician services furnished after July 31, 1983 and before August 6, 1990, are payable only under the MCP method described in §414.314.

§ 414.313 Initial method of payment.

(a) Basic rule. Under this method, the intermediary pays the facility for routine professional services furnished by physicians. Payment is in the form of an add-on to the facility’s composite rate payment, which is described in part 413, subpart H of this subchapter.

(b) Services for which payment is not included in the add-on payment. (1) Physician administrative services are considered to be facility services and are paid for as part of the facility’s composite rate.

(2) The carrier pays the physician or the beneficiary (as appropriate) under the reasonable charge criteria set forth in subpart E of part 405 of this chapter for the following services:

(i) Physician services that must be furnished at a time other than during the dialysis session (excluding pre-dialysis and post-dialysis examinations and examinations that could have been furnished on a pre-dialysis or post-dialysis basis), such as monthly and semi-annual examinations to review health status and treatment.

(ii) Physician surgical services other than insertion of catheters for patients who are on peritoneal dialysis and do not have indwelling catheters.

(iii) Physician services furnished to hospital inpatients who were not admitted solely to receive maintenance dialysis.

(iv) Administration of hepatitis B vaccine.

(c) Physician election of the initial method. (1) Each physician in a facility
§ 414.314 Monthly capitation payment method.

(a) Basic rules. (1) Under the monthly capitation payment (MCP) method, the carrier pays an MCP amount for each patient, to cover all professional services furnished by the physician, except those listed in paragraph (b) of this section.

(2) The carrier pays the MCP amount, subject to the deductible and coinsurance provisions, either to the physician if the physician accepts assignment or to the beneficiary if the physician does not accept assignment.

(3) The MCP method recognizes the need of maintenance dialysis patients for physician services furnished periodically over relatively long periods of time, and the capitation amounts are consistent with physicians’ charging patterns in their localities.

(4) Payment of the capitation amount for any particular month is contingent upon the physician furnishing to the patient all physician services required by the patient during the month, except those listed in paragraph (b) of this section.

(b) Services not included in the MCP.

(1) Services that are not included in the MCP and which may be paid in accordance with the reasonable charge rules set forth in subpart E of part 405 of this chapter are limited to the following:

(i) Administration of hepatitis B vaccine.

(ii) Covered physician services furnished by another physician when the patient is not available to receive, or the attending physician is not available to furnish, the outpatient services as usual (see paragraph (b)(3) of this section).

(iii) Covered physician services furnished to hospital inpatients, including services related to inpatient dialysis, by a physician who elects not to continue to receive the MCP during the period of inpatient stay.

(iv) Surgical services, including declotting of shunts, other than the insertion of catheters for patients on maintenance peritoneal dialysis who do not have indwelling catheters.

(v) Needed physician services that are—
§ 414.320 Determination of reasonable charges for physician renal transplantation services.

(a) Comprehensive payment for services furnished during a 60-day period. (1) The comprehensive payment is subject to the deductible and coinsurance provisions and is for all surgeon services furnished during a period of 60 days in connection with a renal transplantation, including the usual preoperative and postoperative care, and for immunosuppressant therapy if supervised by the transplant surgeon.

(2) Additional sums, in amounts established on the basis of program experience, may be included in the comprehensive payment for other surgery performed concurrently with the transplant operation.

(3) The amount of the comprehensive payment may not exceed the lower of the following:

(i) The actual charges made for the services.

(ii) Overall national payment levels established under the ESRD program and adjusted to give effect to variations in physician's charges throughout the nation. (These adjusted amounts are the maximum allowances in a carrier's service area for renal transplantation surgery and related services by surgeons.)

(4) Maximum allowances computed under these instructions are revised at the beginning of each calendar year to the extent permitted by the lesser of the following:

(i) Changes in the economic index as described in §405.504(a)(3)(i) of this chapter.

(ii) Percentage changes in the weighted average of the carrier's prevailing charges (before adjustment by the economic index) for—

(A) A unilateral nephrectomy; or

(B) Another medical or surgical service designated by HCFA for this purpose.

(b) Other payments. Payments for covered medical services furnished to the transplant recipient by other specialists, as well as for services by the

§ 414.316 Payment for physician services to patients in training for self-dialysis and home dialysis.

(a) For each patient, the carrier pays a flat amount that covers all physician services required to create the capacity for self-dialysis and home dialysis.

(b) HCFA determines the amount on the basis of program experience and reviews it periodically.

(c) The payment is made at the end of the training course, is subject to the deductible and coinsurance provisions, and is in addition to any amounts payable under the initial or MCP methods set forth in §§414.313 and 414.314, respectively.

(d) If the training is not completed, the payment amount is proportionate to the time spent in training.

§ 414.312 Determination of reasonable charges for diagnostic radiology services.

(a) Payment is made under the Medicare fee schedule for all services furnished to patients for the purposes of providing diagnostic radiology services. (The fee schedule is described in §§414.300 through 414.304, inclusive.)

(b) Exception. The amount of payment for X-ray services in connection with transplant surgery is determined under §414.320.
transplant surgeon after the 60-day period covered by the comprehensive payment, are made under the reasonable charge criteria set forth in §405.502 (a) through (d) of this chapter. The payments for physicians’ services in connection with renal transplantations are changed on the basis of program experience and the expected advances in the medical art for this operation.

§414.330 Payment for home dialysis equipment, supplies, and support services.

(a) Equipment and supplies—(1) Basic rule. Except as provided in paragraph (a)(2) of this section, Medicare pays for home dialysis equipment and supplies only under the prospective payment rates established at §413.170.

(2) Exception. If the conditions in subparagraphs (a)(2)(i) through (iv) of this section are met, Medicare pays for home analysis equipment and supplies on a reasonable charge basis in accordance with subpart E (Criteria for Determination of Reasonable Charges; Reimbursement for Services of Hospital Interns, Residents, and Supervising Physicians) of part 405, but the amount of payment may not exceed the limit for equipment and supplies in paragraph (c)(2) of this section.

(i) The patient elects to obtain home dialysis equipment and supplies from a supplier that is not a Medicare approved dialysis facility.

(ii) The patient certifies to HCFA that he or she has only one supplier for all home dialysis equipment and supplies. This certification is made on HCFA Form 382 (the “ESRD Beneficiary Selection” form).

(iii) In writing, the supplier—

(A) Agrees to receive Medicare payment for home dialysis supplies and equipment only on an assignment-related basis; and

(B) Certifies to HCFA that it has a written agreement with one Medicare approved dialysis facility or, if the beneficiary is also entitled to military or veteran’s benefits, one military or Veterans Administration hospital, for each patient. (See subpart U of part 405 of this chapter for the requirements for a Medicare approved dialysis facility.) Under the agreement, the facility or military or VA hospital agrees to the following:

(1) To furnish all home dialysis support services for each patient in accordance with subpart U (Conditions for Coverage of Suppliers of ESRD Services) of this chapter. (§410.52 sets forth the scope and conditions of Medicare Part B coverage of home dialysis services, supplies, and equipment.)

(2) To furnish institutional dialysis services and supplies. (§410.50 sets forth the scope and conditions for Medicare Part B coverage of institutional dialysis services and supplies.)

(3) To furnish dialysis-related emergency services.

(4) To arrange for a Medicare approved laboratory to perform dialysis-related laboratory tests that are covered under the composite rate established at §413.170 and to arrange for the laboratory to seek payment from the facility. The facility then includes these laboratory services in its claim for payment for home dialysis support services.

(5) To arrange for a Medicare approved laboratory to perform dialysis-related laboratory tests that are not covered under the composite rate established at §413.170 and for which the laboratory files a Medicare claim directly.

(6) To furnish all other necessary dialysis services and supplies (that is, those which are not home dialysis equipment and supplies).

(7) To satisfy all documentation, recordkeeping and reporting requirements in subpart U (Conditions for Coverage of Suppliers of ESRD Services) of this chapter. This includes maintaining a complete medical record of ESRD related items and services furnished by other parties. The facility must report, on the forms required by HCFA or the ESRD network, all data for each patient in accordance with subpart U.

(iv) The facility with which the agreement is made must be located within a reasonable distance from the patient’s home (that is, located so that the facility can actually furnish the needed services in a practical and timely manner, taking into account variables like the terrain, whether the patient’s home is located in an urban or
rural area, the availability of transportation, and the usual distances traveled by patients in the area to obtain health care services).

(b) Support services—(1) Basic rule. Except as provided in paragraph (b)(2) of this section, Medicare pays for support services only under the prospective payment rates established in §413.170 of this chapter.

(2) Exceptions. If the patient elects to obtain home dialysis equipment and supplies from a supplier that is not an approved ESRD facility, Medicare pays for support services, other than support services furnished by military or VA hospitals referred to in paragraph (a)(1)(i)(B) of this section, under paragraphs (b)(2)(i) and (ii) of this section but in no case may the amount of payment exceed the limit for support services in paragraph (c)(1) of this section:

(i) For support services furnished by a hospital-based ESRD facility, Medicare pays on a reasonable cost basis in accordance with part 413 of this chapter.

(ii) For support services furnished by an independent ESRD facility, Medicare pays on the basis of reasonable charges that are related to costs and allowances that are reasonable when the services are furnished in an effective and economical manner.

(c) Payment limits—(1) Support services. The amount of payment for home dialysis support services is limited to the national average Medicare-allowed charge per patient per month for home dialysis support services, as determined by HCFA, plus the median cost per treatment for all dialysis facilities for laboratory tests included under the composite rate, as determined by HCFA, multiplied by the national average number of treatments per month.

(2) Equipment and supplies. Payment for home dialysis equipment and supplies is limited to an amount equal to the result obtained by subtracting the support services payment limit in paragraph (c)(1) of this section from the amount (or, in the case of continuous cycling peritoneal dialysis, 130 percent) of the national median payment as determined by HCFA that would have been made under the prospective payment rates established in §413.170 of this chapter for hospital-based facilities.

(3) Notification of changes to the payment limits. Updated data are incorporated into the payment limits when the prospective payment rates established at §413.170 of this chapter are updated, and changes are announced by notice in the Federal Register without a public comment period. Revisions of the methodology for determining the limits are published in the Federal Register in accordance with the Department’s established rulemaking procedures.

§ 414.335 Payment for EPO furnished to a home dialysis patient for use in the home.

(a) Payment for EPO used at home by a home dialysis patient is made only to either a Medicare approved ESRD facility or a supplier of home dialysis equipment and supplies.

(b) Payment is made in accordance with the rules set forth in §413.170 of this chapter.

Section 415.335—Payment for EPO furnished to a home dialysis patient for use in the home.

(1) Payment for EPO used at home by a home dialysis patient is made only to either a Medicare approved ESRD facility or a supplier of home dialysis equipment and supplies.

(2) Payment is made in accordance with the rules set forth in §413.170 of this chapter.

Subparts F–H—[Reserved]

PART 415—SERVICES FURNISHED BY PHYSICIANS IN PROVIDERS, SUPERVISING PHYSICIANS IN TEACHING SETTINGS, AND RESIDENTS IN CERTAIN SETTINGS

Subpart A—General Provisions

Sec. 415.1 Basis and scope.

Subpart B—Fiscal Intermediary Payments to Providers for Physician Services

415.50 Scope.
415.55 General payment rules.
415.60 Allocation of physician compensation costs.
415.70 Limits on compensation for physician services in providers.

Subpart C—Part B Carrier Payments for Physician Services to Beneficiaries in Providers

415.100 Scope.
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§ 415.1 Basis and scope.

(a) Basis. This part is based on the provisions of the following sections of the Act: Section 1848 establishes a fee schedule for payment for physician services. Section 1861(q) specifies what is included in the term “physician services” covered under Medicare. Section 1862(a)(14) sets forth the exclusion of nonphysician services furnished to hospital patients under Part B of Medicare. Section 1866(d)(5)(B) provides for a payment adjustment under the prospective payment system for the operating costs of inpatient hospital services furnished to Medicare beneficiaries in cost reporting periods beginning on or after October 1, 1983, to account for the indirect costs of medical education. Section 1886(h) establishes the methodology for Medicare payment of the cost of direct GME activities.

(b) Scope. This part sets forth rules for fiscal intermediary payments to providers for physician services, Part B carrier payments for physician services to beneficiaries in providers, physician services in teaching settings, and services of residents.

Subpart B—Fiscal Intermediary Payments to Providers for Physician Services

§ 415.50 Scope.

This subpart sets forth rules for payment by fiscal intermediaries to providers for services furnished by physicians. Payment for covered services is made either under the prospective payment system (PPS) to PPS-participating providers in accordance with part 412 of this chapter or under the reasonable cost method to non-PPS participating providers in accordance with part 413 of this chapter.

§ 415.55 General payment rules.

(a) Allowable costs. Except as specified otherwise in §§ 413.102 of this chapter (concerning compensation of owners), 415.60 (concerning allocation of physician compensation costs), and 415.162 (concerning payment for physician services furnished to beneficiaries in teaching hospitals), costs a provider incurs for services of physicians are allowable only if the following conditions are met:

(1) The services do not meet the conditions in § 415.102(a) regarding fee schedule payment for services of physicians to a beneficiary in a provider.
(2) The services include a surgeon's supervision of services of a qualified anesthetist, but do not include physician availability services, except for reasonable availability services furnished for emergency rooms and the services of standby surgical team physicians.

(3) The provider has incurred a cost for salary or other compensation it furnished the physician for the services.

(4) The costs incurred by the provider for the services meet the requirements in §413.9 of this chapter regarding costs related to patient care.

(5) The costs do not include supervision of interns and residents unless the provider elects reasonable cost payment as specified in §415.160, or any other costs incurred in connection with an approved GME program that are payable under §413.86 of this chapter.

(b) Allocation of allowable costs. The provider must follow the rules in §415.60 regarding allocation of physician compensation costs to determine its costs of services.

(c) Limits on allowable costs. The intermediary must apply the limits on compensation set forth in §415.70 to determine its payments to a provider for the costs of services.

§415.60 Allocation of physician compensation costs.

(a) Definition. For purposes of this subpart, physician compensation costs means monetary payments, fringe benefits, deferred compensation, and any other items of value (excluding office space or billing and collection services) that a provider or other organization furnishes a physician in return for the physician services. Other organizations are entities related to the provider within the meaning of §413.17 of this chapter or entities that furnish services for the provider under arrangements within the meaning of the Act.

(b) General rule. Except as provided in paragraph (d) of this section, each provider that incurs physician compensation costs must allocate those costs, in proportion to the percentage of total time that is spent in furnishing each category of services, among—

(1) Physician services to the provider (as described in §415.55); and

(2) Physician services to patients (as described in §415.102); and

(3) Activities of the physician, such as funded research, that are not paid under either Part A or Part B of Medicare.

(c) Allowable physician compensation costs. Only costs allocated to payable physician services to the provider (as described in §415.55) are allowable costs to the provider under this subpart.

(d) Allocation of all compensation to services to the provider. Generally, the total physician compensation received by a physician is allocated among all services furnished by the physician, unless—

(1) The provider certifies that the compensation is attributable solely to the physician services furnished to the provider; and

(2) The physician bills all patients for the physician services he or she furnishes to them and personally receives the payment from or on behalf of the patients. If returned directly or indirectly to the provider or an organization related to the provider within the meaning of §413.17 of this chapter, these payments are not compensation for physician services furnished to the provider.

(e) Assumed allocation of all compensation to beneficiary services. If the provider and physician agree to accept the assumed allocation of all the physician services to direct services to beneficiaries as described under §415.102(a), HCFA does not require a written allocation agreement between the physician and the provider.

(f) Determination and payment of allowable physician compensation costs. (1) Except as provided under paragraph (e) of this section, the intermediary pays the provider for these costs only if—

(i) The provider submits to the intermediary a written allocation agreement between the provider and the physician that specifies the respective amounts of time the physician spends in furnishing physician services to the provider, physician services to patients, and services that are not payable under either Part A or Part B of Medicare; and

(ii) The compensation is reasonable in terms of the time devoted to these services.
(2) In the absence of a written allocation agreement, the intermediary assumes, for purposes of determining reasonable costs of the provider, that 100 percent of the physician compensation cost is allocated to services to beneficiaries as specified in paragraph (b)(2) of this section.

(g) Recordkeeping requirements. Except for services furnished in accordance with the assumed allocation under paragraph (e) of this section, each provider that claims payment for services of physicians under this subpart must meet all of the following requirements:

(1) Maintain the time records or other information it used to allocate physician compensation in a form that permits the information to be validated by the intermediary or the carrier.

(2) Report the information on which the physician compensation allocation is based to the intermediary or the carrier on an annual basis and promptly notify the intermediary or carrier of any revisions to the compensation allocation.

(3) Retain each physician compensation allocation, and the information on which it is based, for at least 4 years after the end of each cost reporting period to which the allocation applies.

§ 415.70 Limits on compensation for physician services in providers.

(a) Principle and scope. (1) Except as provided in paragraphs (a)(2) and (a)(3) of this section, HCFA establishes reasonable compensation equivalency limits on the amount of compensation paid to physicians by providers. These limits are applied to a provider’s costs incurred in compensating physicians for services to the provider, as described in §415.55(a).

(2) Limits established under this section do not apply to costs of physician compensation attributable to furnishing inpatient hospital services that are paid for under the prospective payment system implemented under part 412 of this chapter or to costs of physician compensation attributable to approved GME programs that are payable under §413.86 of this chapter.

(3) Compensation that a physician receives for activities that may not be paid for under either Part A or Part B of Medicare is not considered in applying these limits.

(b) Methodology for establishing limits. HCFA establishes a methodology for determining annual reasonable compensation equivalency limits and, to the extent possible, considers average physician incomes by specialty and type of location using the best available data.

(c) Application of limits. If the level of compensation exceeds the limits established under paragraph (b) of this section, Medicare payment is based on the level established by the limits.

(d) Adjustment of the limits. The intermediary may adjust limits established under paragraph (b) of this section to account for costs incurred by the physician or the provider related to malpractice insurance, professional memberships, and continuing medical education.

(1) For the costs of membership in professional societies and continuing medical education, the intermediary may adjust the limit by the lesser of—

(i) The actual cost incurred by the provider or the physician for these activities; or

(ii) Five percent of the appropriate limit.

(2) For the cost of malpractice expenses incurred by either the provider or the physician, the intermediary may adjust the reasonable compensation equivalency limit by the cost of the malpractice insurance expense related to the physician service furnished to patients in providers.

(e) Exception to limits. An intermediary may grant a provider an exception to the limits established under paragraph (b) of this section only if the provider can demonstrate to the intermediary that it is unable to recruit or maintain an adequate number of physicians at a compensation level within these limits.

(f) Notification of changes in methodologies and payment limits. (1) Before the start of a cost reporting period to which limits established under this section will be applied, HCFA publishes a notice in the Federal Register that sets forth the amount of the limits and explains how it calculated the limits.

(2) If HCFA proposes to revise the methodology for establishing payment
§ 415.102 Conditions for fee schedule payment for physician services to beneficiaries in providers.

(a) General rule. If the physician furnishes services to beneficiaries in providers, the carrier pays on a fee schedule basis provided the following requirements are met:

(1) The services are personally furnished for an individual beneficiary by a physician.

(2) The services contribute directly to the diagnosis or treatment of an individual beneficiary.

(3) The services ordinarily require performance by a physician.

(4) In the case of radiology or laboratory services, the additional requirements in §415.120 or §415.130, respectively, are met.

(b) Exception. If a physician furnishes services in a provider that do not meet the requirements in paragraph (a) of this section, but are related to beneficiary care furnished by the provider, the intermediary pays for those services, if otherwise covered. The intermediary follows the rules in §§415.55 and 415.60 for payment on the basis of reasonable cost or PPS, as appropriate.

(c) Effect of billing charges for physician services to a provider.

(1) If a physician furnishes services that may be paid under the reasonable cost rules in §415.55 or §415.60, and paid by the intermediary, or would be paid under those rules except for the PPS rules in part 412 of this chapter, and under the payment rules for GME established by §413.86 of this chapter, neither the provider nor the physician may seek payment from the carrier, beneficiary, or another insurer.

(2) If a physician furnishes services to an individual beneficiary that do not meet the applicable conditions in §§415.120 (concerning conditions for payment for radiology services) and 415.130 (concerning conditions for payment for physician pathology services), the carrier does not pay on a fee schedule basis.

(3) If the physician, the provider, or another entity bills the carrier or the beneficiary or another insurer for physician services furnished to the provider, as described in §415.55(a), HCFA considers the provider to which the services are furnished to have violated its provider participation agreement, and may terminate that agreement. See part 489 of this chapter for rules governing provider agreements.

(d) Effect of physician assumption of operating costs. If a physician or other entity enters into an agreement (such as a lease or concession) with a provider, and the physician (or entity) assumes some or all of the operating costs of the provider department in which the physician furnishes physician services, the following rules apply:

(1) If the conditions set forth in paragraph (a) of this section are met, the carrier pays for the physician services under the physician fee schedule in part 414 of this chapter.
(2) To the extent the provider incurs a cost payable on a reasonable cost basis under part 413 of this chapter, the intermediary pays the provider on a reasonable cost basis for the costs associated with producing these services, including overhead, supplies, equipment costs, and services furnished by nonphysician personnel.

(3) The physician (or other entity) is treated as being related to the provider within the meaning of §413.17 of this chapter (concerning cost to related organizations).

(4) The physician (or other entity) must make its books and records available to the provider and the intermediary as necessary to verify the nature and extent of the costs of the services furnished by the physician (or other entity).

§ 415.105 Amounts of payment for physician services to beneficiaries in providers.

(a) General rule. The carrier determines amounts of payment for physician services to beneficiaries in providers in accordance with the general rules governing the physician fee schedule payment in part 414 of this chapter, except as provided in paragraph (b) of this section.

(b) Application in certain settings—(1) Teaching hospitals. The carrier applies the rules in subpart D of this part (concerning physician services in teaching settings), in addition to those in this section, in determining whether fee schedule payment should be made for physician services to individual beneficiaries in a teaching hospital.

(2) Hospital-based ESRD facilities. The carrier applies §§414.310 through 414.314 of this chapter, which set forth determination of reasonable charges under the ESRD program, to determine the amount of payment for physician services furnished to individual beneficiaries in a hospital-based ESRD facility approved under part 405 subpart U.

§ 415.110 Conditions for payment: Medically directed anesthesia services.

(a) General payment rule. Medicare pays for the physician's medical direction of anesthesia services for one service or two through four concurrent anesthesia services furnished after December 31, 1998, only if each of the services meets the condition in §415.102(a) and the following additional conditions:

(i) Performs a pre-anesthetic examination and evaluation;

(ii) Prescribes the anesthesia plan;

(iii) Personally participates in the most demanding aspects of the anesthesia plan including, if applicable, induction and emergence;

(iv) Ensures that any procedures in the anesthesia plan that he or she does not perform are performed by a qualified individual as defined in operating instructions;

(v) Monitors the course of anesthesia administration at frequent intervals;

(vi) Remains physically present and available for immediate diagnosis and treatment of emergencies; and

(vii) Provides indicated post-anesthesia care.

(2) The physician directs no more than four anesthesia services concurrently and does not perform any other services while he or she is directing the single or concurrent services so that one or more of the conditions in paragraph (a)(1) of this section are not violated.

(3) If the physician personally performs the anesthesia service, the payment rules in §414.46(c) of this chapter apply (Physician personally performs the anesthesia procedure).

(b) Medical documentation. The physician alone inclusively documents in the patient's medical record that the conditions set forth in paragraph (a)(1) of this section have been satisfied, specifically documenting that he or she performed the pre-anesthetic exam and evaluation, provided the indicated post-anesthesia care, and was present during the most demanding procedures, including induction and emergence where applicable.

[63 FR 58912, Nov. 2, 1998]

§ 415.120 Conditions for payment: Radiology services.

(a) Services to beneficiaries. The carrier pays for radiology services furnished by a physician to a beneficiary
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(2) Relate to a test result that lies outside the clinically significant normal or expected range in view of the condition of the beneficiary.

(3) Result in a written narrative report included in the beneficiary's medical record.

(4) Require the exercise of medical judgment by the consultant physician.

(c) Physician pathology services furnished by an independent laboratory. The technical component of physician pathology services furnished by an independent laboratory to a hospital inpatient before January 1, 2001, or to an outpatient are paid on a fee schedule basis under this subpart. On or after January 1, 2001, payment is made only to the hospital for the technical component of physician pathology services furnished to a hospital inpatient.

[60 FR 63178, Dec. 8, 1995, as amended at 64 FR 59442, Nov. 2, 1999]

Subpart D—Physician Services in Teaching Settings

§ 415.130 Conditions for payment: Physician pathology services.

(a) Physician pathology services. The carrier pays for pathology services furnished by a physician to an individual beneficiary on a fee schedule basis only if the services meet the conditions for payment in §415.102(a) and are one of the following services:

(1) Surgical pathology services.

(2) Specific cytopathology, hematology, and blood banking services that have been identified to require performance by a physician and are listed in program operating instructions.

(3) Clinical consultation services that meet the requirements in paragraph (b) of this section.

(4) Clinical laboratory interpretative services that meet the requirements of paragraphs (b)(1), (b)(3), and (b)(4) of this section and that are specifically listed in program operating instructions.

(b) Clinical consultation services. For purposes of this section, clinical consultation services must meet the following requirements:

(1) Be requested by the beneficiary's attending physician.
§ 415.160 Election of reasonable cost payment for direct medical and surgical services of physicians in teaching hospitals: General provisions.

(a) Scope. A teaching hospital may elect to receive payment on a reasonable cost basis for the direct medical and surgical services of its physicians in lieu of fee schedule payments that might otherwise be made for these services.

(b) Conditions. A teaching hospital may elect to receive these payments only if—

(1) The hospital notifies its intermediary in writing of the election and meets the conditions of either paragraph (b)(2) or paragraph (b)(3) of this section;

(2) All physicians who furnish services to Medicare beneficiaries in the hospital agree not to bill charges for these services; or

(3) All physicians who furnish services to Medicare beneficiaries in the hospital are employees of the hospital and, as a condition of employment, are precluded from billing for these services.

(c) Effect of election. If a teaching hospital elects to receive reasonable cost payment for physician direct medical and surgical services furnished to beneficiaries—

(1) Those services and the supervision of interns and residents furnishing care to individual beneficiaries are covered as hospital services, and

(2) The intermediary pays the hospital for those services on a reasonable cost basis under the rules in § 415.162. (Payment for other physician compensation costs related to approved GME programs is made as described in § 413.86 of this chapter.)

(d) Election declined. If the teaching hospital does not make this election, payment is made—

(1) For physician services furnished to beneficiaries on a fee schedule basis as described in part 414 subject to the rules in this subpart, and

(2) For the supervision of interns and residents as described in § 413.86.

§ 415.162 Determining payment for physician services furnished to beneficiaries in teaching hospitals.

(a) General rule. Payments for direct medical and surgical services of physicians furnished to beneficiaries and supervision of interns and residents furnishing care to beneficiaries is made by Medicare on the basis of reasonable cost if the hospital exercises the election as provided for in §415.160. If this election is made, the following occurs:

(1) Physician services furnished to beneficiaries and supervision of interns and residents furnishing care to beneficiaries are paid on a reasonable-cost basis, as provided for in paragraph (b) of this section.

(2) Payment for certain medical school costs may be made as provided for in paragraph (c) of this section.

(3) Payments for services donated by volunteer physicians to beneficiaries are made to a fund designated by the organized medical staff of the teaching hospital or medical school as provided for in paragraph (d) of this section.

(b) Reasonable cost of physician services and supervision of interns and residents.

(1) Physician services furnished to beneficiaries and supervision of interns and residents furnishing care to beneficiaries in a teaching hospital are payable as provider services on a reasonable-cost basis.

(2) For purposes of this paragraph, reasonable cost is defined as the direct salary paid to these physicians, plus applicable fringe benefits.

(3) The costs must be allocated to the services as provided by paragraph (j) of this section and apportioned to program beneficiaries as provided by paragraph (g) of this section.

(4) Other allowable costs incurred by the provider related to the services described in this paragraph are payable subject to the requirements applicable to all other provider services.

(c) Reasonable costs for the services furnished by a medical school or related organization in a hospital. An amount is payable to the hospital by HCFA under the Medicare program provided that the costs would be payable if incurred directly by the hospital rather than under the arrangement. The amount must not be in excess of the reasonable costs (as defined in paragraphs (c)(1) and (c)(2) of this section) incurred by a teaching hospital for services furnished by a medical school or organization as described in §413.17 of this chapter for certain costs to the medical school (or a related organization) in furnishing services in the hospital.

(1) Reasonable costs of physician services.

(i) When the medical school and the hospital are related organizations. If the medical school (or organization related to the medical school) and the hospital are related by common ownership or control as described in §413.17 of this chapter—

(A) The costs of these services are allowable costs to the hospital under the provisions of §413.17 of this chapter; and

(B) The reimbursable costs to the hospital are determined under the provisions of this section in the same manner as the costs incurred for physicians on the hospital staff and without regard to payments made to the medical school by the hospital.

(ii) When the medical school and the hospital are not related organizations.

(A) If the medical school and the hospital are not related organizations under the provisions of §413.17 of this chapter and the hospital makes payment to the medical school for the costs of those services furnished to all patients, payment is made by Medicare to the hospital for the reasonable cost incurred by the hospital for its payments to the medical school for services furnished to beneficiaries.

(B) Costs incurred under an arrangement must be allocated to the full range of services furnished to the hospital by the medical school physicians on the same basis as provided for under paragraph (j) of this section, and costs allocated to direct medical and surgical services furnished to hospital patients must be apportioned to beneficiaries as provided for under paragraph (g) of this section.

(C) If the medical school and the hospital are not related organizations under the provisions of §413.17 of this chapter and the hospital makes payment to the medical school only for the costs of those services furnished to beneficiaries, costs of the medical...
school not to exceed 105 percent of the sum of physician direct salaries, applicable fringe benefits, employer’s portion of FICA taxes, Federal and State unemployment taxes, and workmen’s compensation paid by the medical school or an organization related to the medical school may be recognized as allowable costs of the medical school.

(D) These allowable medical school costs must be allocated to the full range of services furnished by the physicians of the medical school or organization related as provided by paragraph (j) of this section.

(E) Costs allocated to direct medical and surgical services furnished to hospital patients must be apportioned to beneficiaries as provided by paragraph (g) of this section.

(2) Reasonable costs of other than direct medical and surgical services. These costs are determined in accordance with paragraph (c)(1) of this section except that—

(i) If the hospital makes payment to the medical school for other than direct medical and surgical services furnished to beneficiaries and supervision of interns and residents furnishing care to beneficiaries, these payments are subject to the required cost-finding and apportionment methods applicable to the cost of other hospital services (except for direct medical and surgical services furnished to beneficiaries); or

(ii) If the hospital makes payment to the medical school only for these services furnished to beneficiaries, the cost of these services is not subject to cost-finding and apportionment otherwise provided by this subpart, and the reasonable cost paid by Medicare must be determined on the basis of the health insurance ratio(s) used in the apportionment of all other provider costs (excluding physician direct medical and surgical services furnished to beneficiaries) applied to the allowable medical school costs incurred by the medical school for the services furnished to all patients of the hospital.

(d) “Salary equivalent” payments for direct medical and surgical services furnished by physicians on the voluntary staff of the hospital.

(1) HCFA makes payments under the Medicare program to a fund as defined in §415.164 for direct medical and surgical services furnished to beneficiaries on a regularly scheduled basis by physicians on the unpaid voluntary medical staff of the hospital (or medical school under arrangement with the hospital).

(i) These payments represent compensation for contributed medical staff time which, if not contributed, would have to be obtained through employed staff on a payable basis.

(ii) Payments for volunteer services are determined by applying to the regularly scheduled contributed time an hourly rate not to exceed the equivalent of the average direct salary (exclusive of fringe benefits) paid to all full-time, salaried physicians (other than interns and residents) on the hospital staff or, if the number of full-time salaried physicians is minimal in absolute terms or in relation to the number of physicians on the voluntary staff, to physicians at like institutions in the area.

(iii) This “salary equivalent” is a single hourly rate covering all physicians regardless of specialty and is applied to the actual regularly scheduled time contributed by the physicians in furnishing direct medical and surgical services to beneficiaries including supervision of interns and residents in that care.

(iv) A physician who receives any compensation from the hospital or a medical school related to the hospital by common ownership or control (within the meaning of §413.17 of this chapter) for direct medical and surgical services furnished to any patient in the hospital is not considered an unpaid voluntary physician for purposes of this paragraph.

(v) If, however, a physician receives compensation from the hospital or related medical school or organization only for services that are other than direct medical and surgical services, a salary equivalent payment for the physician’s regularly scheduled direct medical and surgical services to beneficiaries in the hospital may be imputed. However, the sum of the imputed value for volunteer services and the physician’s actual compensation from the hospital and the related medical school (or organization) may not
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exceed the amount that would have been imputed if all of the physician’s hospital and medical school services (compensated and volunteer) had been volunteer services, or paid at the rate of $30,000 per year, whichever is less.

(2) The following examples illustrate how the allowable imputed value for volunteer services is determined. In each example, it has been assumed that the average salary equivalent hourly rate is equal to the hourly rate for the individual physician’s compensated services.

Example No. 1. Dr. Jones received $3,000 a year from Hospital X for services other than direct medical services to all patients, for example, utilization review and administrative services. Dr. Jones also voluntarily furnished direct medical services to beneficiaries. The imputed value of the volunteer services amounted to $10,000 for the cost reporting period. The full imputed value of Dr. Jones’ volunteer direct medical services would be allowed since the total amount of the imputed value ($10,000) and the compensated services ($3,000) does not exceed $30,000.

Example No. 2. Dr. Smith received $25,000 from Hospital X for services as a department head in a teaching hospital. Dr. Smith also voluntarily furnished direct medical services to beneficiaries. The imputed value of the volunteer services amounted to $10,000. Only $5,000 of the imputed value of volunteer services would be allowed since the total amount of the imputed value ($10,000) and the compensated services ($25,000) exceeds the $30,000 maximum amount allowable for all of Dr. Smith’s services.

(3) The amount of the imputed value for volunteer services applicable to beneficiaries and payable to a fund is determined in accordance with the aggregate per diem method described in paragraph (g) of this section.

(4) Medicare payments to a fund must be used by the fund solely for improvement of care of hospital patients or for educational or charitable purposes (which may include but are not limited to medical and other scientific research).

(i) No personal financial gain, either direct or indirect, from benefits of the fund may inure to any of the hospital staff physicians, medical school faculty, or physicians for whom Medicare imputes costs for purposes of payment into the fund.

(ii) Expenses met from contributions made to the hospital from a fund are not included as a reimbursable cost when expended by the hospital, and depreciation expense is not allowed with respect to equipment or facilities donated to the hospital by a fund or purchased by the hospital from monies in a fund.

(e) Requirements for payment—(1) Physicians on the hospital staff. The requirements under which the costs of physician direct medical and surgical services (including supervision of interns and residents) to beneficiaries are the same as those applicable to the cost of all other covered provider services except that the costs of these services are separately determined as provided by this section and are not subject to cost-finding as described in § 413.24 of this chapter.

(2) Physicians on the medical school faculty. Payment is made to a hospital for the costs of services of physicians on the medical school faculty, provided that if the medical school is not related to the hospital (within the meaning of § 413.17 of this chapter, concerning cost to related organizations), the hospital does not make payment to the medical school for services furnished to all patients and the following requirements are met: If the hospital makes payment to the medical school for services furnished to all patients,
these requirements do not apply. (See paragraph (c)(1)(ii) of this section.)

(i) There is a written agreement between the hospital and the medical school or organization, specifying the types and extent of services to be furnished by the medical school and specifying that the hospital must pay to the medical school an amount at least equal to the reasonable cost (as defined in paragraph (c) of this section) of furnishing the services to beneficiaries.

(ii) The costs are paid to the medical school by the hospital no later than the date on which the cost report covering the period in which the services were furnished is due to HCFA.

(iii) Payment for the services furnished under an arrangement would have been made to the hospital had the services been furnished directly by the hospital.

(3) Physicians on the voluntary staff of the hospital (or medical school under arrangement with the hospital). If the conditions for payment to a fund outlined in §415.164 are met, payments are made on a "salary equivalent" basis (as defined in paragraph (d) of this section) to a fund.

(f) Requirements for payment for medical school faculty services other than physician direct medical and surgical services. If the requirements for payment for physician direct medical and surgical services furnished to beneficiaries in a teaching hospital is determined on the basis of an average per diem, as defined in paragraph (h)(1) of this section, for physician direct medical and surgical services to all patients except that the average per diem is derived from the imputed value of the physician direct medical and surgical services furnished to all patients.

(h) Definitions.

(1) Average cost per diem for physician direct medical and surgical services (including supervision of interns and residents) furnished in a teaching hospital to patients in each category of physician services described in paragraph (g)(1) of this section means the amount computed by dividing total reasonable costs of these services in each category by the sum of—

(i) Inpatient days (as defined in paragraph (h)(2) of this section); and

(ii) Outpatient visit days (as defined in paragraph (h)(3) of this section).

(2) Inpatient days are determined by counting the day of admission as 3.5 days and each day after a patient's day of admission, except the day of discharge, as 1 day.

(3) Outpatient visit days are determined by counting only one visit day for each calendar day that a patient visits an outpatient department or multiple outpatient departments.

(i) Application. (1) The following illustrates how apportionment based on the aggregate per diem method for costs of physician direct medical and surgical services furnished in a teaching hospital to patients is determined.

TEACHING HOSPITAL Y

Statistical and financial data:

| Total inpatient days as defined in paragraph (h)(2) of this section and outpatient visit days as defined in paragraph (h)(3) of this section | 75,000 |
| Total inpatient Part A days | 20,000 |
| Total inpatient Part B days where Part A coverage is not available | 1,000 |
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Total outpatient Part B visit days ......................... 5,000
Total cost of direct medical and surgical services furnished to all patients by physicians on the medical school faculty as determined in accordance with paragraph (i) of this section .................................. $1,650,000
Computation of cost applicable to program for physicians on the medical school faculty:
Average cost per diem for direct medical and surgical services to patients by physicians on the medical school faculty: $1,650,000 ÷ 75,000 = $22 per diem.
Cost of physician direct medical and surgical services furnished to inpatient beneficiaries covered under Part B: $22 per diem × 1,000 ...................... $22,000
Cost of physician direct medical and surgical services furnished to outpatient beneficiaries covered under Part B: $22 per diem × 5,000 ...................... $110,000

(2) The following illustrates how the imputed value of physician volunteer direct medical and surgical services furnished in a teaching hospital to beneficiaries is determined.
Example: The physicians on the medical staff of Teaching Hospital Y donated a total of 5,000 hours in furnishing direct medical and surgical services to patients of the hospital during a cost reporting period and did not receive any compensation from either the hospital or the medical school. Also, the imputed value for any physician volunteer services did not exceed the rate of $30,000 per year per physician.

Statistical and Financial Data:
Total salaries paid to the full-time salaried physicians by the hospital (excluding interns and residents) ....................... $800,000
Total physicians who were paid for an average of 40 hours per week or 2,080 (52 weeks x 40 hours per week) hours per year ........ 20
Average hourly rate equivalent: $800,000 ÷ 41,600 (2,080 × 20) ........................................ $19.23
Computation of total imputed value of physician volunteer services applicable to all patients:
(Total donated hours × average hourly rate equivalent): 5,000 × $19.23 ...................... $96,150
Total inpatient days (as defined in paragraph (h)(2) of this section) and outpatient visit days (as defined in paragraph (h)(3) of this section) ...................... 75,000
Total inpatient Part A days 20,000
Total inpatient Part B days if Part A coverage is not available ......................... 1,000
Total outpatient Part B visit days ......................... 5,000
Computation of imputed value of physician volunteer direct medical and surgical services furnished to Medicare beneficiaries:

Average per diem for physician direct medical and surgical services to all patients:

$96,150 \div 75,000 = $1.28 per diem

Imputed value of physician direct medical and surgical services furnished to inpatient beneficiaries covered under Part A:

$1.28 per diem × 20,000 = $25,600

Imputed value of physician direct medical and surgical services furnished to inpatient beneficiaries covered under Part B:

$1.28 per diem × 1,000 = $1,280

Imputed value of physician direct medical and surgical services furnished to outpatient beneficiaries covered under Part B:

$1.28 per diem × 3,000 = $6,400

Total ................................... $33,280

(j) Allocation of compensation paid to physicians in a teaching hospital.

(1) In determining reasonable cost under this section, the compensation paid by a teaching hospital, or a medical school or related organization under arrangement with the hospital, to physicians in a teaching hospital must be allocated to the full range of services implicit in the physician compensation arrangements. (However, see paragraph (d) of this section for the computation of the "salary equivalent" payments for volunteer services furnished to patients.)

(2) This allocation must be made and must be capable of substantiation on the basis of the proportion of each physician's time spent in furnishing each type of service to the hospital or medical school.

§415.164 Payment to a fund.

(a) General rules. Payment for certain voluntary services by physicians in teaching hospitals (as these services are described in §415.160) is made on a salary equivalent basis (as described in §415.162(d)) subject to the conditions and limitations contained in parts 405 and 413 of this chapter and this part 415, to a single fund (as defined in paragraph (b) of this section) designated by the organized medical staff of the hospital (or, if the services are furnished in the hospital by the faculty of a medical school, to a fund as may be designated by the faculty), if the following conditions are met:

(1) The hospital (or medical school furnishing the services under arrangement with the hospital) incurs no actual cost in furnishing the services.

(2) The hospital has an agreement with HCFA under part 489 of this chapter.

(3) The intermediary, or HCFA as appropriate, has received written assurances that—

(i) The payment is used solely for the improvement of care of hospital patients or for educational or charitable purposes; and

(ii) Neither the individuals who are furnished the services nor any other persons are charged for the services (and if charged, provision is made for the return of any monies incorrectly collected).

(b) Definition of a fund. For purposes of paragraph (a) of this section, a fund is an organization that meets either of the following requirements:

(1) The organization has and retains exemption, as a governmental entity or under section 501(c)(3) of the Internal Revenue Code (nonprofit educational, charitable, and similar organizations), from Federal taxation.

(2) The organization is an organization of physicians who, under the terms of their employment by an entity that meets the requirements of paragraph (b)(1) of this section, are required to turn over to that entity all income that the physician organization derives from the physician services.

(c) Status of a fund. A fund approved for payment under paragraph (a) of this section has all the rights and responsibilities of a provider under Medicare except that it does not enter into an agreement with HCFA under part 489 of this chapter.

§415.170 Conditions for payment on a fee schedule basis for physician services in a teaching setting.

Services meeting the conditions for payment in §415.102(a) furnished in teaching settings are payable under the physician fee schedule if—
(a) The services are personally furnished by a physician who is not a resident; or
(b) The services are furnished by a resident in the presence of a teaching physician except as provided in §415.172 (concerning physician fee schedule payment for services of teaching physicians), §415.174 (concerning an exception for services furnished in hospital outpatient and certain other ambulatory settings), §415.176 (concerning renal dialysis services), and §415.184 (concerning psychiatric services), as applicable.

§ 415.172 Physician fee schedule payment for services of teaching physicians.

(a) General rule. If a resident participates in a service furnished in a teaching setting, physician fee schedule payment is made only if a teaching physician is present during the key portion of any service or procedure for which payment is sought.

(1) In the case of surgical, high-risk, or other complex procedures, the teaching physician must be present during all critical portions of the procedure and immediately available to furnish services during the entire service or procedure.

(i) In the case of surgery, the teaching physician's presence is not required during opening and closing of the surgical field.

(ii) In the case of procedures performed through an endoscope, the teaching physician must be present during the entire viewing.

(2) In the case of evaluation and management services, the teaching physician must be present during the portion of the service that determines the level of service billed. (However, in the case of evaluation and management services furnished in hospital outpatient departments and certain other ambulatory settings, the requirements of §415.174 apply.)

(b) Documentation. Except for services furnished as set forth in §§415.174 (concerning an exception for services furnished in hospital outpatient and certain other ambulatory settings), 415.176 (concerning renal dialysis services), and 415.184 (concerning psychiatric services), the medical records must document the teaching physician was present at the time the service is furnished. The presence of the teaching physician during procedures may be demonstrated by the notes in the medical records made by a physician, resident, or nurse. In the case of evaluation and management procedures, the teaching physician must personally document his or her participation in the service in the medical records.

(c) Payment level. In the case of services such as evaluation and management for which there are several levels of service codes available for reporting purposes, the appropriate payment level must reflect the extent and complexity of the service when fully furnished by the teaching physician.

§ 415.174 Exception: Evaluation and management services furnished in certain centers.

(a) In the case of certain evaluation and management codes of lower and mid-level complexity (as specified by HCFA in program instructions), carriers may make physician fee schedule payment for a service furnished by a resident without the presence of a teaching physician. For the exception to apply, all of the following conditions must be met:

(1) The services must be furnished in a center that is located in an outpatient department of a hospital or another ambulatory care entity in which the time spent by residents in patient care activities is included in determining intermediary payments to a hospital under §413.86.

(2) Any resident furnishing the service without the presence of a teaching physician must have completed more than 6 months of an approved residency program.

(3) The teaching physician must not direct the care of more than four residents at any given time and must direct the care from such proximity as to constitute immediate availability. The teaching physician must—

(i) Have no other responsibilities at the time;

(ii) Assume management responsibility for those beneficiaries seen by the residents;

(iii) Ensure that the services furnished are appropriate;
§ 415.176 Renal dialysis services.

(iii) Review with each resident during or immediately after each visit, the beneficiary’s medical history, physical examination, diagnosis, and record of tests and therapies; and

(v) Document the extent of the teaching physician’s participation in the review and direction of the services furnished to each beneficiary.

(4) The range of services furnished by residents in the center includes all of the following:

(i) Acute care for undifferentiated problems or chronic care for ongoing conditions.

(ii) Coordination of care furnished by other physicians and providers.

(iii) Comprehensive care not limited by organ system, or diagnosis.

(5) The patients seen must be an identifiable group of individuals who consider the center to be the continuing source of their health care and in which services are furnished by residents under the medical direction of teaching physicians.

(b) Nothing in paragraph (a) of this section may be construed as providing a basis for the coverage of services not determined to be covered under Medicare, such as routine physical checkups.

[60 FR 63178, Dec. 8, 1995; 61 FR 59554, Nov. 22, 1996]

§ 415.178 Anesthesia services.

(a) General rule. An unreduced physician fee schedule payment may be made if a physician is involved in a single anesthesia procedure involving an anesthesia resident. In the case of anesthesia services, the teaching physician must be present during all critical portions of the procedure and immediately available to furnish services during the entire service or procedure. The teaching physician cannot receive an unreduced fee if he or she performs services involving other patients during the period the anesthesia resident is furnishing services in a single case.

(b) Documentation. Documentation must indicate the physician’s presence or participation in the administration of the anesthesia.

[60 FR 63178, Dec. 8, 1995; 61 FR 42385, Aug. 15, 1996]

§ 415.180 Teaching setting requirements for the interpretation of diagnostic radiology and other diagnostic tests.

(a) General rule. Physician fee schedule payment is made for the interpretation of diagnostic radiology and other diagnostic tests if the interpretation is performed or reviewed by a physician other than a resident.

(b) Documentation. Documentation must indicate that the physician personally performed the interpretation or reviewed the resident’s interpretation with the resident.

§ 415.184 Psychiatric services.

To qualify for physician fee schedule payment for psychiatric services furnished under an approved GME program, the physician must meet the requirements of §§ 415.170 and 415.172, including documentation, except that the requirement for the presence of the teaching physician during the service in which a resident is involved may be met by observation of the service by use of a one-way mirror, video equipment, or similar device.

§ 415.190 Conditions of payment: Assistants at surgery in teaching hospitals.

(a) Basis, purpose, and scope. This section describes the conditions under which Medicare pays on a fee schedule basis for the services of an assistant at surgery in a teaching hospital. This section is based on section 1842(b)(7)(D)(I) of the Act and applies only to hospitals with an approved GME residency program. Except as specified in paragraph (c) of this section, fee schedule payment is not available for assistants at surgery in hospitals with—
(1) A training program relating to the medical specialty required for the surgical procedure; and
(2) A resident in a training program relating to the specialty required for the surgery available to serve as an assistant at surgery.

(b) Definition. Assistant at surgery means a physician who actively assists the physician in charge of a case in performing a surgical procedure.

(c) Conditions for payment for assistants at surgery. Payment on a fee schedule basis is made for the services of an assistant at surgery in a teaching hospital only if the services meet one of the following conditions:

(1) Are required as a result of exceptional medical circumstances.
(2) Are complex medical procedures performed by a team of physicians, each performing a discrete, unique function integral to the performance of a complex medical procedure that requires the special skills of more than one physician.
(3) Constitute concurrent medical care relating to a medical condition that requires the presence of, and active care by, a physician of another specialty during surgery.
(4) Are medically required and are furnished by a physician who is primarily engaged in the field of surgery, and the primary surgeon does not use interns and residents in the surgical procedures that the surgeon performs (including preoperative and postoperative care).
(5) Are not related to a surgical procedure for which HCFA determines that assistants are used less than 5 percent of the time.

Subpart E—Services of Residents

§ 415.204 Services of residents in approved GME programs.

(a) General rules. Services furnished in hospitals by residents in approved GME programs are specifically excluded from being paid as “physician services” defined in §414.2 of this chapter and are payable as hospital services. This exclusion applies whether or not the resident is licensed to practice under the laws of the State in which he or she performs the service. The payment methodology for services of residents in hospitals and hospital-based providers is set forth in §413.86 of this chapter.

(b) Exception. For low and mid-level evaluation and management services furnished under certain conditions in centers located in hospital outpatient departments and other ambulatory settings, see §415.174.

(c) Definitions. See §415.152 for definitions of terms used in this subpart E.

§ 415.202 Services of residents not in approved GME programs.

(a) General rules. For services of a physician employed by a hospital who is authorized to practice only in a hospital setting and for the services of a resident who is not in any approved GME program, payment is made to the hospital on a Part B reasonable cost basis regardless of whether the services are furnished to hospital inpatients or outpatients.

(b) Payment. For services described in paragraph (a) of this section, payment is made under Part B by reducing the reasonable costs of furnishing the services by the beneficiary deductible and paying 80 percent of the remaining amount. No payment is made for other costs of unapproved programs, such as administrative costs related to teaching activities of physicians.

§ 415.204 Services of residents in skilled nursing facilities and home health agencies.

(a) Medicare Part A payment. Payment is made under Medicare Part A for interns’ and residents’ services furnished in the following settings that meet the specified requirements:

(1) Skilled nursing facility. Payment to a participating skilled nursing facility may include the cost of services of an intern or resident who is in an approved GME program in a hospital with which the skilled nursing facility has a transfer agreement that provides, in part, for the transfer of patients and the interchange of medical records.
(2) Home health agency. A participating home health agency may receive payment for the cost of the services of an intern or resident who is under an approved GME program of a hospital with which the home health agency is affiliated or under common
§ 415.206 Services of residents in non-provider settings.

These activities are furnished as part of the services if the beneficiary is a Medicare beneficiary. (Nevertheless, see § 413.86 of this chapter for the costs of approved GME programs in hospital-based providers.)

(b) Medicare Part B payment. Medical services of a resident of a hospital that are furnished by a skilled nursing facility or home health agency are paid under Medicare Part B if payment is not provided under Medicare Part A. Payment is made under Part B for a resident's services by reducing the reasonable costs of furnishing the services by the beneficiary deductible and paying 80 percent of the remaining amount.

§ 415.208 Services of moonlighting residents.

(a) Definition. For purposes of this section, the term services of moonlighting residents refers to services that licensed residents perform that are outside the scope of an approved GME program.

(b) Services in GME program hospitals.

(1) The services of residents to inpatients of hospitals in which the residents have their approved GME program are not covered as physician services and are payable under § 413.86 regarding direct GME payments.

(2) Services of residents that are not related to their approved GME programs and are performed in an outpatient department or emergency department of a hospital in which they have their training program are covered as physician services and payable under the physician fee schedule if any of the following criteria are met:

(i) The services are identifiable physician services and meet the conditions for payment of physician services to beneficiaries in providers in § 415.102(a).

(ii) The resident is fully licensed to practice medicine, osteopathy, dentistry, or podiatry by the State in which the services are performed.

(iii) The services performed can be separately identified from those services that are required as part of the approved program.

(3) If the criteria specified in paragraph (b)(2) of this section are met, the services of the moonlighting resident are considered to have been furnished by the individual in his or her capacity as a physician, rather than in the capacity of a resident. The carrier must review the contracts and agreements for these services to ensure compliance with the criteria specified in paragraph (b)(2) of this section.
(4) No payment is made for services of a "teaching physician" associated with moonlighting services, and the time spent furnishing these services is not included in the teaching hospital’s full-time equivalency count for the indirect GME payment (§412.105 of this chapter) and for the direct GME payment (§413.86 of this chapter).

(c) Other settings. Moonlighting services of a licensed resident in an approved GME program furnished outside the scope of that program in a hospital or other setting that does not participate in the approved GME program are payable under the physician fee schedule as set forth in §415.206(b)(1).

PART 416—AMBULATORY SURGICAL SERVICES

Subpart A—General Provisions and Definitions

§ 416.1 Basis and scope.

(a) Statutory basis. (1) Section 1832(a)(2)(F)(i) of the Act provides for Medicare Part B coverage of facility services furnished in connection with surgical procedures specified by the Secretary under section 1833(i)(1) of the Act.

(2) Section 1833(i)(1)(A) of the Act requires the Secretary to specify the surgical procedures that can be performed safely on an ambulatory basis in an ambulatory surgical center, or a hospital outpatient department.

(3) Section 1833(i)(1)(B) and (3) specify the amounts to be paid for facility services furnished in connection with the specified surgical procedures when they are performed, respectively, in an ASC, or in a hospital outpatient department.

(b) Scope. This part sets forth—

(1) The conditions that an ASC must meet in order to participate in the Medicare program;

(2) The scope of covered services; and
§ 416.2 Definitions.

As used in this part:

(3) The conditions for Medicare payment for facility services.


§ 416.25 Basic requirements.

Participation as an ASC is limited to facilities that—

(a) Meet the definition in §416.2; and

(b) Have in effect an agreement obtained in accordance with this subpart.

[56 FR 8843, Mar. 1, 1991]

§ 416.26 Qualifying for an agreement.

(a) Deemed compliance. HCFA may deem an ASC to be in compliance with any or all of the conditions set forth in subpart C of this part if—

1 The ASC authorizes the release to HCFA, of the findings of the accreditation survey.

(b) Survey of ASCs. (1) Unless HCFA deems the ASC to be in compliance with the conditions set forth in subpart C of this part, the State survey agency must survey the facility to ascertain compliance with those conditions, and report its findings to HCFA.

(2) HCFA surveys deemed ASCs on a sample basis as part of HCFA’s validation process.

(c) Acceptance of the ASC as qualified to furnish ambulatory surgical services. If HCFA determines, after reviewing the survey agency recommendation and other evidence relating to the qualification of the ASC, that the facility meets the requirements of this part, it sends to the ASC—

1 Written notice of the determination; and

(2) Two copies of the ASC agreement.

(d) Filing of agreement by the ASC. If the ASC wishes to participate in the program, it must—

1 Have both copies of the ASC agreement signed by its authorized representative; and

(2) File them with HCFA.

(e) Acceptance by HCFA. If HCFA accepts the agreement filed by the ASC, it returns the ASC one copy of the agreement, with a notice of acceptance specifying the effective date.

(f) Appeal rights. If HCFA refuses to enter into an agreement or if HCFA terminates an agreement, the ASC is entitled to a hearing in accordance with part 498 of this chapter.

[56 FR 8843, Mar. 1, 1991]

§ 416.30 Terms of agreement with HCFA.

As part of the agreement under §416.26 the ASC must agree to the following:

(a) Compliance with coverage conditions. The ASC agrees to meet the conditions for coverage specified in subpart C of this part and to report promptly to HCFA any failure to do so.

(b) Limitation on charges to beneficiaries.1 The ASC agrees to charge the

1 For facility services furnished before July 1987, the ASC had to agree to make no charge to the beneficiary, since those services were
§ 416.35 Termination of agreement.

(a) Termination by the ASC—(1) Notice to HCFA. An ASC that wishes to terminate its agreement must send HCFA written notice of its intent.

(2) Date of termination. The notice may state the intended date of termination which must be the first day of a calendar month.

(i) If the notice does not specify a date, or the date is not acceptable to HCFA, HCFA may set a date that will not be more than 6 months from the date on the ASC's notice of intent.

(ii) HCFA may accept a termination date that is less than 6 months after the date on the ASC's notice if it determines that to do so would not unduly disrupt services to the community or otherwise interfere with the effective and efficient administration of the Medicare program.

(3) Voluntary termination. If an ASC ceases to furnish services to the community, that shall be deemed to be a voluntary termination of the agreement by the ASC, effective on the last day of business with Medicare beneficiaries.

(b) Termination by HCFA—(1) Cause for termination. HCFA may terminate an agreement if it determines that the ASC—

(i) No longer meets the conditions for coverage as specified under § 416.26; or

(ii) Is not in substantial compliance with the provisions of the agreement, the requirements of this subpart, and other applicable regulations of subchapter B of this chapter, or any applicable provisions of title XVIII of the Act.

(2) Notice of termination. HCFA sends notice of termination to the ASC at
§ 416.40 Condition for coverage—Compliance with State licensure law.

The ASC must comply with State licensure requirements.

§ 416.41 Condition for coverage—Governing body and management.

The ASC must have a governing body, that assumes full legal responsibility for determining, implementing, and monitoring policies governing the ASC’s total operation and for ensuring that these policies are administered so as to provide quality health care in a safe environment. When services are provided through a contract with an outside resource, the ASC must assure that these services are provided in a safe and effective manner. Standard: Hospitalization. The ASC must have an effective procedure for the immediate transfer to a hospital, of patients requiring emergency medical care beyond the capabilities of the ASC. This hospital must be a local, Medicare participating hospital or a local, non-participating hospital that meets the requirements for payment for emergency services under §482.2 of this chapter. The ASC must have a written transfer agreement with such a hospital, or all physicians performing surgery in the ASC must have admitting privileges at such a hospital.

§ 416.42 Condition for coverage—Surgical services.

Surgical procedures must be performed in a safe manner by qualified physicians who have been granted clinical privileges by the governing body of the ASC in accordance with approved policies and procedures of the ASC.

(a) Standard: Anesthetic risk and evaluation. A physician must examine the patient immediately before surgery to evaluate the risk of anesthesia and of the procedure to be performed. Before discharge from the ASC, each patient must be evaluated by a physician for proper anesthesia recovery.

(b) Standard: Administration of anesthesia. Anesthetics must be administered by only—

(1) A qualified anesthetist; or

(2) A physician qualified to administer anesthesia, a certified registered nurse anesthetist or anesthesiologist’s assistant as defined in §410.68(b) of this chapter, or a supervised trainee in an approved educational program. In those cases in which a non-physician administers the anesthesia, the anesthetist must be under the supervision of the operating physician, and in the case of an anesthesiologist’s assistant, under the supervision of an anesthesiologist.

(c) Standard: Discharge. All patients are discharged in the company of a responsible adult, except those exempted by the attending physician.

§ 416.43 Condition for coverage—Evaluation of quality.

The ASC, with the active participation of the medical staff, must conduct
§ 416.44 Condition for coverage—Environment.

The ASC must have a safe and sanitary environment, properly constructed, equipped, and maintained to protect the health and safety of patients.

(a) Standard: Physical environment. The ASC must provide a functional and sanitary environment for the provision of surgical services.

(1) Each operating room must be designed and equipped so that the types of surgery conducted can be performed in a manner that protects the lives and assures the physical safety of all individuals in the area.

(2) The ASC must have a separate recovery room and waiting area.

(3) The ASC must establish a program for identifying and preventing infections, maintaining a sanitary environment, and reporting the results to appropriate authorities.

(b) Standard: Safety from fire. (1) Except as provided in paragraphs (b) (2) and (3) of this section, the ASC must meet the provisions of the 1985 edition of the Life Safety Code of the National Fire Protection Association (which is incorporated by reference)¹ that are applicable to ambulatory surgical centers.

(2) In consideration of a recommendation by the State survey agency, HCFA may waive, for periods deemed appropriate, specific provisions of the Life Safety Code which, if rigidly applied, would result in unreasonable hardship upon an ASC, but only if the waiver will not adversely affect the health and safety of the patients.

(3) Any ASC that, on May 9, 1988, complies with the requirements of the 1981 edition of the Life Safety Code, with or without waivers, will be considered to be in compliance with this standard, so long as the ASC continues to remain in compliance with that edition of the Life Safety Code.

(c) Standard: Emergency equipment. Emergency equipment available to the operating rooms must include at least the following:

(1) Emergency call system.

(2) Oxygen.

(3) Mechanical ventilatory assistance equipment including airways, manual breathing bag, and ventilator.

(4) Cardiac defibrillator.

(5) Cardiac monitoring equipment.

(6) Tracheostomy set.

(7) Laryngoscopes and endotracheal tubes.

(8) Suction equipment.

(9) Emergency medical equipment and supplies specified by the medical staff.

(d) Standard: Emergency personnel. Personnel trained in the use of emergency equipment and in cardiopulmonary resuscitation must be available whenever there is a patient in the ASC.


§ 416.45 Condition for coverage—Medical staff.

The medical staff of the ASC must be accountable to the governing body.

(a) Standard: Membership and clinical privileges. Members of the medical staff must be legally and professionally qualified for the positions to which they are appointed and for the performance of privileges granted. The ASC grants privileges in accordance with recommendations from qualified medical personnel.

(b) Standard: Reappraisals. Medical staff privileges must be periodically reappraised by the ASC. The scope of procedures performed in the ASC must be periodically reviewed and amended as appropriate.

(c) Standard: Other practitioners. If the ASC assigns patient care responsibilities to practitioners other than physicians, it must have established policies and procedures, approved by the governing body, for overseeing and evaluating their clinical activities.

¹See footnote to §405.1134(a) of this chapter.
§ 416.46 Condition for coverage—Nursing services.

The nursing services of the ASC must be directed and staffed to assure that the nursing needs of all patients are met.

(a) Standard: Organization and staffing. Patient care responsibilities must be delineated for all nursing service personnel. Nursing services must be provided in accordance with recognized standards of practice. There must be a registered nurse available for emergency treatment whenever there is a patient in the ASC.

(b) [Reserved]

§ 416.47 Condition for coverage—Medical records.

The ASC must maintain complete, comprehensive, and accurate medical records to ensure adequate patient care.

(a) Standard: Organization. The ASC must develop and maintain a system for the proper collection, storage, and use of patient records.

(b) Standard: Form and content of record. The ASC must maintain a medical record for each patient. Every record must be accurate, legible, and promptly completed. Medical records must include at least the following:

(1) Patient identification.
(2) Significant medical history and results of physical examination.
(3) Pre-operative diagnostic studies (entered before surgery), if performed.
(4) Findings and techniques of the operation, including a pathologist's report on all tissues removed during surgery, except those exempted by the governing body.
(5) Any allergies and abnormal drug reactions.
(6) Entries related to anesthesia administration.
(7) Documentation of properly executed informed patient consent.
(8) Discharge diagnosis.

§ 416.48 Condition for coverage—Pharmaceutical services.

The ASC must provide drugs and biologicals in a safe and effective manner, in accordance with accepted professional practice, and under the direction of an individual designated responsible for pharmaceutical services.

(a) Standard: Administration of drugs. Drugs must be prepared and administered according to established policies and acceptable standards of practice.

(1) Adverse reactions must be reported to the physician responsible for the patient and must be documented in the record.

(2) Blood and blood products must be administered by only physicians or registered nurses.

(3) Orders given orally for drugs and biologicals must be followed by a written order, signed by the prescribing physician.

(b) [Reserved]

§ 416.49 Condition for coverage—Laboratory and radiologic services.

If the ASC performs laboratory services, it must meet the requirements of part 493 of this chapter. If the ASC does not provide its own laboratory services, it must have procedures for obtaining routine and emergency laboratory services from a certified laboratory in accordance with part 493 of this chapter. The referral laboratory must be certified in the appropriate specialties and subspecialties of service to perform the referred tests in accordance with the requirements of part 493 of this chapter. The ASC must have procedures for obtaining radiologic services from a Medicare approved facility to meet the needs of patients.

[57 FR 7135, Feb. 28, 1992]

Subpart D—Scope of Benefits

§ 416.60 General rules.

(a) The services payable under this part are facility services furnished to Medicare beneficiaries, by a participating facility, in connection with covered surgical procedures specified in §416.65.

(b) The surgical procedures, including all preoperative and post-operative services that are performed by a physician, are covered as physician services under part 410 of this chapter.

[56 FR 8844, Mar. 1, 1991]

§ 416.61 Scope of facility services.

(a) Included services. Facility services include, but are not limited to—
§ 416.120 Basis for payment.

The basis for payment depends on where the services are furnished.

(a) Hospital outpatient department. Payment is in accordance with part 413 of this chapter.

(b) [Reserved]

(c) ASC—(1) General rule. Payment is based on a prospectively determined rate. This rate covers the cost of services such as supplies, nursing services, equipment, etc., as specified in §416.61. The rate does not cover physician services or other medical services covered under part 410 of this chapter (for example, X-ray services or laboratory services) which are not directly related...
§ 416.125 ASC facility services payment rate.

(a) The payment rate is based on a prospectively determined standard overhead amount per procedure derived from an estimate of the costs incurred by ambulatory surgical centers generally in providing services furnished in connection with the performance of that procedure.

(b) The payment must be substantially less than would have been paid under the program if the procedure had been performed on an inpatient basis in a hospital.


§ 416.130 Publication of revised payment methodologies.

Whenever HCFA proposes to revise the payment rate for ASCs, HCFA publishes a notice in the Federal Register describing the revision. The notice also explains the basis on which the rates were established. After reviewing public comments, HCFA publishes a notice establishing the rates authorized by this section. In setting these rates, HCFA may adopt reasonable classifications of facilities and may establish different rates for different types of surgical procedures.


§ 416.140 Surveys.

(a) Timing, purpose, and procedures. (1) No more often than once a year, HCFA conducts a survey of a randomly selected sample of participating ASCs to collect data for analysis or reevaluation of payment rates.

(2) HCFA notifies the selected ASCs by mail of their selection and of the form and content of the report the ASCs are required to submit within 60 days of the notice.

(3) If the facility does not submit an adequate report in response to HCFA’s survey request, HCFA may terminate the agreement to participate in the Medicare program as an ASC.

(4) HCFA may grant a 30-day postponement of the due date for the survey report if it determines that the facility has demonstrated good cause for the delay.

(b) Requirements for ASCs. ASCs must—

(1) Maintain adequate financial records, in the form and containing the data required by HCFA, to allow determination of the payment rates for covered surgical procedures furnished to Medicare beneficiaries under this subpart.

(2) Within 60 days of a request from HCFA submit, in the form and detail as may be required by HCFA, a report of—

(i) Their operations, including the allowable costs actually incurred for the period and the actual number and kinds of surgical procedures furnished during the period; and

(ii) Their customary charges for each surgical procedure furnished for the period.


§ 416.150 Beneficiary appeals.

A beneficiary (or ASC as his or her assignee) may request a hearing by a carrier (subject to the limitations and conditions set forth in part 405, subpart H of this chapter) if the beneficiary or the ASC—
(a) Is dissatisfied with a carrier’s denial of a request for payment made on his or her behalf by an ASC;
(b) Is dissatisfied with the amount of payment; or
(c) Believes the request for payment is not being acted upon with reasonable promptness.

Subpart F—Adjustment in Payment Amounts for New Technology Intraocular Lenses Furnished by Ambulatory Surgical Centers

SOURCE: 64 FR 32205, June 16, 1999, unless otherwise noted.

§ 416.180 Definitions.
As used in this subpart, the following definitions apply:
Class of new technology intraocular lenses (IOLs) means all of the IOLs, collectively, that HCFA determines meet the definition of “new technology IOL” under the provisions of this subpart.
Interested party means any individual, partnership, corporation, association, society, scientific or academic establishment, professional or trade organization, or any other legal entity.
New technology IOL means an IOL that HCFA determines has been approved by the FDA for use in labeling and advertising the IOL’s claims of specific clinical advantages and superiority over existing IOLs with regard to reduced risk of intraoperative or postoperative complication or trauma, accelerated postoperative recovery, reduced induced astigmatism, improved postoperative visual acuity, more stable postoperative vision, or other comparable clinical advantages.
New technology subset means a group of IOLs that HCFA determines meet the criterion for being treated as new technology IOLs and that share a common feature or features that distinguish them from other IOLs. For example, all new technology IOLs that are made of a particular bioengineered material could comprise one subset, while all that rely on a particular optical innovation could comprise another.

§ 416.185 Payment review process.
(a) HCFA publishes a FEDERAL REGISTER notice announcing the deadline and requirements for submitting a request for HCFA to review payment for an IOL.
(b) HCFA receives a request to review the appropriateness of the payment amount for an IOL.
(c) HCFA compiles a list of the requests it receives and identifies the IOL manufacturer’s name, the model number of the IOL to be reviewed, the interested party or parties that submit requests, and a summary of the interested party’s grounds for requesting review of the appropriateness of the IOL payment amount.
(d) HCFA publishes the list of requests in a FEDERAL REGISTER notice with comment period, giving the public 30 days to comment on the IOLs for which review was requested.
(e) HCFA reviews the information submitted with the request to review, any timely public comments that are submitted regarding the list of IOLs published in the FEDERAL REGISTER, and any other timely information that HCFA deems relevant to decide whether to provide a payment adjustment as specified in §416.200. HCFA makes a determination of whether the IOL meets the definition of a new technology IOL in §416.180.
(f) If HCFA determines that a lens is a new technology IOL, HCFA establishes a payment adjustment as follows:
(1) Before July 16, 2002—$50.
(2) After July 16, 2002—$50 or the amount announced through proposed and final rulemaking in connection with ambulatory surgical center services.
(g) HCFA designates a predominant characteristic of a new technology IOL that both sets it apart from other IOLs and links it with other similar IOLs with the same characteristic to establish a specific subset of new technology within the “class of new technology IOLs.”
(h) Within 90 days of the end of the comment period following the FEDERAL REGISTER notice identified in paragraph (d) of this section, HCFA publishes in the FEDERAL REGISTER its determinations with regard to IOLs that
§416.190 It has determined are “new technology” lenses that qualify for a payment adjustment.

(i) Payment adjustments are effective beginning 30 days after the publication of HCFA’s determinations in the Federal Register.

§416.190 Who may request a review.

Any party who is able to furnish the information required in §416.195 may request that HCFA review the appropriateness of the payment amount provided under section 1833(i)(2)(A)(iii) of the Act with respect to an IOL that meets the definition of a new technology IOL in §416.180.

§416.195 A request to review.

(a) Content of a request. The request must include all of the following information:

(1) The name of the manufacturer, the model number, and the trade name of the IOL.

(2) A copy of the FDA’s summary of the IOL’s safety and effectiveness.

(3) A copy of the labeling claims of specific clinical advantages approved by the FDA for the IOL.

(4) A copy of the IOL’s original FDA approval notification.

(5) Reports of modifications made after the original FDA approval.

(6) Other information that HCFA finds necessary for identification of the IOL.

(b) Confidential information. To the extent that information received from an IOL manufacturer can reasonably be characterized as a trade secret or as privileged or confidential commercial or financial information, HCFA maintains the confidentiality of the information and protects it from disclosure not otherwise authorized or required by Federal law as allowed under Exemption 4 of the Freedom of Information Act (5 U.S.C. 552(b)(4)) and, with respect to trade secrets, the Trade Secrets Act (18 U.S.C. 1905).

§416.200 Application of the payment adjustment.

(a) HCFA recognizes the IOL(s) that define a new technology subset for purposes of this subpart as belonging to the class of new technology IOLs for a period of 5 years effective from the date that HCFA recognizes the first new technology IOL for a payment adjustment.

(b) Any IOL that HCFA subsequently recognizes as belonging to a new technology subset receives the new technology payment adjustment for the remainder of the 5-year period established with HCFA’s recognition of the first IOL in the subset.

(c) Beginning 5 years after the effective date of HCFA’s initial recognition of a new technology subset, payment adjustments cease for all IOLs that HCFA designates as belonging to that subset and payment reverts to the standard payment rate set under section 1833(i)(2)(A)(iii) of the Act for IOL insertion procedures performed in ASCs.

(d) ASCs that furnish an IOL designated by HCFA as belonging to the class of new technology IOLs must submit claims using specific billing codes to receive the new technology IOL payment adjustment.

PART 417—HEALTH MAINTENANCE ORGANIZATIONS, COMPETITIVE MEDICAL PLANS, AND HEALTH CARE PREPAYMENT PLANS

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AUTHORITY: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh), secs. 1301, 1306, and 1310 of the Public Health Service Act (42 U.S.C. 300e, 300e±5, and 300e±9), and 31 U.S.C. 9701.

§ 417.1 42 CFR Ch. IV (10–1–00 Edition)

1301 of the PHS Act and the regulations in subparts B and C of this part.

Health professionals means physicians (doctors of medicine and doctors of osteopathy), dentists, nurses, podiatrists, optometrists, physicians' assistants, clinical psychologists, social workers, pharmacists, nutritionists, occupational therapists, physical therapists, and other professionals engaged in the delivery of health services who are licensed, practice under an institutional license, are certified, or practice under authority of the HMO, a medical group, individual practice association, or other authority consistent with State law.

Individual practice association (IPA) means a partnership, association, corporation, or other legal entity that delivers or arranges for the delivery of health services and which has entered into written services arrangement or arrangements with health professionals, a majority of whom are licensed to practice medicine or osteopathy. The written services arrangement must provide:

(1) That these health professionals will provide their professional services in accordance with a compensation arrangement established by the entity; and

(2) To the extent feasible, for the sharing by these health professionals of health (including medical) and other records, equipment, and professional, technical, and administrative staff.

Medical group means a partnership, association, corporation, or other group:

(1) That is composed of health professionals licensed to practice medicine or osteopathy and of such other licensed health professionals (including dentists, optometrists, and podiatrists) as are necessary for the provision of health services for which the group is responsible;

(2) A majority of the members of which are licensed to practice medicine or osteopathy; and

(3) The members of which:

(i) After the end of the 48 month period beginning after the month in which the HMO for which the group provides health services becomes a qualified HMO, as their principal professional activity (over 50 percent individually) engage in the coordinated practice of their profession and as a group responsibility have substantial responsibility (over 35 percent in the aggregate of their professional activity) for the delivery of health services to enrollees of an HMO;

(ii) Pool their income from practice as members of the group and distribute it among themselves according to a prearranged salary or drawing account or other similar plan unrelated to the provision of specific health services;

(iii) Share health (including medical) records and substantial portions of major equipment and of professional, technical, and administrative staff;

(iv) Establish an arrangement whereby an enrollee's enrollment status is not known to the health professional who provides health services to the enrollee.

Medical group members means (1) a health professional engaged as a partner, associate, or shareholder in the medical group, or (2) any other health professional employed by the group who may be designated as a medical group member by the medical group.

Medically underserved population means the population of an urban or rural area as described in Sec. 417.912(d).

Nonmetropolitan area means an area no part of which is within a standard metropolitan statistical area as designated by the Office of Management and Budget and which does not contain a city whose population exceeds 50,000 individuals.

Party in interest means: (1) Any director, officer, partner, or employee responsible for management or administration of an HMO, any person who is directly or indirectly the beneficial owner of more than 5 percent of the equity of the HMO, any person who is the beneficial owner of a mortgage, deed of trust, note, or other interest secured by, and valuing more than 5 percent of the assets of the HMO, and, in the case of an HMO organized as a nonprofit corporation, an incorporator or member of the corporation under applicable State corporation law;

(2) Any entity in which a person described in paragraph (1):

(i) Is an officer or director;
§ 417.2 Basis and scope.

(a) Subparts B through F of this part pertain to the Federal qualification of HMOs under title XIII of the Public Health Service (PHS) Act.

(b) Subparts G through R of this part set forth the rules for Medicare contracts with, and payment to, HMOs and competitive medical plans (CMPs) under section 1876 of the Act.
§ 417.101 Health benefits plan: Basic health services.

(a) An HMO must provide or arrange for the provision of basic health services to its enrollees as needed and without limitations as to time and cost other than those prescribed in the PHS Act and these regulations, as follows:

1. Physician services (including consultant and referral services by a physician), which must be provided by a licensed physician, or if a service of a physician may also be provided under applicable State law by other health professionals, an HMO may provide the service through these other health professionals;

2. (i) Outpatient services, which must include diagnostic services, treatment services and x-ray services, for patients who are ambulatory and may be provided in a non-hospital based health care facility or at a hospital;

   (ii) Inpatient hospital services, which must include but not be limited to, room and board, general nursing care, meals and special diets when medically necessary, use of operating room and related facilities, use of intensive care unit and services, x-ray services, laboratory, and other diagnostic tests, drugs, medications, biologicals, anesthesia and oxygen services, special duty nursing when medically necessary, radiation therapy, inhalation therapy, and administration of whole blood and blood plasma;

   (iii) Outpatient services and inpatient hospital services must include short-term rehabilitation services and physical therapy, the provision of which the HMO determines can be expected to result in the significant improvement of a member's condition within a period of two months;

   (iv) Instructions to its enrollees on procedures to be followed to secure medically necessary emergency health services both in the service area and out of the service area;

   (v) Twenty outpatient visits per enrollee per year, as may be necessary and appropriate for short-term evaluative or crisis intervention mental health services, or both;

   (vi) Diagnosis, medical treatment and referral services (including referral services to appropriate ancillary services) for the abuse of or addiction to alcohol and drugs:

   (i) Diagnosis and medical treatment for the abuse of or addiction to alcohol and drugs must include detoxification for alcoholism or drug abuse on either an outpatient or inpatient basis, whichever is medically determined to be appropriate, in addition to the other required basic health services for the treatment of other medical conditions;

   (ii) Referral services may be either for medical or for nonmedical ancillary services. Medical services must be a part of basic health services; nonmedical ancillary services (such as vocational rehabilitation and employment counseling) and prolonged rehabilitation services in a specialized inpatient or residential facility need not be a part of basic health services;

   (vii) Diagnostic laboratory and diagnostic and therapeutic radiologic services in support of basic health services;

   (viii) Home health services provided at an enrollee's home by health care personnel, as prescribed or directed by the responsible physician or other authority designated by the HMO; and

   (ix) Preventive health services, which must be made available to members and must include at least the following:

       (i) A broad range of voluntary family planning services;

       (ii) Services for infertility;

       (iii) Well-child care from birth;

       (iv) Periodic health evaluations for adults;

       (v) Eye and ear examinations for children through age 17, to determine the need for vision and hearing correction; and

Subpart B—Qualified Health Maintenance Organizations: Services

Subpart B—Qualified Health Maintenance Organizations: Services

Subpart B—Qualified Health Maintenance Organizations: Services

2 CFR Ch. IV (10–1–00 Edition)
§ 417.103 Providers of basic and supplemental health services.

(a)(1) The HMO must provide that the services of health professionals that are provided as basic health services will, except as provided in paragraph (c) of this section, be provided or arranged for through (i) health professionals who are staff of the HMO, (ii) a medical group or groups, (iii) an IPA or

health service by the policymaking body of the HMO;

(12) Personal or comfort items and private rooms, unless medically necessary during inpatient hospitalization;

(13) Whole blood and blood plasma;

(14) Long-term physical therapy and rehabilitation;

(15) Durable medical equipment for home use (such as wheel chairs, surgical beds, respirators, dialysis machines); and

(16) Health services that are unusual and infrequently provided and not necessary for the protection of individual health, as approved by HCFA upon application by the HMO.

(e) An HMO may not offer to provide or arrange for the provision of basic health services on a prepayment basis that do not include all the basic health services set forth in paragraph (a) of this section or that are limited as to time and cost except in a manner prescribed by this subpart.


§ 417.102 Health benefits plan: Supplemental health services.

(a) An HMO may provide to its enrollees any health service that is not included as a basic health service under §417.101(a). These health services may be limited as to time and cost.

(b) An HMO must determine the level and scope of supplemental health services included with basic health services provided to its enrollees for a basic health services payment or those services offered to its enrollees as supplemental health services.

IPAs, (iv) physicians or other health professionals under direct service contracts with the HMO for the provision of these services, or (v) any combination of staff, medical group or groups, IPA or IPAs, or physicians or other health professionals under direct service contracts with the HMO.

(2) A staff or medical group model HMO may have as providers of basic health services physicians who have also entered into written services arrangements with an IPA or IPAs, but only if either (i) these physicians number less than 50 percent of the physicians who have entered into arrangements with the IPA or IPAs, or (ii) if the sharing is 50 percent or greater, HCFA approves the sharing as being consistent with the purposes of section 1310(b) of the PHS Act.

(3) After the 4 year period beginning with the month following the month in that an HMO becomes a qualified HMO, an entity that meets the requirements of the definition of medical group in §417.100, except for subdivision (3)(i) of that definition, may be considered a medical group if HCFA determines that the principal professional activity (over 50 percent individually) of the entity’s members is the coordinated practice of their profession, and if the HMO has demonstrated to the satisfaction of HCFA that the entity is committed to the delivery of medical services on a prepaid group practice basis by either:

(i) Presenting a reasonable time-phased plan for the entity to achieve compliance with the “substantial responsibility” requirement of subdivision (3)(i) of the definition of “medical group” in §417.100. The HMO must update the plan annually and must demonstrate to the satisfaction of HCFA that the entity is making continuous efforts and progress towards compliance with the requirements of the definition of “medical group,” or

(ii) Demonstrating that compliance by the entity with the “substantial responsibility” requirement is unreasonable or impractical because (A) the HMO serves a non-metropolitan or rural area as defined in §417.100, or (B) the entity is a multi-speciality group that provides medical consultation upon referral on a regional or national basis, or (C) the majority of the residents of the HMO’s service area are not eligible for employer-employee health benefits plans and the HMO has an insufficient number of enrollees to require utilization of at least 35 percent of the entity’s services.

(b) HMOs must have effective procedures to monitor utilization and control cost of basic and supplemental health services and to achieve utilization goals, which may include mechanisms such as risk sharing, financial incentives, or other provisions agreed to by providers.

(c) Paragraph (a) of this section does not apply to the provision of the services of a physician:

(1) Which the HMO determines are unusual or infrequently used services; or

(2) Which, because of an emergency, it was medically necessary to provide to the enrollee other than as required by paragraph (a) of this section; or

(3) Which are provided as part of the inpatient hospital services by employees or staff of a hospital or provided by staff of other entities such as community mental health centers, home health agencies, visiting nurses’ associations, independent laboratories, or family planning agencies.

(d) Supplemental health services must be provided or arranged for by the HMO and need not be provided by providers of basic health services under contract with the HMO.

(e) Each HMO must:

(1) Pay the provider, or reimburse its enrollees for the payment of reasonable charges for basic health services (or supplemental health services that the HMO agreed to provide on a prepayment basis) for which its enrollees have contracted, which were medically necessary and immediately required to be obtained other than through the HMO because of an unforeseen illness, injury, or condition, as determined by the HMO;

(2) Adopt procedures to review promptly all claims from enrollees for reimbursement for the provision of health services described in paragraph (e)(1) of this section, including a procedure for the determination of the medical necessity for obtaining the services other than through the HMO; and
§ 417.104 Payment for basic health services.

(a) Basic health services payment. Each HMO must provide or arrange for the provision of basic health services for a basic health services payment that:

(1) Is to be paid on a periodic basis without regard to the dates these services are provided;

(2) Is fixed without regard to the frequency, extent, or kind of basic health services actually furnished;

(3) Except as provided in paragraph (c) of this section, is fixed under a community rating system, as described in paragraph (b) of this section; and

(4) May be supplemented by nominal copayments which may be required for the provision of specific basic health services. Each HMO may establish one or more copayment options calculated on the basis of a community rating system.

(i) An HMO may not impose copayment charges that exceed 50 percent of the total cost of providing any single service to its enrollees, nor in the aggregate more than 20 percent of the total cost of providing all basic health services.

(ii) To insure that copayments are not a barrier to the utilization of health services or enrollment in the HMO, an HMO may not impose copayment charges on any subscriber (or enrollees covered by the subscriber’s contract with the HMO) in any calendar year, when the copayments made by the subscriber (or enrollees) in that calendar year total 200 percent of the total annual premium cost which that subscriber (or enrollees) would be required to pay if he (or they) were enrolled under an option with no copayments. This limitation applies only if the subscriber (or enrollees) demonstrates that copayments in that amount have been paid in that year.

(b) Community rating system. Under a community rating system, rates of payment for health services may be determined on a per person or per family basis, as described in paragraph (b)(1) of this section or on a per group basis as described in paragraph (b)(2) of this section. An HMO may fix its rates of payment under the system described in paragraph (b)(1) or (b)(2) of this section or under both such systems, but an HMO may use only one such system for fixing its rates of payment for any one group.

(1) A system of fixing rates of payment for health services may provide that the rates will be fixed on a per person or per family basis and may vary with the number of persons in a family. Except as otherwise authorized in this paragraph, these rates must be equivalent for all individuals and for all families of similar composition. Rates of payment may be based on either a schedule of rates charged to each subscriber group or on a per-enrollee-per-month (or per-subscriber-per-month) revenue requirement for the HMO. In the former event, rates may vary from group to group if the projected total revenue from each group is substantially equivalent to the revenue that would be derived if the schedule of rates were uniform for all groups. In the latter event, the payments from each group of subscribers must be calculated to yield revenues substantially equivalent to the product of the total number of enrollees (or subscribers) expected to be enrolled from the group and the per-enrollee-per-month (or per-subscriber-per-month) revenue requirement for the HMO. Under the system described in this paragraph, rates of payment may not vary because of actual or anticipated utilization of services by individuals associated with any specific group of subscribers. These provisions do not preclude changes in the rates of payment that are established for new enrollments or re-enrollments and that do not apply to existing contracts until the renewal of these contracts.

(2) A system of fixing rates of payment for health services may provide
that the rates will be fixed for individuals and families by groups. Except as otherwise authorized in this paragraph, such rates must be equivalent for all individuals in the same group and for all families of similar composition in the same group. If an HMO is to fix rates of payment for individuals and families by groups, it must:

(i) Classify all of the enrollees of the organization into classes based on factors that the HMO determines predict the differences in the use of health services by the individuals or families in each class and which have not been disapproved by HCFA,

(ii) Determine its revenue requirements for providing services to the enrollees of each class established under paragraph (b)(2)(i) of this section, and

(iii) Fix the rates of payment for the individuals and families of a group on the basis of a composite of the organization’s revenue requirements determined under paragraph (b)(2)(ii) of this section for providing services to them as members of the classes established under paragraph (b)(2)(i) of this section. HCFA will review the factors used by each HMO to establish classes under paragraph (b)(2)(i) of this section. If HCFA determines that any such factor may not reasonably be used to predict the use of the health services by individuals and families, HCFA will disapprove the factor for that purpose.

(3)(i) Nominal differentials in rates may be established to reflect differences in marketing costs and the different administrative costs of collecting payments from the following categories of potential subscribers:

(A) Individual (non-group) subscribers (including their families).

(B) Small groups of subscribers (100 subscribers or fewer).

(C) Large groups of subscribers (over 100 subscribers).

(ii) Differentials in rates may be established for subscribers enrolled in an HMO: (A) Under a contract with a governmental authority under section 1079 (“Contracts for Medical Care for Spouses and Children: Plans”) or section 1086 (“Contracts for Health Benefits for Certain Members, Former Members and their Dependents”) of title 10 (“Armed Forces”), United States Code; or (B) under any other governmental program (other than the health benefits program authorized by chapter 89 (“Health Insurance”) of title 5 (“Government Organization and Employees”), United States Code; or (C) under any health benefits program for employees of States, political subdivisions of states, and other public entities.

(4) An HMO may establish a separate community rate for separate regional components of the organization upon satisfactory demonstration to HCFA of the following:

(i) Each regional component is geographically distinct and separate from any other regional component; and

(ii) Each regional component provides substantially the full range of basic health services to its enrollees, without extensive referral between components of the organization for these services, and without substantial utilization by any two components of the same health care facilities. The separate community rate for each regional component of the HMO must be based on the different costs of providing health services in the respective regions.

(c) Exceptions to community rating requirement. (1) In the case of an HMO that provided comprehensive health services on a prepaid basis before it became a qualified HMO, the requirement of community rating shall not apply to the HMO during the forty-eight month period beginning with the month following the month in which it became a qualified HMO.

(2) The requirement of community rating does not apply to the basic health services payment for basic health services provided an enrollee who is a full-time student at an accredited institution of higher education.

(d) Late payment penalty. HMOs may charge a late payment penalty on accounts receivable that are in arrears.

(e) Review procedures for evaluating the community rating by class system under paragraph (b)(2). An HMO may establish a community rating system under paragraph (b)(2) of this section

1Further information entitled “Guidelines for Rating by Class” may be obtained from the Office of Prepaid Health Care, Division of Qualification Analysis, HHS Cohen Bldg., room 4360, 330 Independence Ave. SW., Washington, DC. 20020.
Health Care Financing Administration, HHS

§ 417.106 Quality assurance program; Availability, accessibility, and continuity of basic and supplemental health services.

(a) Quality assurance program. Each HMO or CMP must have an ongoing quality assurance program for its health services that meets the following conditions:

1. Stresses health outcomes to the extent consistent with the state of the art.

2. Provides review by physicians and other health professionals of the process followed in the provision of health services.

3. Uses systematic data collection of performance and patient results, provides interpretation of these data to its practitioners, and institutes needed change.

4. Includes written procedures for taking appropriate remedial action whenever, as determined under the quality assurance program, inappropriate or substandard services have been provided or services that ought to have been furnished have not been provided.

(b) Availability and accessibility of health care services. Basic health services and those supplemental health services for which enrollees have contracted must be provided or arranged for by the HMO in accordance with the following rules:

1. Except as provided in paragraph (b)(2) of this section, the services must
be available to each enrollee within the HMO's service area.

(2) Exception. If the HMO's service area is located wholly within a non-metropolitan area, the HMO may make available outside its service area any basic health service that is not a primary care or emergency care service, if the number of providers of that basic health service who will provide the service to the HMO's enrollees is insufficient to meet the demand. As used in this paragraph, primary care includes general practice, family practice, general internal medicine, general pediatrics, and general obstetrics and gynecology. An HMO that provides the services covered by these fields through at least a general or family practitioner, or a pediatrician and a general internist, is considered to be providing primary care.

(3) The services must be available and accessible with reasonable promptness to each of the HMO's enrollees as ensured through—

(i) Staffing patterns within generally accepted norms for meeting the projected enrollment needs; and

(ii) Geographic location, hours of operation, and arrangements for after-hours services. (Medically necessary emergency services must be available 24 hours a day, 7 days a week.)

(c) Continuity of care. The HMO must ensure continuity of care through arrangements that include but are not limited to the following:

(1) Use of a health professional who is primarily responsible for coordinating the enrollee's overall health care.

(2) A system of health and medical records that accumulates pertinent information about the enrollee's health care and makes it available to appropriate professionals.

(3) Arrangements made directly or through the HMO's providers to ensure that the HMO or the health professional who coordinates the enrollee's overall health care is kept informed about the services that the referral resources furnish to the enrollee.

(d) Confidentiality of health records. Each HMO must establish adequate procedures to ensure the confidentiality of the health and medical records of its enrollees.

[58 FR 38068, July 15, 1993]
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(C) Sources and uses of funds statements; and
(D) Balance sheets.

(b) Assumption of financial risk. Each HMO must assume full financial risk on a prospective basis for the provision of basic health services, except that it may obtain insurance or make other arrangements as follows:
(1) For the cost of providing to any enrollee basic health services with an aggregate value of more than $5,000 in any year.
(2) For the cost of basic health services obtained by its enrollees from sources other than the HMO because medical necessity required that they be furnished before they could be secured through the HMO.
(3) For not more than 90 percent of the amount by which its costs for any of its fiscal years exceed 115 percent of its income for that fiscal year.
(4) For physicians or other health professionals, health care institutions, or any other combination of such individuals or institutions to assume all or part of the financial risk on a prospective basis for their furnishing of basic health services to the HMO's enrollees.

§ 417.122 Protection of enrollees.

(a) Liability protection. (1) Each HMO must adopt and maintain arrangements satisfactory to HCFA to protect its enrollees from incurring liability for payment of any fees that are the legal obligation of the HMO. These arrangements may include any of the following:
(i) Contractual arrangements that prohibit health care providers used by the enrollees from holding any enrollee liable for payment of any fees that are the legal obligation of the HMO.
(ii) Insurance, acceptable to HCFA.
(iii) Financial reserves, acceptable to HCFA, that are held for the HMO and restricted for use only in the event of insolvency.
(iv) Any other arrangements acceptable to HCFA.
(2) The requirements of this paragraph do not apply to an HMO if HCFA determines that State law protects the HMO enrollees from liability for payment of any fees that are the legal obligation of the HMO.

(b) Protection against loss of benefits if the HMO becomes insolvent. The insolvency protection plan required under §417.120(a) must provide for continuation of benefits as follows:
(1) For all enrollees, for the duration of the contract period for which payment has been made.
(2) For enrollees who are in an inpatient facility on the date of insolvency, until they are discharged from the facility.

§ 417.124 Administration and management.

(a) General requirements. Each HMO must have administrative and managerial arrangements satisfactory to HCFA, as demonstrated by at least the following:
(1) A policymaking body that exercises oversight and control over the HMO's policies and personnel to ensure that management actions are in the best interest of the HMO and its enrollees.
(2) Personnel and systems sufficient for the HMO to organize, plan, control and evaluate the financial, marketing, health services, quality assurance program, administrative and management aspects of the HMO.
(3) At a minimum, management by an executive whose appointment and removal are under the control of the HMO's policymaking body.
(b) Full and fair disclosure—(1) Basic rule. Each HMO must prepare a written description of the following:
(i) Benefits (including limitations and exclusions).
(ii) Coverage (including a statement of conditions on eligibility for benefits).
(iii) Procedures to be followed in obtaining benefits and a description of circumstances under which benefits may be denied.
(iv) Rates.
(v) Grievance procedures.
(vi) Service area.
(vii) Participating providers.
(viii) Financial condition including at least the following most recently audited information: Current assets, other assets, total assets; current liabilities, long term liabilities; and net worth.
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(2) Requirements for the description. (i) The description must be written in a way that can be easily understood by the average person who might enroll in the HMO.

(ii) The description of benefits and coverage may be in general terms if reference is made to a detailed statement of benefits and coverage that is available without cost to any person who enrolls in the HMO or to whom the opportunity for enrollment is offered.

(iii) The HMO must provide the description to any enrollee or person who is eligible to elect the HMO option and who requests the material from the HMO or the administrator of a health benefits plan. For purposes of this requirement, “administrator” (of a health benefits plan) has the meaning it is given in the Employment Retirement Income Security Act of 1974 (ERISA) at 29 U.S.C. 1002(16)(A).

(iv) If the HMO provides health services through individual practice associations (IPAs), the HMO must specify the number of member physicians by specialty, and a listing of the hospitals where HMO enrollees will receive basic and supplemental health services.

(v) If the HMO provides health services other than through IPAs, the HMO must specify, for each ambulatory care facility, the facility’s address, days and hours of operation, and the number of physicians by specialty, and a listing of the hospitals where HMO enrollees will receive basic and supplemental health services.

(c) Broadly representative enrollment. (1) Each HMO must offer enrollment to persons who are broadly representative of the various age, social, and income groups within its service area.

(2) If an HMO has a medically underserved population located in its service area, not more than 75 percent of its enrollees may be from the medically underserved population unless the area in which that population resides is a rural area.

(d) Health status and enrollment. (1) The HMO may not, on the basis of health status, health care needs, or age of the individual—

(i) Expel or refuse to reenroll any enrollee; or

(ii) Refuse to enroll individual members of a group.

(2) For purposes of this paragraph, a “group” is composed of individuals who enroll in the HMO under a contract or other arrangement that covers two or more subscribers. Examples of groups are employees who enroll under a contract between their employer and the HMO, or members of an organization that arranges coverage for its membership.

(3) Nothing in this subpart prohibits an HMO from requiring that, as a condition for continued eligibility for enrollment, enrolled dependent children, upon reaching a specified age, convert to individual enrollment, consistent with paragraph (e) of this section.

(e) Conversion of enrollment. (1) Each HMO must offer individual enrollment to the following:

(i) Each enrollee (and his or her enrolled dependents) leaving a group.

(ii) Each enrollee who otherwise cease to be eligible for HMO enrollment because of his or her age, or the death or divorce of an enrollee.

(2) The individual enrollment offered must meet the conditions of subpart B of this part and this subpart C.

(3) The HMO is not required to offer individual enrollment except to the enrollees specified in this paragraph.

(4) The HMO must offer the enrollment on the same terms and conditions that it makes available to other nongroup enrollees.

(f) [Reserved]

(g) Grievance procedures. Each HMO must have and use meaningful procedures for hearing and resolving grievances between the HMO’s enrollees and the HMO, including the HMO staff and medical groups and IPAs that furnish services. These procedures must ensure that:

(1) Grievances and complaints are transmitted in a timely manner to appropriate HMO decisionmaking levels that have authority to take corrective action; and

(2) Appropriate action is taken promptly, including a full investigation if necessary and notification of concerned parties as to the results of the HMO’s investigation.

(h) Certification of institutional providers. Each HMO must ensure that its affiliated institutional providers meet one of the following conditions:
(1) In the case of hospitals, are either accredited by the Joint Commission on Accreditation of Health Care Organizations, or certified by Medicare.

(2) In the case of laboratories, are either CLIA-exempt, or have in effect a valid certificate of one of the following types, issued by HCFA in accordance with section 353 of the PHS Act and part 493 of this chapter:
   (i) Registration certificate.
   (ii) Certificate.
   (iii) Certificate of waiver.
   (iv) Certificate of accreditation.

(3) In the case of other affiliated institutional providers, are certified for participation in Medicare and Medicaid in accordance with part 405, 416, 418, 488, or 491 of this chapter, as appropriate.


§ 417.126 Recordkeeping and reporting requirements.

(a) General reporting and disclosure requirements. Each HMO must have an effective procedure to develop, compile, evaluate, and report to HCFA, to its enrollees, and to the general public, at the times and in the manner that HCFA requires, and while safeguarding the confidentiality of the doctor-patient relationship, statistics and other information with respect to the following:
   (1) The cost of its operations.
   (2) The patterns of utilization of its services.
   (3) The availability, accessibility, and acceptability of its services.
   (4) The extent practical, developments in the health status of its enrollees.
   (5) Information demonstrating that the HMO has a fiscally sound operation.
   (6) Other matters that HCFA may require.

(b) Significant business transactions. Each HMO must report to HCFA annually, within 120 days of the end of its fiscal year (unless for good cause shown, HCFA authorizes an extension of time), the following:
   (1) A description of significant business transactions (as defined in paragraph (c) of this section) between the HMO and a party in interest.
   (2) With respect to those transactions—
      (i) A showing that the costs of the transactions listed in paragraph (c) of this section do not exceed the costs that would be incurred if these transactions were with someone who is not a party in interest; or
      (ii) If they do exceed, a justification that the higher costs are consistent with prudent management and fiscal soundness requirements.
   (3) A combined financial statement for the HMO and a party in interest if either of the following conditions is met:
      (i) Thirty-five percent or more of the costs of operation of the HMO go to a party in interest.
      (ii) Thirty-five percent or more of the revenue of a party in interest is from the HMO.

(c) Significant business transaction defined. As used in paragraph (b) of this section—
   (1) Business transaction means any of the following kinds of transactions:
      (i) Sale, exchange or lease of property.
      (ii) Loan of money or extension of credit.
      (iii) Goods, services, or facilities furnished for a monetary consideration, including management services, but not including—
         (A) Salaries paid to employees for services performed in the normal course of their employment; or
         (B) Health services furnished to the HMO’s enrollees by hospitals and other providers, and by HMO staff, medical groups, or IPAs, or by any combination of those entities.
   (2) Significant business transaction means any business transaction or series of transactions of the kind specified in paragraph (c)(1) of this section that, during any fiscal year of the HMO, have a total value that exceeds $25,000 or 5 percent of the HMO’s total operating expenses, whichever is less.
   (d) Requirements for combined financial statements. (1) The combined financial statements required by paragraph (b)(3) of this section must display in separate columns the financial information for the HMO and each of these parties in interest.
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(2) Inter-entity transactions must be eliminated in the consolidated column.

(3) These statements must have been examined by an independent auditor in accordance with generally accepted accounting principles, and must include appropriate opinions and notes.

(4) Upon written request from an HMO showing good cause, HCFA may waive the requirement that its combined financial statement include the financial information required in this paragraph (d) with respect to a particular entity.

(e) Reporting and disclosure under ERISA. (1) For any employees' health benefits plan that includes an HMO in its offerings, the HMO must furnish, upon request, the information the plan needs to fulfill its reporting and disclosure obligations (with respect to the particular HMO) under the Employee Retirement Income Security Act of 1974 (ERISA).

(i) The HMO must furnish the information to the employer or the employer's designee, or to the plan administrator, as the term “administrator” is defined in ERISA.

(ii) Loan of money or extension of credit.

(iii) Goods, services, or facilities furnished for a monetary consideration, including management services, but not including—

(A) Salaries paid to employees for services performed in the normal course of their employment; or

(B) Health services furnished to the HMO's enrollees by hospitals and other providers, and by HMO staff, medical groups, or IPAs, or by any combination of those entities.

(2) Significant business transaction means any business transaction or series of transactions of the kind specified in paragraph (c)(1) of this section that, during any fiscal year of the HMO, have a total value that exceeds $25,000 or 5 percent of the HMO's total operating expenses, whichever is less.

(d) Requirements for combined financial statements. (1) The combined financial statements required by paragraph (b)(3) of this section must display in separate columns the financial information for the HMO and each of these parties in interest.

(2) Inter-entity transactions must be eliminated in the consolidated column.

(3) These statements must have been examined by an independent auditor in accordance with generally accepted accounting principles, and must include appropriate opinions and notes.

(4) Upon written request from an HMO showing good cause, HCFA may waive the requirement that its combined financial statement include the financial information required in this paragraph (d) with respect to a particular entity.

(e) Reporting and disclosure under ERISA. (1) For any employees' health benefits plan that includes an HMO in its offerings, the HMO must furnish, upon request, the information the plan needs to fulfill its reporting and disclosure obligations (with respect to the particular HMO) under the Employee Retirement Income Security Act of 1974 (ERISA).

(2) The HMO must furnish the information to the employer or the employer's designee, or to the plan administrator, as the term “administrator” is defined in ERISA.

Subpart D—Application for Federal Qualification

§ 417.142

This subpart sets forth—

(a) The requirements for—

(1) Entities that seek qualification as HMOs under title XIII of the PHS Act; and

(2) HMOs that seek—

(i) Qualification for their regional components; or

(ii) Expansion of their service areas; or

(b) The procedures that HCFA follows to make determinations; and

(c) Other related provisions, including application fees.

[59 FR 40896, Sept. 30, 1994]
(2) HCFA determines whether the entity is an HMO on the basis of the entity’s application and any additional information and investigation (including site visits) that HCFA may require.

(3) HCFA may determine that an entity is any of the following:

(i) An operational qualified HMO.

(ii) A preoperational qualified HMO.

(iii) A transitional qualified HMO.

(b) Operational qualified HMO. HCFA determines that an entity is an operational qualified HMO if—

(1) HCFA finds that the entity meets the requirements of subparts B and C of this part.

(2) The entity, within 30 days of HCFA’s determination, provides written assurances, satisfactory to HCFA, that it—

(i) Provides and will provide basic health services (and any supplemental health services included in any contract) to its enrollees;

(ii) Provides and will provide these services in the manner prescribed in sections 1301(b) and 1301(c) of the PHS Act and subpart B of this part;

(iii) Is organized and operated and will continue to be organized and operated in the manner prescribed in section 1301(c) of the PHS Act and subpart C of this part;

(iv) Under arrangements that safeguard the confidentiality of patient information and records, will provide access to HCFA and the Comptroller General or any of their duly authorized representatives for the purpose of audit, examination or evaluation to any books, documents, papers, and records of the entity relating to its operation as an HMO, and to any facilities that it operates; and

(v) Will continue to comply with any other assurances that it has given to HCFA.

(c) Preoperational qualified HMO. (1) HCFA may determine that an entity is a preoperational qualified HMO if it provides, within 30 days of HCFA’s determination, satisfactory assurances that it will become operational within 60 days following that determination and will, when it becomes operational, meet the requirements of subparts B and C of this part.

(2) Within 30 days after receiving notice that the entity has begun operation, HCFA determines whether it is an operational qualified HMO. In the absence of this determination, the entity is not an operational qualified HMO even though it becomes operational.

(d) Transitional qualified HMO: General rules—(1) Basic requirements. HCFA may determine that an entity is a transitional qualified HMO if the entity—

(i) Meets the requirements of paragraph (d)(2) through (d)(4) of this section; and

(ii) Provides the assurances specified in paragraphs (d)(5) through (d)(7) of this section within 30 days of HCFA’s determination.

(2) Organization and operation. The entity is organized and operated in accordance with subpart C of this part, except that it need not—

(i) Assume full financial risk for the provision of basic health services as required by §417.120(b); or

(ii) Comply with the limitations that are imposed on insurance by §417.120(b)(1).

(3) Range of services. The entity is currently providing the following services on a prepaid basis:

(i) Physician services.

(ii) Outpatient services and inpatient hospital services. (The entity need not provide or pay for hospital inpatient or outpatient services that it can show are being provided directly, through insurance, or under arrangements, by other entities.)

(iii) Medically necessary emergency services.

(iv) Diagnostic laboratory services and diagnostic and therapeutic radiologic services.

These services must meet the requirements of §417.101, but may be limited in time and cost without regard to the constraints imposed by §417.101(a).

(4) Payment for services—(i) General rule. The entity pays for basic health services in accordance with §417.104, except that it need not comply with the copayments limitations imposed by §417.104(a)(4).

(ii) Determination of payment rates. In determining payment rates, the entity need not comply with the community rating requirements of §§417.104(b) and 417.105(b).
§ 417.143 Application requirements.

(a) General requirements. This section sets forth application requirements for entities that seek qualification as HMOs; HMOs that seek expansion of their service areas; and HMOs that seek qualification of their regional components as HMOs.

(b) Completion of an application form.

(1) In order to receive a determination concerning whether an entity is a qualified regional component, an HMO that has more than one regional component is considered qualified for those regional components for which assurances have been signed in accordance with this section.

(g) Special rules. Enrollees entitled to Medicare or Medicaid. For an HMO that accepts enrollees entitled to Medicare or Medicaid, the following rules apply:

(1) The requirements of title XVIII and XIX of the Act, as appropriate, take precedence over conflicting requirements of sections 1301(b) and 1301(c) of the PHS Act.

(2) An HMO that complies with paragraph (g)(2) of this section may obtain and retain Federal qualification if, for its other enrollees, the HMO meets the requirements of sections 1301(b) and 1301(c) of the PHS Act and implementing regulations in this subpart D and subparts B and C of this part.

(h) Special rules: Enrollees under the Federal employee health benefits program (FEHBP). An HMO that accepts enrollees under the FEHBP (Chapter 89 of title 5 of the U.S.C.) may obtain and retain Federal qualification if, for its other enrollees, it complies with the requirements of section 1301(b) and 1301(c) of the PHS Act and implementing regulations in this subpart D and subparts B and C of this part.

[59 FR 48636, Sept. 30, 1994]
qualified HMO, an individual authorized to act for the entity (the applicant) must complete an application form provided by HCFA.

(2) The authorized individual must describe thoroughly how the entity meets, or will meet, the requirements for qualified HMOs described in the PHS Act and in subparts B and C of this part, this subpart D, and 417.168 and 417.169 of subpart F.

(c) Collection of an application fee. In accordance with the requirements of 31 U.S.C. 9701, Fees and charges for Government services and things of value, HCFA determines the amount of the application fee that must be submitted with each type of application.

(1) The fee is reasonably related to the Federal government’s cost of qualifying an entity and may vary based on the type of application.

(2) Each type of application has one set fee rather than a charge based on the specific cost of each determination. (For example, each Federally qualified HMO applicant seeking Federal qualification of one of its regional components as an HMO is charged the same amount, unless the amount of the fee has been changed under paragraph (f) of this section.)

(d) Application fee amounts. The application fee amounts for applications completed on or after July 13, 1987 are as follows:

(1) $18,400 for an entity seeking qualification as an HMO or qualification of a regional component of an HMO.

If, in the case of an HMO seeking qualification of a regional component, HCFA determines that there is no need for a site visit, $8,000 will be returned to the applicant.

(2) $6,900 for an HMO seeking expansion of its service area.

(3) $3,100 for a CMP seeking qualification as an HMO.

(e) Refund of an application fee. HCFA refunds an application fee only if the entity withdraws its application within 10 working days after receipt by HCFA. Application fees are not returned in any other circumstance, even if qualification or certification is denied.

(f) Procedure for changing the amount of an application fee. If HCFA determines that a change in the amount of a fee is appropriate, HCFA issues a notice of proposed rulemaking in the FEDERAL REGISTER to announce the proposed new amount.

(g) New application after denial. An entity may not submit another application under this subpart for the same type of determination for four full months after the date of the notice in which HCFA denied the application.

(h) Disclosure of application information under the Freedom of Information Act. An applicant submitting material that he or she believes is protected from disclosure under 5 U.S.C. 552, the Freedom of Information Act, or because of exceptions provided in 45 CFR part 5, the Department’s regulations providing exceptions to disclosure, should label the material “privileged” and include an explanation of the applicability of an exception described in 45 CFR part 5.


§ 417.144 Evaluation and determination procedures.

(a) Basis for evaluation and determination. (1) HCFA evaluates an application for Federal qualification on the basis of information contained in the application itself and any additional information that HCFA obtains through on-site visits, public hearings, and any other appropriate procedures.

(2) If the application is incomplete, HCFA notifies the entity and allows 60 days from the date of the notice for the entity to furnish the missing information.

(b) Notice of determination. (1) If HCFA finds that the entity does not appear to meet the requirements for qualification and appears to be able to meet those requirements within 60 days, HCFA gives the entity notice of intent to deny.
qualification and a summary of the basis for this preliminary finding.

(2) Within 60 days from the date of the notice, the entity may respond in writing to the issues or other matters that were the basis for HCFA’s preliminary finding, and may revise its application to remedy any defects identified by HCFA.

(d) Denial and reconsideration of denial.

(1) If HCFA denies an application for qualification under this subpart, HCFA gives the entity written notice of the denial and an opportunity to request reconsideration of that determination.

(2) A request for reconsideration must—

(i) Be submitted in writing, within 60 days following the date of the notice of denial;

(ii) Be addressed to the HCFA officer or employee who denied the application; and

(iii) Set forth the grounds upon which the entity requests reconsideration, specifying the material issues of fact and of law upon which the entity relies.

(3) HCFA bases its reconsideration upon the record compiled during the qualification review proceedings, materials submitted in support of the request for reconsideration, and other relevant materials available to HCFA.

(4) HCFA gives the entity written notice of the reconsidered determination and the basis for the determination.

(e) Information on qualified HMOs—FEDERAL REGISTER notices. In quarterly FEDERAL REGISTER notices, HCFA gives the names, addresses, and service areas of newly qualified HMOs and describes the expanded service areas of other qualified HMOs.

(2) Listings. A cumulative list of qualified HMOs is available from the following office, which is open from 8:30 a.m. to 5 p.m., Monday through Friday: Office of Managed Care, room 4360, Cohen Building, 400 Independence Avenue S.W., Washington, DC 20201.

§ 417.150 Definitions.

As used in this subpart, unless the context indicates otherwise—

Agreement means a collective bargaining agreement.

Bargaining representative means an individual or entity designated or selected, under any applicable Federal, State, or local law, or public entity collective bargaining agreement, to represent employees in collective bargaining, or any other employee representative designated or selected under any law.

Carrier means a voluntary association, corporation, partnership, or other organization that is engaged in providing, paying for, or reimbursing all or part of the cost of health benefits under group insurance policies or contracts, medical or hospital service agreements, enrollment or subscription contracts, or similar group arrangements, in consideration of premiums or other periodic charges payable to the carrier.

Collective bargaining agreement means an agreement entered into between an employing entity and the bargaining representative of its employees.

Contract means an employer-employee or public entity-employee contract, or a contract for health benefits.

Designee means any person or entity authorized to act on behalf of an employing entity or a group of employing entities to offer the option of enrollment in a qualified health maintenance organization to their eligible employees.

Eligible employee means an employee who meets the employer’s requirements for participation in the health benefits plan.

Employee means any individual employed by an employer or public entity on a full-time or part-time basis.
Employer has the meaning given that term in section 3(d) of the Fair Labor Standards Act of 1938, except that it—
(1) includes non-appropriated fund instrumentalities of the United States Government; and
(2) excludes the following:
(i) The governments of the United States, the District of Columbia and the territories and possessions of the United States, the 50 States and their political subdivisions, and any agencies or instrumentalities of any of the foregoing, including the United States Postal Service and Postal Rate Commission.
(ii) Any church, or convention or association of churches, and any organization operated, supervised, or controlled by a church, or convention or association of churches that meets the following conditions:
(A) Is an organization that is described in section 501(c)(3) of the Internal Revenue Code of 1954.
(B) Does not discriminate, in the employment, compensation, promotion or termination of employment of any personnel, or in the granting of staff and other privileges to physicians or other health personnel, on the grounds that the individuals obtain health care through HMOs, or participate in furnishing health care through HMOs.
Employing entity means an employer or public entity.
Employing entity-employee contract means a legally enforceable agreement (other than a collective bargaining agreement) between an employing entity and its employees for the provision of, or payment for, health benefits for its employees, or for its employees and their eligible dependents.
Group enrollment period means the period of at least 10 working days each calendar year during which each eligible employee is given the opportunity to select among the alternatives included in a health benefits plan.
Health benefits means health benefits and services.
Health benefits contract means a contract or other agreement between an employing entity or a designee and a carrier for the provision of, or payment for, health benefits to eligible employees or to eligible employees and their eligible dependents.
Health benefits plan means any arrangement, to provide or pay for health services, that is offered to eligible employees, or to eligible employees and their eligible dependents, by or on behalf of an employing entity.
Public entity means the 50 states, Puerto Rico, Guam, the Virgin Islands, the Northern Mariana Islands and American Samoa and their political subdivisions, the District of Columbia, and any agency or instrumentality of the foregoing, and political subdivisions include counties, parishes, townships, cities, municipalities, towns, villages, and incorporated villages.
Qualified HMO means an HMO that has in effect a determination, made under subpart D of this part, that the HMO is an operational, preoperational, or transitional qualified HMO.
To offer a health benefits plan means to make participation in a health benefits plan available to eligible employees, or to eligible employees and their eligible dependents regardless of whether the employing entity makes a financial contribution to the plan on behalf of these employees, directly or indirectly, for example, through payments on any basis into a health and welfare trust fund.

§ 417.151 Applicability.

(a) Basic rule. Effective October 24, 1995,1 this subpart applies to any employing entity that offers a health benefits plan to its employees, meets the conditions specified in paragraphs (b) through (e) of this section, and elects to include one or more qualified HMOs in the health plan alternatives it offers its employees.

1Before October 24, 1995, an employing entity that met the conditions specified in §417.151 was required to include one or more qualified HMOs, if it received from at least one qualified HMO a written request for inclusion and that request met the timing, content, and procedural requirements specified in §417.152.
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(b) Number of employees. During any calendar quarter of the preceding calendar year, the employer or public entity employed an average of not less than 25 employees.

(c) Minimum wage. During any calendar quarter of the preceding calendar year, the employer was required to pay the minimum wage specified in section 6 of the Fair Labor Standards Act of 1938, or would have been required to pay that wage but for section 13(a) of that Act.

(d) Federal assistance under section 317 of the PHS Act. The public entity has a pending application for, or is receiving, assistance under section 317 of the PHS Act.

(e) Employees in HMO's service area. At least 25 of the employing entity's employees reside within the HMO's service area.


§ 417.153 Offer of HMO alternative.

(a) Basic rule. An employing entity that is subject to this subpart and that elects to include one or more qualified HMOs must offer the HMO alternative in accordance with this section.

(b) Employees to whom the HMO option must be offered. Each employing entity must offer the option of enrollment in a qualified HMO to each eligible employee and his or her eligible dependents who reside in the HMO's service area.

(c) Manner of offering the HMO option.

(1) For employees who are represented by a bargaining representative, the option of enrollment in a qualified HMO—

(i) Must first be presented to the bargaining representative; and

(ii) If the representative accepts the option, must then be offered to each represented employee.

(2) For employees not represented by a bargaining representative, the option must be offered directly to those employees.


§ 417.155 How the HMO option must be included in the health benefits plan.

(a) HMO access to employees—(1) Purpose and timing.

(i) Purpose. The employing entity must provide each HMO included in its health benefits plan fair and reasonable access to all employees specified in §417.153(b), so that the HMO can explain its program in accordance with §417.124(b).

(ii) Timing. The employing entity must provide access beginning at least 30 days before, and continuing during, the group enrollment period.

(b) Review of HMO offering materials.

(1) The HMO must give the employing entity or designee opportunity to review, revise, and approve HMO educational and offering materials before distribution.

(2) Revisions must be limited to correcting factual errors and misleading or ambiguous statements, unless—

(i) The HMO and the employing entity agree otherwise; or

(ii) Other revisions are required by law.

(3) The employing entity or designee must complete revision of the materials promptly so as not to delay or otherwise interfere with their use during the group enrollment period.

(c) Group enrollment period; prohibition of restrictions; effective date of HMO coverage—(1) Prohibition of restrictions. If an employing entity or designee includes the option of enrollment in a qualified HMO in the health benefits plan offered to its eligible employees, it must provide a group enrollment period before the effective date of HMO coverage. The employing entity may not impose waiting periods as a condition of enrollment in the HMO or of transfer from HMO to non-HMO coverage, or exclusions, or limitations based on health status.
(2) Effective date of coverage. Unless otherwise agreed to by the employing entity, or designee, and the HMO, coverage under the HMO contract for employees selecting the HMO option begins on the day the non-HMO contract expires or is renewed without lapse.

(3) Coordination of benefits. Nothing in this subpart precludes the uniform application of coordination of benefits agreements between the HMOs and the other carriers that are included in the health benefits plan.

(d) Continued eligibility for “free-standing” health benefits—(1) Basic requirement. At the request of a qualified HMO, the employing entity or its designee must provide that employees selecting the option of HMO membership will not, because of this selection, lose their eligibility for free-standing dental, optical, or prescription drug benefits for which they were previously eligible or would be eligible if selecting a non-HMO option and that are not included in the services provided by the HMO to its enrollees as part of the HMO prepaid benefit package.

(2) “Free-standing” defined. For purposes of this paragraph, the term “free-standing” refers to a benefit that—

(i) Is not integrated or incorporated into a basic health benefits package or major medical plan, and

(ii) Is—

(A) Offered by a carrier other than the one offering the basic health benefits package or major medical plan; or

(B) Subject to a premium separate from the premium for the basic health benefits package or major medical plan.

(3) Examples of the employing entity’s obligation with respect to the continued eligibility—(i) The health benefits plan includes a free-standing dental benefit. The HMO does not offer any dental coverage as part of its health services provided to members on a prepaid basis. The employing entity must provide employees who select the HMO option continue to be eligible for dental coverage. (If the dental coverage is not optional for employees selecting the non-HMO option, nothing in this regulation requires that the coverage be mandatory for employees selecting the non-HMO option.)

(ii) The non-HMO option provides free-standing coverage for optical services (such as refraction and the provision of eyeglasses), and the HMO does not. The employing entity must provide that employees who select the HMO option continue to be eligible for optical coverage.

(iii) The non-HMO option includes dental coverage in its major medical package, with a common deductible applied to dental as well as non-dental benefits. The HMO provides no dental coverage as part of its pre-paid health services. Because the dental coverage is not free-standing, the employing entity is not required to provide that employees who select the HMO option continue to be eligible for dental coverage, but is free to do so.

(e) Opportunity to select among coverage options: Requirement for affirmative written selection—(1) Opportunity other than during a group enrollment period. The employing entity or designee must provide opportunity (in addition to the group enrollment period) for selection among coverage options, by eligible employees who meet any of the following conditions:

(i) Are new employees.

(ii) Have been transferred or have changed their place of residence, resulting in—

(A) Eligibility for enrollment in a qualified HMO for which they were not previously eligible by place of residence; or

(B) Residence outside the service area of a qualified HMO in which they were previously enrolled.

(iii) Are covered by any coverage option that ceases operation.

(2) Prohibition of restrictions. When the employees specified in paragraph (e)(1) of this section are eligible to participate in the health benefits plan, the employing entity or designee must make available, without waiting periods or exclusions based on health status as a condition, the opportunity to enroll in an HMO, or transfer from HMO coverage to non-HMO coverage.
(3) **Affirmative written selection.** The employing entity or designee must require that the eligible employee make an affirmative written selection in any of the following circumstances:

(i) Enrollment in a particular qualified HMO is offered for the first time.

(ii) The eligible employee elects to change from one option to another.

(iii) The eligible employee is one of those specified in paragraph (e)(1) of this section.

(f) **Determination of copayment levels and supplemental health services.** The selection of a copayment level and of supplemental health services to be contracted for must be made as follows:

(1) For employees represented by a collective bargaining representative, the selection of copayment levels and supplemental health services is subject to the collective bargaining process.

(2) For employees not represented by a bargaining representative, the selection of copayment levels and supplemental health services is subject to the same decisionmaking process used by the employing entity with respect to the non-HMO option in its health benefits plan.

(3) In all cases, the HMO has the right to include, with the basic benefits package it provides to its enrollees for a basic health services payment, on a non-negotiable basis, those supplemental health services that meet the following conditions:

(i) Are required to be offered under State law.

(ii) Are included uniformly by the HMO in its prepaid benefit package.

(iii) Are available to employees who select the non-HMO option but not available to those who select the HMO option.


§ 417.156 When the HMO must be offered to employees.

(a) General rules. (1) The employing entity or designee must offer eligible employees the option of enrollment in a qualified HMO at the earliest date permitted under the terms of existing agreements or contracts.

(2) If the HMO's request for inclusion in a health benefits plan is received at a time when existing contracts or agreements do not provide for inclusion, the employing entity must include the HMO option in the health benefits plan at the time that new agreements or contracts are offered or negotiated.

(b) Specific requirements. Unless mutually agreed otherwise, the following rules apply:

(1) Collective bargaining agreement. The employing entity or designee must raise the HMO's request during the collective bargaining process—

(i) When a new agreement is negotiated;

(ii) At the time prescribed, in an agreement with a fixed term of more than 1 year, for discussion in change in health benefits; or

(iii) In accordance with a specific process for review of HMO offers.

(2) Contracts. For employees not covered by a collective bargaining agreement, the employing entity or designee must include the HMO option in any health benefits plan offered to eligible employees when the existing contract is renewed or when a new health benefits contract or other arrangement is negotiated.

(i) If a contract has no fixed term or has a term in excess of 1 year, the contract must be treated as renewable on its earliest anniversary date.

(ii) If the employing entity or designee is self-insured, the budget year must be treated as the term of the existing contract.

(3) Multiple arrangements. In the case of a health benefits plan that includes multiple contracts or other arrangements with varying expiration or renewal dates, the employing entity must include the HMO option, in accordance with paragraphs (b)(1) and (b)(2) of this section,—

(i) At the time each contract or arrangement is renewed or reissued; or

(ii) The benefits provided under the contract or arrangement are offered to employees.

[59 FR 49841, Sept. 30, 1994]

§ 417.157 Contributions for the HMO alternative.

(a) General principles—(1) Non-discrimination. The employer contribution to an HMO must be in an amount that does not discriminate financially.
against an employee who enrolls in an
HMO. A contribution does not discrimi-
nate financially if the method of deter-
mimming the contribution is reasonable
and is designed to ensure that employ-
ees have a fair choice among health
benefits plan alternatives.

(2) Effect of agreements or contracts.
The employing entity or designee is
not required to pay more for health
benefits as a result of offering the HMO
alternative than it would otherwise be
required to pay under a collective barg-
ing agreement or contract that
provides for health benefits and is in
effect at the time the HMO alternative
is included.

(3) Examples of acceptable employer
contributions. The following are meth-
ods that are considered nondiscrim-
inatory:
(i) The employer contribution to the
HMO is the same, per employee, as the
contribution to non-HMO alternatives.
(ii) The employer contribution re-
ffects the composition of the HMO's en-
rollment in terms of enrollee at-
tributes that can reasonably be used to
predict utilization, experience, costs,
or risk. For each enrollee in a given
class established on the basis of those
attributes, the employer contributes
an equal amount, regardless of the
health benefits plan chosen by the em-
ployee.
(iii) The employer contribution is a
fixed percentage of the premium for
each of the alternatives offered.
(iv) The employer contribution is de-
termined under a mutually acceptable
arrangement negotiated by the HMO
and the employer. In negotiating the
arrangement, the employer may not in-
sist on terms that would cause the
HMO to violate any of the require-
ments of this part.

(4) Adjustment of employer contribu-
tion. An employer contribution deter-
mimed by an acceptable method may in
some cases be adjusted if it would re-
sult in a nominal payment or no pay-
ment at all by HMO enrollees (because
the HMO premium is lower than the
premiums for the other alternatives of-
fered). If, for example, the employer has
a policy of requiring all employees to
contribute to their health benefits plan,
the employer may require HMO
enrollees who would otherwise pay lit-
tle or nothing at all, to make a pay-
ment that does not exceed 50 percent
of the employee contribution to the prin-
cipal non-HMO alternative. The prin-
cipal non-HMO alternative is the one
that covers the largest number of en-
rollees from the particular employer.

(b) Administrative expenses. (1) In de-
termining the amount of its contribu-
tion to the HMO, the employing entity
or designee may not consider adminis-
trative expenses incurred in connection
with offering any alternative in the
health benefits plan.

(2) However, if the employing entity
or designee has special requirements
for other than standard solicitation
brochures and enrollment literature, it
must, in the case of the HMO alter-
native, determine and distribute any
administrative costs attributable to
those requirements in a manner con-
sistent with its method of determining
and distributing those costs for the
non-HMO alternatives.

(c) Exclusion for contribution for cer-
tain benefits. In determining the
amount of the employing entity's con-
tribution or the designee's cost for the
HMO alternative, the employing entity
or designee may exclude those portions
of the contribution allocable to bene-
fits (such as life insurance or insurance
for supplemental health benefits)—
(1) For which eligible employees and
their eligible dependents are covered
notwithstanding selection of the HMO
alternative; and
(2) That are not offered on a prepay-
ment basis by the HMO to the employ-
ing entity's employees.

(d) Contributions determined by agree-
ments or contracts or by law. If the spe-
cific amount of the employing entity's
contribution for health benefits is fixed
by an agreement or contract, or by
law, that amount constitutes the em-
ploying entity's obligation for con-
tribution toward the HMO premiums.

(e) Allocation of portion of a contribu-
tion determined by an agreement. In some
cases, the employing entity's contribu-
tion for health benefits is determined
by an agreement that also provides for
benefits other than health benefits. In
that case, the employing entity must
determine, or instruct its designee to
§ 417.158 Payroll deductions.

Each employing entity that provides payroll deductions as a means of paying employees' contributions for health benefits or provides a health benefits plan that does not require an employee contribution must, with the consent of an employee who selects the HMO option, arrange for the employee's contribution, if any, to be paid through payroll deductions.

§ 417.159 Relationship of section 1310 of the Public Health Service Act to the National Labor Relations Act and the Railway Labor Act.

The decision of an employing entity subject to this subpart to include the HMO alternative in any health benefits plan offered to its eligible employees must be carried out consistently with the obligations imposed on that employing entity under the National Labor Relations Act, the Railway Labor Act, and other laws of similar effect.

Subpart F—Continued Regulation of Federally Qualified Health Maintenance Organizations

§ 417.160 Applicability.

This subpart applies to any entity that has been determined to be a qualified HMO under subpart D of this part.

§ 417.161 Compliance with assurances.

Any entity subject to this subpart must comply with the assurances that it gave under §417.166.

§ 417.162 Reporting requirements.

Entities subject to this subpart must submit:

(a) The reports that may be required by HCFA under §417.126, and

(b) Any additional reports HCFA may reasonably require.

§ 417.163 Enforcement procedures.

(a) Complaints. Any person, group, association, corporation, or other entity may file with HCFA a written complaint with respect to an HMO's compliance with assurances it gave under subpart D of this part. A complaint must—

(1) State the grounds and underlying facts of the complaint;

(2) Give the names of all persons involved; and

(3) Assure that all appropriate grievance and appeals procedures established by the HMO and available to the complainant have been exhausted.

(b) Investigations. (1) HCFA may initiate investigations when, based on a report, a complaint, or any other information, HCFA has reason to believe that a Federally qualified HMO is not in compliance with any of the assurances it gave under subpart D of this part.

(2) When HCFA initiates an investigation, it gives the HMO written notice that includes a full statement of
the pertinent facts and of the matters being investigated and indicates that the HMO may submit, within 30 days of the date of the notice, a written report concerning these matters.

(3) HCFA obtains any information it considers necessary to resolve issues related to the assurances, and may use site visits, public hearings, or any other procedures that HCFA considers appropriate in seeking this information.

(c) Determination and notice by HCFA—(1) Determination. (i) On the basis of the investigation, HCFA determines whether the HMO has failed to comply with any of the assurances it gave under subpart D of this part.

(ii) HCFA publishes in the Federal Register a notice of each determination of non-compliance.

(2) Notice of determination: Corrective action. (i) HCFA gives the HMO written notice of the determination.

(ii) The notice specifies the manner in which the HMO has not complied with its assurances and directs the HMO to initiate the corrective action that HCFA considers necessary to bring the HMO into compliance.

(iii) The HMO must initiate this corrective action within 30 days of the date of the notice from HCFA, or within any longer period that HCFA determines to be reasonable and specifies in the notice. The HMO must carry out the corrective action within the time period specified by HCFA in the notice.

(iv) The notice may provide the HMO an opportunity to submit, for HCFA’s approval, proposed methods for achieving compliance.

(d) Remedy: Revocation of qualification. If HCFA determines that a qualified HMO has failed to initiate or to carry out corrective action in accordance with paragraph (c)(2) of this section—

(1) HCFA revokes the HMO’s qualification and notifies the HMO of this action.

(2) In the notice, HCFA provides the HMO with an opportunity for reconsideration of the revocation, including, at the HMO’s election, a fair hearing.

(3) The revocation of qualification is effective on the tenth calendar day after the day of the notice unless HCFA receives a request for reconsideration by that date.

(4) If after reconsideration HCFA again determines to revoke the HMO’s qualification, this revocation is effective on the tenth calendar day after the date of the notice of reconsidered determination.

(5) HCFA publishes in the Federal Register each determination it makes under this paragraph (d).

(6) A revocation under this paragraph (d) has the effect described in §417.164.

(e) Notice by the HMO. Within 15 days after the date HCFA issues a notice of revocation, the HMO must prepare a notice that explains, in readily understandable language, the reasons for the determination that it is not a qualified HMO, and send the notice to the following:

(1) The HMO’s enrollees.

(2) Each employer or public entity that has offered enrollment in the HMO in accordance with subpart E of this part.

(3) Each lawfully recognized collective bargaining representative or other representative of the employees of the employer or public entity.

(f) Reimbursement of enrollees for services improperly denied, or for charges improperly imposed. (1) If HCFA determines, under paragraph (c)(1) of this section, that an HMO is out of compliance, HCFA may require the HMO to reimburse its enrollees for the following—

(i) Expenses for basic or supplemental health services that the enrollee obtained from other sources because the HMO failed to provide or arrange for them in accordance with its assurances.

(ii) Any amounts the HMO charged the enrollee that are inconsistent with its assurances. (Rules applicable to charges for all enrollees are set forth in §§417.104 and 417.105. The additional rules applicable to Medicare enrollees are in §415.454.)

(2) This paragraph applies regardless of when the HMO failed to comply with the appropriate assurances.

(g) Remedy: Civil suit—(1) Applicability. This paragraph applies to any HMO or other entity to which a grant, loan, or loan guarantee was awarded, as set forth in subpart V of this part, on the basis of its assurances regarding
§ 417.164 Effect of revocation of qualification on inclusion in employee's health benefit plans.

When an HMO's qualification is revoked under §417.163(d), the following rules apply:

(a) The HMO may not seek inclusion in employees health benefits plans under subpart E of this part.

(b) Inclusion of the HMO in an employer's health benefit plan—

(1) Is disregarded in determining whether the employer is subject to the requirements of subpart E of this part; and

(2) Does not constitute compliance with subpart E of this part by the employer.


§ 417.165 Reapplication for qualification.

An entity whose qualification as an HMO has been revoked by HCFA for purposes of section 1330 of the PHS Act may, after completing the corrective action required under §417.163(c)(2), reapply for a determination of qualification in accordance with the procedures specified in subpart D of this part.


§ 417.166 Waiver of assurances.

(a) General rule. HCFA may release an HMO from compliance with any assurances the HMO gives under subpart D of this part if—

(1) The qualification requirements are changed by Federal law; or

(2) The HMO shows good cause, consistent with the purposes of title XIII of the PHS Act.

(b) Basis for finding of good cause. (1) Grounds upon which HCFA may find good cause include but are not limited to the following:

(i) The HMO has filed for reorganization under Federal bankruptcy provisions and the reorganization can only be approved with the waiver of the assurances.

(ii) State laws governing the entity have been changed after it signed the assurances so as to prohibit the HMO from being organized and operated in a manner consistent with the signed assurances.

(2) Changes in State laws do not constitute good cause to the extent that the changes are preempted by Federal law under section 1311 of the PHS Act.

(c) Consequences of waiver. If HCFA waives any assurances regarding compliance with section 1301 of the PHS Act, HCFA concurrently revokes the HMO's qualification unless the waiver is based on paragraph (a)(1) of this section.


Subparts G—[Reserved]

Subpart J—Qualifying Conditions for Medicare Contracts

SOURCE: 50 FR 1346, Jan. 10, 1985, unless otherwise noted.

§ 417.400 Basis and scope.

(a) Statutory basis. The regulations in this subpart implement section 1876 of the Act, which authorizes Medicare payment to HMOs and CMPs that contract with HCFA to furnish covered services to Medicare beneficiaries.

(b) Scope. (1) This subpart sets forth the requirements an HMO or CMP must meet in order to enter into a contract with HCFA under section 1876 of the Act. It also specifies the procedures that HCFA follows to evaluate applications and make determinations.

(2) The rules for payment to HMOs and CMPs are set forth in subparts N, O, and P of this part.
(3) The rules for HCPP participation in Medicare under section 1833(a)(1)(A) of the Act are set forth in subpart U of this part.

§ 417.401 Definitions.

As used in this subpart and subparts K through R of this part, unless the context indicates otherwise—

Adjusted average per capita cost (AAPCC) means an actuarial estimate made by HCFA in advance of an HMO’s or CMP’s contract period that represents what the average per capita cost to the Medicare program would be for each class of the HMO’s or CMP’s Medicare enrollees if they had received covered services other than through the HMO or CMP in the same geographic area or in a similar area.

Adjusted community rate (ACR) is the equivalent of the premium that a risk HMO or CMP would charge Medicare enrollees independently of Medicare payments if the HMO or CMP used the same rates it charges non-Medicare enrollees for a benefit package limited to covered Medicare services.

Arrangement means a written agreement between an HMO or CMP and another entity, under which—

(1) The other entity agrees to furnish specified services to the HMO’s or CMP’s Medicare enrollees;

(2) The HMO or CMP retains responsibility for the services; and

(3) Medicare payment to the HMO or CMP discharges the beneficiary’s obligation to pay for the services.

Benefit stabilization fund means a fund established by HCFA, at the request of a risk HMO or CMP, to withhold a portion of the per capita payments available to the HMO or CMP and pay that portion in a subsequent contract period for the purpose of stabilizing fluctuations in the availability of the additional benefits the HMO or CMP provides to its Medicare enrollees.

Cost contract means a Medicare contract under which HCFA pays the HMO or CMP on a reasonable cost basis.

Cost HMO or CMP means an HMO or CMP that has in effect a cost contract with HCFA under section 1876 of the Act and subpart L of this part.

Demonstration project means a demonstration project under section 402 of the Social Security Amendments of 1967 (42 U.S.C. 1395b–1) or section 222(a) of the Social Security Amendments of 1972 (42 U.S.C. 1395b–1 (note)), relating to the provision of services for which payment is made under Medicare on a prospectively determined basis.

Emergency services means covered inpatient or outpatient services that are furnished by an appropriate source other than the HMO or CMP and that meet the following conditions:

(1) Are needed immediately because of an injury or sudden illness.

(2) Are such that the time required to reach the HMO’s or CMP’s providers or suppliers (or alternatives authorized by the HMO or CMP) would mean risk of permanent damage to the enrollee’s health.

Once initiated, the services continue to be considered emergency services as long as transfer of the enrollee to the HMO’s or CMP’s source of health care or authorized alternative is precluded because of risk to the enrollee’s health or because transfer would be unreasonable, given the distance and the nature of the medical condition.

Geographic area means the area found by HCFA to be the area within which the HMO or CMP furnishes, or arranges for furnishing, the full range of services that it offers to its Medicare enrollees.

Medicare enrollee means a Medicare beneficiary who has been identified on HCFA records as an enrollee of an HMO orCMP that has a contract with HCFA under section 1876 of the Act and subpart L of this part.

New Medicare enrollee means a Medicare beneficiary who—

(1) Enrolls with an HMO or CMP after the date on which the HMO or CMP first enters into a risk contract under subpart L of this part; and

(2) Was not enrolled with the HMO or CMP at the time he or she became entitled to benefits under Part A or eligible to enroll in Part B of Medicare.

Risk contract means a Medicare contract under which HCFA pays the HMO or CMP on a risk basis for Medicare covered services.

Risk HMO or CMP means an HMO or CMP that has in effect a risk contract with HCFA under section 1876 of the Act and subpart L of this part.
§ 417.402 Urgently needed services

Urgently needed services means covered services that are needed by an enrollee who is temporarily absent from the HMO’s or CMP’s geographic area and that—

(1) Are required in order to prevent serious deterioration of the enrollee’s health as a result of unforeseen injury or illness; and

(2) Cannot be delayed until the enrollee returns to the HMO’s or CMP’s geographic area.


§ 417.403 Effective date of initial regulations.

(a) The changes made to section 1876 of the Act by section 114 of the Tax Equity and Fiscal Responsibility Act of 1982 became effective on February 1, 1985, the effective date of the initial implementing regulations.

(b) The changes made to section 1876 of the Act by section 4002 of the Balanced Budget Act (BBA) of 1997 are incorporated in section 422 except for 1876 cost contracts. Upon enactment of the BBA (August 5, 1997) no new cost contracts or service area expansions are accepted by HCFA except for current Health Care Prepayment Plans that may convert to 1876 cost contracts. Also, 1876 cost contracts may not be extended or renewed beyond December 31, 2004.


§ 417.404 General requirements.

(a) In order to contract with HCFA under the Medicare program, an entity must—

(1) Be determined by HCFA to be an HMO or CMP (in accordance with §§117.142 and 417.407, respectively); and

(2) Comply with the contract requirements set forth in subpart L of this part.

(b) HCFA enters into or renews a contract only if it determines that action would be consistent with the effective and efficient implementation of section 1876 of the Act.

[60 FR 45675, Sept. 1, 1995]

§ 417.406 Application and determination.

(a) Responsibility for making determinations. HCFA is responsible for determining whether an entity meets the requirements to be an HMO or CMP.

(b) Application requirements. (1) The application requirements for HMOs are set forth in §417.143.

(2) The requirements of §417.143 also apply to CMPs except that there are no application fees.

(c) Determination. HCFA uses the procedures set forth in §417.144(a) through (d) to determine whether an entity is an HMO or CMP.

(d) Oversight of continuing compliance. (1) HCFA oversees an entity’s continued compliance with the requirements for an HMO as defined in §417.1 or for a CMP as set forth in §417.407.

(2) If an entity no longer meets those requirements, HCFA terminates the contract of that entity in accordance with §417.494.

[60 FR 45675, Sept. 1, 1995]

§ 417.407 Requirements for a Competitive Medical Plan (CMP).

(a) General rule. To qualify as a CMP, an entity must be organized under the laws of a State and must meet the requirements of paragraphs (b) through (f) of this section.

(b) Required services. (1) Basic rule. Except as provided in paragraph (b)(2) of this section, the entity furnishes to its enrollees at least the following services:

(i) Physicians’ services performed by physicians.

(ii) Laboratory, x-ray, emergency, and preventive services.

(iii) Out-of-area coverage.

(iv) Inpatient hospital services.

(2) Exception for Medicaid prepayment risk contracts. An entity that had, before 1970, a Medicaid prepayment risk contract that did not include provision of inpatient hospital services is not required to provide those services.

(c) Compensation for services. The entity receives compensation (except for deductibles, coinsurance, and copayments) for the health care services it provides to enrollees on a periodic, prepaid capitation basis regardless of the
frequency, extent, or kind of services provided to any enrollee.

d) Source of physicians' services. The entity provides physicians' services primarily through—

(1) Physicians who are employees or partners of the entity; or

(2) Physicians or groups of physicians (organized on a group or individual practice basis) under contract with the entity to provide physicians' services.

e) Assumption of financial risk. The rules set forth in §417.120(b) for HMOs apply also to CMPs except that reference to "basic services" must be read as reference to the required services listed in paragraph (b) of this section.

(f) Protection of enrollees. The entity provides adequately against the risk of insolvency by meeting the requirements of §§417.120(a) and 417.122 for protection of enrollees against loss of benefits and liability for payment of any fees that are the legal responsibility of the entity.

§417.408 Contract application process.

(a) Contents of application. (1) The application for a contract must include supporting information in the form and detail required by HCFA. (2) Whenever feasible, HCFA exempts the HMO or CMP from resubmittal of information it has already submitted to HCFA in connection with a determination made under the provisions of §417.406.

(b) Approval of application. (1) If HCFA approves the application, it gives written notice to the HMO or CMP, indicating that it meets the requirements for either a risk or reasonable cost contract or only for a reasonable cost contract.

(2) If the HMO or CMP is dissatisfied with a determination that it meets the requirements only for a reasonable cost contract, it may request reconsideration in accordance with the procedures specified in subpart R of this part.

(c) Denial of application. If HCFA denies the application, it gives written notice to the HMO or CMP indicating—

(1) That it does not meet the contract requirements under section 1876 of the Act;

(2) The reasons why the HMO or CMP does not meet the contract requirements; and

(3) The HMO's or CMP's right to request reconsideration in accordance with the procedures specified in subpart R of this part.

§417.410 Qualifying conditions: General rules.

(a) Basic requirement. In order to qualify for a contract with HCFA under this subpart, an HMO or CMP must demonstrate its ability to enroll Medicare beneficiaries and other individuals and groups and to deliver a specified comprehensive range of high quality services efficiently, effectively, and economically to its Medicare enrollees.

(b) Other qualifying conditions. An HMO or CMP must meet qualifying conditions that pertain to operating experience, enrollment, range of services, furnishing of services, and a quality assurance program.

(c) Standards. Generally, each qualifying condition is interpreted by a series of standards that are used in surveying an HMO or CMP to determine its qualifications for a Medicare contract.

(d) Application of standards. Application of the standards enables the surveyor to determine—

(1) The HMO's or CMP's activities;

(2) The extent to which the HMO or CMP complies with each condition;

(3) The nature and extent of any deficiencies; and

(4) The need for improvement if HCFA should enter into a contract with the HMO or CMP.

(e) Requirements for a risk contract. An HMO or CMP may enter into a risk contract with HCFA if it—

(1) Meets all the applicable requirements in the statute and regulations;

(2) Has at least 5,000 enrollees or 1,500 enrollees if it serves a primarily rural area as defined in §417.413(b)(3); and

(3) Has at least 75 Medicare enrollees or has an acceptable plan to achieve this Medicare membership within 2 years;
§ 417.412 Qualifying condition: Administration and management.

The HMO or CMP must demonstrate that it—
(a) Has sufficient administrative capability to carry out the requirements of the contract; and
(b) Does not have any agents or management staff or persons with ownership or control interests who have been convicted of criminal offenses related to their involvement in Medicaid, Medicare, or social service programs under title XX of the Act.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38078, July 15, 1993; 60 FR 45676, Sept. 1, 1995]

§ 417.413 Qualifying condition: Operating experience and enrollment.

(a) Condition. The HMO or CMP must demonstrate that it has operating experience and an enrolled population sufficient to provide a reasonable basis for establishing a prospective per capita reimbursement rate or a reasonable cost reimbursement rate, as appropriate.

(b) Standard: Enrollment and operating experience for HMOs or CMPs to contract on a risk basis. To be eligible to contract on a risk basis—
(1) A nonrural HMO or CMP must currently have the following:
   (i) At least 5,000 enrollees; and
   (ii) At least 75 Medicare enrollees or a plan acceptable to HCFA for achieving a Medicare enrollment of 75 within 2 years from the beginning of its initial contract period.
(2) A rural HMO or CMP must currently have—
   (i) At least 1,500 enrollees; and
   (ii) At least 75 Medicare enrollees or a plan acceptable to HCFA for achieving a Medicare enrollment of 75 within 2 years from the beginning of its initial contract period.
(3) For purposes of this paragraph, an HMO or CMP is considered rural if at least 50 percent of its enrollees reside in nonmetropolitan areas. A nonmetropolitan area is an area—
   (i) No part of which is within a metropolitan statistical area (MSA) as designated by the Executive Office of Management and Budget; and
   (ii) That does not contain a city whose population exceeds 50,000 individuals.
(4) A subdivision or subsidiary of an HMO or CMP that meets the requirements of paragraph (b)(1) or (b)(2) of this section need not demonstrate that it meets those requirements as an independent unit if the HMO or CMP assumes responsibility for the financial risk, and adequate management and supervision of health care services furnished by its subdivision or subsidiary.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38078, July 15, 1993; 60 FR 45676, Sept. 1, 1995]
(d) Standard: Composition of enrollment—(1) Requirement. Except as specified in paragraphs (d)(2) and (e) of this section, not more than 50 percent of an HMO’s or CMP’s enrollment may be Medicare beneficiaries.

(2) Waiver of composition of enrollment standard. HCFA may waive compliance with the requirements of paragraph (d)(1) of this section if the HMO or CMP has made and is making reasonable efforts to enroll individuals who are not Medicare beneficiaries and it meets one of the following requirements:

(i) The HMO or CMP serves a geographic area in which Medicare beneficiaries and Medicaid recipients constitute more than 50 percent of the population. (HCFA does not grant a waiver that would permit the percentage of Medicare and Medicaid enrollees to exceed the percentage of Medicare beneficiaries and Medicaid recipients in the population of the geographic area.)

(ii) The HMO or CMP is owned and operated by a government entity. The waiver may be for a period up to three years after the date the HMO or CMP first enters into a contract under this subpart, and may not be extended.

(iii) The HMO or CMP requests waiver of the composition rule because it is in the public interest. The organization provides documentation that supports one of the following:

(A) The organization serves a medically underserved rural or urban area.

(B) The organization demonstrates a long-term business and community service commitment to the area.

(C) The organization believes that a waiver is necessary to promote managed care choices in an area with limited or no managed care choices.

(3) Waiver granted on or before October 21, 1986. An HMO or CMP (or a successor HMO or CMP) that as of October 21, 1986, had been granted an exception, waiver, or modification of the requirements of paragraph (d)(1) of this section, must make and throughout the period of the exception, waiver, or modification continue to make reasonable efforts to meet scheduled enrollment goals, consistent with a schedule of compliance approved by HCFA.

(i) If HCFA determines that the HMO or CMP has complied, or made significant progress toward compliance, with the approved schedule, and that an extension is in the best interest of the Medicare program, HCFA may extend the waiver of modification.

(ii) If HCFA determines that the HMO or CMP has not complied with the approved schedule, HCFA may apply the sanctions described in paragraphs (d)(6) and (d)(7) of this section, as applicable.

(4) Basis for application of sanctions. HCFA may, as an alternative to contract termination, apply the sanctions specified in paragraph (d)(6) of this section if HCFA determines that the HMO or CMP is not complying with the requirements in paragraphs (d)(1), (d)(2), or (d)(3) of this section, as applicable.

(5) Notice of sanction. Before applying the sanctions specified in paragraph (d)(6) of this section, HCFA sends a written notice to the HMO or CMP stating the proposed action and its basis. HCFA gives the HMO or CMP 15 days after the date of the notice to provide evidence establishing the HMO’s or CMP’s compliance with the requirements in paragraph (d)(1), (d)(2), or (d)(3) of this section, as applicable.

(6) Sanctions. If, following review of the HMO’s or CMP’s timely response to HCFA’s notice, HCFA determines that an HMO or CMP does not comply with the requirements of paragraphs (d)(1), (d)(2), or (d)(3) of this section, HCFA may apply either of the following sanctions:

(i) Require the HMO or CMP to stop accepting new enrollment applications after a date specified by HCFA.

(ii) Deny payment for individuals who are formally added or “accreted” to HCFA’s records as Medicare enrollees after a date specified by HCFA.

(7) Termination by HCFA. In addition to the sanctions described in paragraph (d)(6) of this section, HCFA may decline to renew an HMO’s or CMP’s contract in accordance with §417.492(b), or terminate its contract in accordance with §417.494(b) if HCFA determines that the HMO or CMP no longer substantially meets the requirements of paragraphs (d)(1), (d)(2), or (d)(3) of this section.
§ 417.414 Qualifying condition: Range of services.

(a) Condition. The HMO or CMP must demonstrate that it is capable of delivering to Medicare enrollees the range of services required in accordance with this section.

(b) Standard: Range of services furnished by eligible HMOs or CMPs. (1) Basic requirement. Except as specified in paragraph (b)(3) of this section, an HMO or CMP must furnish to its Medicare enrollees (directly or through arrangements with others) all the Medicare services to which those enrollees are entitled to the extent that they are available to Medicare beneficiaries who reside in the HMO’s or CMP’s geographic area but are not enrolled in the HMO or CMP.

(2) Criteria for availability. The services are considered available if—

(i) The sources are located within the HMO’s or CMP’s geographic area; or

(ii) It is common practice to refer patients to sources outside that geographic area.

(3) Exception for hospice care. An HMO or CMP is not required to furnish hospice care as described in part 418 of this chapter. However, HMOs or CMPs must inform their Medicare enrollees about the availability of hospice care if—

(i) A hospice participating in Medicare is located within the HMO’s or CMP’s geographic area; or

(ii) It is common practice to refer patients to hospices outside the geographic area.

(c) Standard: Financial responsibility for services furnished outside the HMO or CMP. (1) An HMO or CMP must assume financial responsibility and provide reasonable reimbursement for emergency services and urgently needed services (as defined in § 417.401) that are obtained by its Medicare enrollees from providers and suppliers outside the HMO or CMP even in the absence of the HMO’s or CMP’s prior approval.

(2) An HMO or CMP must assume financial responsibility for services that the Medicare enrollee attempted to obtain from the HMO or CMP, but that the HMO or CMP failed to furnish or unreasonably denied, and that are found, upon appeal by the enrollee under subpart Q of this part, to be services that the enrollee was entitled to have furnished to him or her by the HMO or CMP.

§ 417.416 Qualifying condition: Furnishing of services.

(a) Condition. The HMO or CMP must furnish the required services to its Medicare enrollees through providers and suppliers that meet applicable Medicare statutory definitions and implementing regulations. The HMO or CMP must also ensure that the required services, additional services, and any other supplemental services for which the Medicare enrollee has contracted are available and accessible.
and are furnished in a manner that ensures continuity.

(b) Standard: Conformance with conditions of participation, conditions for coverage, and conditions for certification. (1) Hospitals, SNFs, HHAs, CORFs, and providers of outpatient physical therapy or speech-language pathology services must meet the applicable conditions of participation in Medicare, as set forth elsewhere in this chapter.

(2) Suppliers must meet the conditions for coverage or conditions for certification of their services, as set forth elsewhere in this chapter.

(3) If more than one type of practitioner is qualified to furnish a particular service, the HMO or CMP may select the type of practitioner to be used.

(c) Standard: Physician supervision. The HMO or CMP must provide for supervision by a physician of other health care professionals who are directly involved in the provision of health care as generally authorized under section 1861 of the Act. Except as specified in paragraph (d) of this section, with respect to medical services furnished in an HMO's or CMP's clinic or the office of a physician with whom the HMO or CMP has a service agreement, the HMO or CMP must ensure that—

(1) Services furnished by paramedical, ancillary, and other nonphysician personnel are furnished under the direct supervision of a physician;

(2) A physician is present to perform medical (as opposed to administrative) services whenever the clinics or offices are open; and

(3) Each patient is under the care of a physician.

(d) Exceptions to physician supervision requirement. The following services may be furnished without the direct personal supervision of a physician:

(1) Services of physician assistants and nurse practitioners (as defined in §401.2 of this chapter), and the services and supplies incident to their services. The conditions for payment, as set forth in §§405.2414 and 405.2415 of this chapter for services furnished by rural health clinics and Federally qualified health centers, respectively, also apply when those services are furnished by an HMO or CMP.

(2) When furnished by an HMO or CMP, services of clinical psychologists who meet the qualifications specified in §410.71(d) of this chapter, and the services and supplies incident to their professional services.

(3) When an HMO or CMP contracts on—

(i) A risk basis, the services of a clinical social worker (as defined at §410.73 of this chapter) and the services and supplies incident to their professional services; or

(ii) A cost basis, the services of a clinical social worker (as defined in §410.73 of this chapter). Services incident to the professional services of a clinical social worker furnished by an HMO or CMP contracting on a cost basis are not covered by Medicare and payment will not be made for these services.

(e) Standard: Accessibility and continuity. (1) The HMO or CMP must ensure that the required services and any other services for which Medicare enrollees have contracted are accessible, with reasonable promptness, to the enrollees with respect to geographic location, hours of operation, and provision of after hours service. Medically necessary emergency services must be available twenty-four hours a day, seven days a week.

(2) The HMO or CMP must maintain a health (including medical) record-keeping system through which pertinent information relating to the health care of its Medicare enrollees is accumulated and is readily available to appropriate professionals.


§ 417.418 Qualifying condition: Quality assurance program.

(a) Condition. The HMO or CMP must make arrangements for a quality assurance program that meets the requirements of this section.

(b) Standard. An HMO or CMP must have an ongoing quality assurance program that meets the requirements set forth in §417.106(a).

[58 FR 38072, July 15, 1993]
§ 417.420 Basic rules on enrollment and entitlement.

(a) Enrollment. Individuals who are entitled to benefits under both Part A and Part B of Medicare or only Part B may elect to receive those benefits through an HMO or CMP that has in effect a contract with HCFA under subpart L of this part.

(b) Entitlement. If a Medicare beneficiary enrolls with an HMO or CMP, HCFA pays the HMO or CMP on his or her behalf for the services to which he or she is entitled.

(c) Beneficiary liability. (1) The HMO or CMP may require payment, in the form of premiums or otherwise, from individuals for services not covered under Medicare, as well as deductible and coinsurance amounts attributable to Medicare covered services.

(2) As described in §417.448, Medicare enrollees of risk HMOs or CMPs are liable for services that they obtain from sources other than the HMO or CMP, unless the services are—
   (i) Emergency or urgently needed; or
   (ii) Determined, on appeal under subpart Q of this part, to be services that should have been furnished by the HMO or CMP.


§ 417.422 Eligibility to enroll in an HMO or CMP.

Except as specified in §§417.423 and 417.424, an HMO or CMP must enroll, either for an indefinite period or for a specified period of at least 12 months, any individual who—

(a) Is entitled to Medicare benefits under Parts A and B or under Part B only;

(b) Lives within the geographic area served by the HMO or CMP;

(c) Is not enrolled in any other HMO or CMP that has entered into a contract under subpart L of this part;

(d) During an enrollment period of the HMO or CMP, completes and signs the HMO’s or CMP’s application form and gives whatever information is required for enrollment;

(e) Agrees to abide by the HMO’s or CMP’s rules after they are disclosed to him or her in connection with the enrollment process;

(f) Is not denied enrollment by the HMO or CMP under a selection policy, if any, that has been approved by HCFA under §417.424(b); and

(g) Is not denied enrollment by the HMO or CMP on the basis of any of the administrative criteria concerning denial of enrollment in §417.424(a).


§ 417.423 Special rules: ESRD and hospice patients.

(a) ESRD patients. (1) A Medicare beneficiary who has been medically determined to have end-stage renal disease is not eligible to enroll in an HMO or CMP.

   (2) However, if a beneficiary is already enrolled in an HMO or CMP when he or she is determined to have end-stage renal disease, the HMO or CMP—
   (i) Must reenroll the beneficiary as required by §417.434; and
   (ii) May not disenroll the beneficiary except as provided in §417.460.

(b) Hospice patients. A Medicare beneficiary who elects hospice care under §418.24 of this chapter is not eligible to enroll in an HMO or CMP as long as the hospice election remains in effect.

[60 FR 45677, Sept. 1, 1995]

§ 417.424 Denial of enrollment.

(a) Basis for denial. An HMO or CMP may deny enrollment to an individual who meets the criteria of §417.422 if acceptance would—

   (1) Cause the number of enrollees who are Medicare or Medicaid beneficiaries to exceed 50 percent of the HMO’s or CMP’s total enrollment;

   (2) Prevent the HMO or CMP from complying with any of the other contract qualifying conditions set forth in subpart J of this part;

   (3) Require the HMO or CMP to exceed its enrollment capacity; or

   (4) Cause the enrollment to become substantially nonrepresentative of the
§ 417.428 Marketing activities.

(a) Required marketing activities. An HMO or CMP must meet the following requirements:

(1) Offer its plan to Medicare beneficiaries and provide to those interested in enrolling, adequate written descriptions of the HMO’s or CMP’s rules, procedures, benefits, fees, and other charges, services, and information necessary for beneficiaries to make an informed decision about enrollment.

(2) Notify the general public of its enrollment period (whether time limited or continuous) in an appropriate manner through appropriate media, throughout its enrollment area.

(3) Submit all marketing materials to HCFA at least 45 days before their planned distribution.

(4) Include in the HMO’s or CMP’s written materials provided to prospective enrollees prior to enrollment, notice that the HMO or CMP is authorized by law to terminate or refuse to renew its contract with HCFA, that HCFA may also choose to terminate or refuse to renew its contact with the HMO or CMP and that termination or nonrenewal may result in termination of the individual’s enrollment in the HMO or CMP.

§ 417.426 Open enrollment requirements.

(a) Basic requirements. (1) HMOs or CMPs must provide open enrollment for Medicare beneficiaries for at least 30 consecutive days during each contract year.

(2) During open enrollment, the HMO or CMP must enroll eligible Medicare beneficiaries in the order in which their applications are received and until its enrollment capacity is reached.

(3) The HMO or CMP may accept applications from Medicare beneficiaries after it has reached capacity if it places those individuals on a waiting list and enrolls them in chronological order as vacancies occur.

(4) An HMO or CMP with a risk contract must accept applications from eligible Medicare beneficiaries during the month of November 1996.

(b) Capacity to accept new enrollees. (1) If an HMO or CMP chooses to limit enrollments because of its capacity, it must notify HCFA at least 90 days before the beginning of its open enrollment period and, at that time, provide HCFA with its reasons for limiting enrollment.

(2) HCFA evaluates the HMO’s or CMP’s submittal under paragraph (b)(1) of this section.

(3) The HMO or CMP must promptly notify HCFA if there is any change in its enrollment capacity.

(c) Reserved vacancies. (1) Subject to HCFA’s approval, an HMO or CMP may set aside a reasonable number of vacancies for an anticipated new group contract or for anticipated new enrollees under an existing group contract that will have its enrollment period after the Medicare open enrollment period during the contract year.

(2) Any set aside vacancies that are not filled within a reasonable time after the beginning of the group contract enrollment period must be made available to Medicare beneficiaries and other nongroup applicants under the requirements of this subpart.

§ 417.428 Marketing activities.

(a) Required marketing activities. An HMO or CMP must meet the following requirements:

(1) Offer its plan to Medicare beneficiaries and provide to those interested in enrolling, adequate written descriptions of the HMO’s or CMP’s rules, procedures, benefits, fees, and other charges, services, and other information necessary for beneficiaries to make an informed decision about enrollment.

(2) Notify the general public of its enrollment period (whether time limited or continuous) in an appropriate manner through appropriate media, throughout its enrollment area.

(3) Submit all marketing materials to HCFA at least 45 days before their planned distribution.

(4) Include in the HMO’s or CMP’s written materials provided to prospective enrollees prior to enrollment, notice that the HMO or CMP is authorized by law to terminate or refuse to renew its contract with HCFA, that HCFA may also choose to terminate or refuse to renew its contact with the HMO or CMP and that termination or nonrenewal may result in termination of the individual’s enrollment in the HMO or CMP.
§ 417.430 Application procedures.

(a) Application forms. (1) The application form must comply with HCFA instructions regarding format and content and must include the beneficiary's signature and authorization for disclosure and exchange of necessary information between HCFA and the HMO or CMP.

(2) The HMO or CMP must file and retain application forms for the period specified in HCFA instructions.

(b) Handling of applications. An HMO or CMP must have an effective system for receiving, controlling, and processing applications from Medicare beneficiaries. The system must meet the following conditions and requirements:

(1) Each application is dated as of the date it is received.

(2) Applications are processed in chronological order by date of receipt.

(3) The HMO or CMP gives the beneficiary prompt written notice of acceptance or rejection of the application.

(4) The notice of acceptance—

(i) Specifies the date on which the HMO or CMP will request HCFA to make the enrollment effective; or

(ii) If the HMO or CMP is currently enrolled to capacity, explains the procedures that will be followed when vacancies occur.

(5) The notice of denial explains the reason for denial.

(6) The HMO or CMP transmits the information necessary for HCFA to add the beneficiary to its records of the HMO's or CMP's Medicare enrollees—

(i) Within 30 days from the date of application or from the date a vacancy occurs for an applicant who was accepted (for future enrollment) while there were no vacancies; or

(ii) Within an additional period of time approved by HCFA on a showing by the HMO or CMP that it needs more time.
§ 417.436 Rules for enrollees.

(a) Maintaining rules. An HMO or CMP must maintain written rules that deal with, but need not be limited to the following:

(1) All benefits provided under the contract, as described in §417.440.

(2) How and where to obtain services from or through the HMO or CMP.

(3) The restrictions on coverage for services furnished from sources outside a risk HMO or CMP, other than emergency services and urgently needed services (as defined in §417.401).

(4) The obligation of the HMO or CMP to assume financial responsibility and provide reasonable reimbursement for emergency services and urgently needed services as required by §417.414(c).

(5) Any services other than the emergency or urgently needed services that the HMO or CMP chooses to provide as permitted by this part, from sources outside the HMO or CMP. A cost HMO or CMP must disclose that the enrollee may receive services through any Medicare providers and suppliers.

(6) Premium information, including the amount (or if the amount cannot be included, the telephone number of the source from which this information may be obtained) and the procedures for paying premiums and other charges for which enrollees may be liable.

(7) Grievance and appeal procedures.

(8) Disenrollment rights.
(9) The obligation of an enrollee who is leaving the HMO's or CMP's geographic area for more than 90 days to notify the HMO or CMP of the move or extended absence and the HMO's or CMP's policies concerning retention of enrollees who leave the geographic area for more than 90 days, as described in §417.460(a)(2).

(10) The expiration date of the Medicare contract with HCFA and notice that both HCFA and the HMO or CMP are authorized by law to terminate or refuse to renew the contract, and that termination or nonrenewal of the contract may result in termination of the individual's enrollment in the HMO or CMP.

(11) Advance directives as specified in paragraph (d) of this section.

(12) Any other matters that HCFA may prescribe.

(b) Availability of rules. The HMO or CMP must furnish a copy of the rules to each Medicare enrollee at the time of enrollment and at least annually thereafter.

(c) Changes in rules. If an HMO or CMP changes its rules, it must submit the changes to HCFA in accordance with §417.428(a)(3), and notify its Medicare enrollees of the changes at least 30 days before the effective date of the changes.

(d) Advance directives. (1) An HMO or CMP must maintain written policies and procedures concerning advance directives, as defined in §489.100 of this chapter, with respect to all adult individuals receiving medical care by or through the HMO or CMP and are required to:

(i) Provide written information to those individuals concerning—

(A) The individual's rights under the law of the State in which the organization furnishes services (whether statutory or recognized by the courts of the State) to make decisions concerning such medical care, including the right to accept or refuse medical or surgical treatment and the right to formulate, at the individual's option, advance directives. Providers are permitted to contract with other entities to furnish this information but are still legally responsible for ensuring that the requirements of this section are met. Such information must reflect changes in State law as soon as possible, but no later than 90 days after the effective date of the State law; and

(B) The HMO's or CMP's written policies respecting the implementation of those rights, including a clear and precise statement of limitation if the HMO or CMP cannot implement an advance directive as a matter of conscience. At a minimum, this statement should:

(1) Clarify any differences between institution-wide conscience objections and those that may be raised by individual physicians;

(2) Identify the state legal authority permitting such objection; and

(3) Describe the range of medical conditions or procedures affected by the conscience objection.

(ii) Provide the information specified in paragraphs (d)(1)(i) of this section to each enrollee at the time of initial enrollment. If an enrollee is incapacitated at the time of initial enrollment and is unable to receive information (due to the incapacitating condition or a mental disorder) or articulate whether or not he or she has executed an advance directive, the HMO or CMP may give advance directive information to the enrollee's family or surrogate in the same manner that it issues other materials about policies and procedures to the family of the incapacitated enrollee or to a surrogate or other concerned persons in accordance with State law. The HMO or CMP is not relieved of its obligation to provide this information to the enrollee once he or she is no longer incapacitated or unable to receive such information. Follow-up procedures must be in place to ensure that the information is given to the individual directly at the appropriate time.

(iii) Document in the individual's medical record whether or not the individual has executed an advance directive.

(iv) Not condition the provision of care or otherwise discriminate against an individual based on whether or not the individual has executed an advance directive;

(v) Ensure compliance with requirements of State law (whether statutory or recognized by the courts of the State) regarding advance directives;
(vi) Provide for education of staff concerning its policies and procedures on advance directives; and

(vii) Provide for community education regarding advance directives that may include material required in paragraph (d)(3)(i)(A) of this section, either directly or in concert with other providers or entities. Separate community education materials may be developed and used, at the discretion of the HMO or CMP. The same written materials are not required for all settings, but the material should define what constitutes an advance directive, emphasizing that an advance directive is designed to enhance an incapacitated individual’s control over medical treatment, and describe applicable State law concerning advance directives. An HMO or CMP must be able to document its community education efforts.

(2) The HMO or CMP—

(i) Is not required to provide care that conflicts with an advance directive.

(ii) Is not required to implement an advance directive if, as a matter of conscience, the HMO or CMP cannot implement an advance directive and State law allows any health care provider or any agent of such provider to conscientiously object.

(3) The HMO or CMP must inform individuals that complaints concerning non-compliance with the advance directive requirements may be filed with the State survey and certification agency.


§ 417.440 Entitlement to health care services from an HMO or CMP.

(a) Basic rules. (1) Subject to the conditions and limitations set forth in this subpart, a Medicare enrollee of an HMO or CMP is entitled to receive health care services and supplies directly from, or through arrangements made by, the HMO or CMP as specified in this section and §§417.442-417.446.

(2) A Medicare enrollee is also entitled to receive timely and reasonable payment directly (or have payment made on his or her behalf) for services he or she obtained from a provider or supplier outside the HMO or CMP if those services are—

(i) Emergency services or urgently needed services as defined §417.401.

(ii) Services denied by the HMO or CMP and found (upon appeal under subpart Q of this part) to be services the enrollee was entitled to have furnished by the HMO or CMP.

(b) Scope of services. (1) Part A and Part B services. Except as specified in paragraphs (c), (d), and (e) of this section, a Medicare enrollee is entitled to receive from an HMO or CMP all the Medicare-covered services that are available to individuals residing in the HMO’s or CMP’s geographic area, as follows:

(i) Medicare Part A and Part B services if the enrollee is entitled to benefits under both programs.

(ii) Medicare Part B services if the enrollee is entitled only under that program.

(2) Supplemental services elected by enrollee. A Medicare enrollee of an HMO or CMP may elect to pay for optional services that are offered by the HMO or CMP in addition to the covered Part A and Part B services, and, for risk HMOs or CMPs, in addition to the additional benefits required under §417.442. The HMO or CMP may not set health status standards for those enrollees whom it will accept for these optional supplemental services.

(3) Supplemental services imposed by a risk HMO or CMP. (i) Subject to HCFA’s approval, a risk HMO or CMP may require Medicare enrollees to accept and pay for services in addition to those covered by Medicare. (ii) If the HMO or CMP elects this option, it must impose the requirement on all Medicare enrollees, without regard to health status. (iii) HCFA approves supplemental benefits of this type if HCFA determines that imposition of the requirements will not discourage other Medicare beneficiaries from enrolling in the risk HMO or CMP.

(4) Additional benefits from risk HMOs or CMPs required by statute. Subject to the conditions stated in §417.442, a new Medicare enrollee or a current nonrisk Medicare enrollee who converts to risk reimbursement under §417.444 is eligible to receive, in addition to the covered Part A and Part B benefits for which he or she is eligible, benefits
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consisting of one or both of the following:

(i) A reduction in the HMO’s or CMP’s premium rate or in other charges for services furnished to Medicare enrollees.

(ii) Provision of health benefits or services beyond the required Part A and Part B coverage.

(5) Special supplemental benefits. Under conditions described in § 417.444(c), current nonrisk Medicare enrollees who are not converted to the risk portion of the contract, may enroll in a special supplemental plan, if offered by the HMO or CMP, for some or all of the additional benefits described in paragraph (b)(4) of this section.

(c) Limitation on hospice care. (1) Extent of limitation. (i) Basic rule. Except as provided in paragraph (c)(1)(ii) of this section, a Medicare enrollee who elects to receive hospice care under § 418.24 of this chapter waives the right to receive from the HMO or CMP any Medicare services (including services equivalent to hospice care) that are related to the terminal condition for which the enrollee elected hospice care, or to a related condition.

(ii) Exception. An enrollee who elects hospice care retains the right to services furnished by his or her attending physician if that physician—

(A) Is an employee or contractor of the HMO or CMP; and

(B) Is not an employee of the designated hospice and does not receive compensation from the hospice for those services.

(2) Effective date of limitation. The limitation in paragraph (c)(1) of this section begins on the effective date of the beneficiary’s election of hospice care and remains in effect until the earlier of the following:

(i) The effective date of the enrollee’s revocation of the election of hospice care as described in § 418.28 of this chapter.

(ii) The date the enrollee exhausts his or her hospice benefits.

(3) Payment to HMO or CMP. For the period that the Medicare enrollee’s election of hospice care is in effect, HCFA pays a cost HMO or CMP only as described in § 417.585.

(d) Limitation on provision of inpatient hospital services. If a beneficiary’s effective date of coverage, as specified in § 417.450, in a risk HMO or CMP occurs during an inpatient stay in a hospital paid for under part 412 of this chapter, the HMO or CMP—

(1) Is not responsible for the provision of any of the inpatient hospital services under Part A during the stay and is not required to pay for those services;

(2) Must assume responsibility for payment for or provision of inpatient hospital services under Part A on the day after the day of discharge from the inpatient stay; and

(3) Is responsible for the full scope of services under paragraph (b) of this section, other than inpatient hospital services under Part A, beginning on the effective date of enrollment.

(e) Extension of provision of inpatient hospital services. If an enrollee’s effective date of disenrollment, as defined by § 417.460, occurs during an inpatient stay in a hospital paid for under part 412 of this chapter and the stay is provided or arranged for by the HMO or CMP, or the HMO or CMP is financially responsible for the hospitalization under paragraph (a)(2) of this section, the HMO or CMP—

(1) Is financially responsible for payment of the inpatient services under Part A through the date the beneficiary is discharged from the inpatient stay; and

(2) Is not responsible for the provision of services, furnished on or after the effective date of disenrollment, other than inpatient hospital services under Part A.

(f) Notice of noncoverage of inpatient hospital care. (1) If an enrollee is an inpatient of a hospital, entitlement to inpatient hospital care continues until he or she receives notice of noncoverage of that care.

(2) Before giving notice of noncoverage, the HMO or CMP must obtain the concurrence of its affiliated physician responsible for the hospital care of the enrollee, or other physician as authorized by the HMO or CMP.

(3) The HMO or CMP must give the enrollee written notice that includes the following:

(i) The reason why inpatient hospital care is no longer needed.
§ 417.444 Special rules for certain enrollees of risk HMOs and CMPs.

(a) Applicability. This section applies to any Medicare enrollee of a risk HMO or CMP who meets the following conditions:

(1) On February 1, 1985, was enrolled—

(i) In an HMO or CMP that had in effect a cost contract entered into under section 1876 of the Act in accordance with regulations in effect before February 1, 1985; or

(ii) In an HCPP that was being reimbursed on a reasonable cost basis under section 1833(a)(1)(A) of the Act.

(2) Has continued enrollment in the same entity without interruption or disenrolled after February 1, 1985, and later reenrolled in the same entity.

(b) Retention of nonrisk status—

(1) A “nonrisk” enrollee is a Medicare beneficiary who meets the conditions of paragraph (a) of this section and is enrolled in an entity that enters into a risk contract as an HMO or CMP. A “nonrisk” enrollee may retain nonrisk status indefinitely unless HCFA determines under paragraph (c)(1) of this section, that the enrollee’s status must be changed, or the enrollee requests the change, as provided in paragraph (c)(2) of this section.

(2) A nonrisk enrollee of a risk HMO or CMP is not entitled to additional benefits under § 417.442.

(c) Conversion to risk status—

(1) Conversion based on HCFA determination. If HCFA determines that, for administrative reasons or because there are fewer than 75 current nonrisk Medicare enrollees remaining in the HMO or CMP, all of its nonrisk Medicare enrollees must be covered under the risk provisions of the contract, the conversion process is as follows:

(i) HCFA notifies each affected enrollee of the decision at least 90 days prior to the effective date.

(ii) The nonrisk Medicare enrollees complete and sign forms stating that they understand and accept the new
§ 417.446  [Reserved]

§ 417.448 Restriction on payments for services received by Medicare enrollees of risk HMOs or CMPs.

(a) Basic rule. Except for emergency and urgently needed services as defined in §417.401, risk HMOs or CMPs are not required to make payments to or on behalf of certain Medicare enrollees, for any services received by the enrollees that are not provided—

(1) Directly by the HMO or CMP; or

(2) Through arrangements made by the HMO or CMP.

(b) Application. The restriction on payments for services imposed by paragraph (a) of this section applies to services received by—

(1) New Medicare enrollees;

(2) Nonrisk Medicare enrollees who convert to risk reimbursement; and

(3) Nonrisk Medicare enrollees who elect special supplemental benefit plans.

(c) End of restriction. The restriction of payments imposed by paragraph (a) of this section ends when a Medicare enrollee leaves the HMO’s or CMP’s geographic area for an extended period as defined in §417.401(a), and the HMO or CMP and the enrollee make arrangements for enrollment to continue as provided in §417.460(a)(2)(iv).

(d) Timing. The effective date for the end of the restriction on payments, as discussed in paragraph (c) of this section is the first day of the first month following the month in which the enrollee notifies the HMO or CMP as required in §417.436(a)(9), that he or she has left the HMO’s or CMP’s geographic area for an extended period.

§ 417.450 Effective date of coverage.

(a) Basic rules. Except as specified in paragraph (b) of this section, and notwithstanding the provisions of §417.440(d).

(1) HCFA’s liability for payments to an HMO or CMP on behalf of a Medicare beneficiary begins on the first day of the month in which he or she is—

(i) Entitled to Medicare benefits; and

(ii) Enrolled in an HMO or CMP;

(2) The effective month of coverage may not be earlier than the first month after, nor later than the third month after the month in which HCFA receives the information necessary to include the beneficiary as a Medicare enrollee of the HMO or CMP in HCFA records.

(b) Exceptions.

(1) HCFA may approve a later month if it is requested by the HMO or CMP and the beneficiary.

(2) If an individual becomes an HMO or CMP enrollee before becoming entitled to Medicare Part B benefits, the effective month of coverage is the first month for which he or she becomes entitled to Medicare Part B benefits.

(c) Notice of effective date of coverage.

For each beneficiary added to HCFA’s records as an enrollee of an HMO or CMP, HCFA gives the HMO or CMP prompt written notice of the month with which HCFA’s liability begins.

§ 417.452 Liability of Medicare enrollees.

(a) Deductibles and coinsurance. (1) A Medicare enrollee of an HMO or CMP is...
§ 417.454 Charges to Medicare enrollees.

(a) Limits on charges. The HMO or CMP must agree to charge its Medicare enrollees only for the—

1. Deductible and coinsurance amounts applicable to furnished covered services;

2. Charges for noncovered services or services for which the enrollee is liable as described in §417.452; and

3. Services for which Medicare is not the primary payor as provided in §417.528.

(b) Limit on charges for inpatient hospital care. If a Medicare enrollee who is an inpatient of a hospital requests immediate PRO review (as provided in §417.605) of any determination by the hospital furnishing services or the HMO or CMP that the inpatient hospital services will no longer be covered, the HMO or CMP may not charge the enrollee for any inpatient care costs incurred before noon of the first working day after the PRO issues its review decision.

(c) Reporting requirements. A risk HMO or CMP must report, within 90 days after the end of the contract period, all premiums, enrollment fees,
§ 417.456 Refunds to Medicare enrollees.

(a) Definitions. As used in this section—

Amounts incorrectly collected means amounts collected that are in excess of those specified in §417.452. It includes amounts collected when the enrollee was believed not entitled to Medicare benefits if the enrollee is later determined to have been entitled to Medicare benefits and HCFA is liable for payments as specified in §417.450.

Other amounts due means amounts due a Medicare enrollee for services obtained outside the HMO or CMP if they were—

(1) Emergency services;

(2) Urgently needed services for which the HMO or CMP has assumed financial responsibility; or

(3) On appeal under subpart Q of this part, found to be services the enrollee was entitled to have furnished by the HMO or CMP.

(b) Basic commitment. An HMO or CMP must agree to refund all amounts incorrectly collected from its Medicare enrollees, or from others on behalf of the enrollees, and any other amounts due the enrollees or others on their behalf.

(c) Refund by lump sum payment. An HMO or CMP must make refunds to its current and former Medicare enrollees, or to others who have made payments on behalf of enrollees, by lump sum payment for the following:

(1) Incorrectly collected amounts that were not collected as premiums.

(2) Other amounts due.

(3) All amounts due, if the HMO or CMP is going out of business.

(d) Refund by premium adjustment or lump sum payment or both. An HMO or CMP may make refund by adjustment of future premiums, by lump sum payment, or by a combination of both methods, for amounts that were incorrectly collected in the form of premiums or through a combination of premium payments and other charges.

(e) Refund when enrollee has died or cannot be located. If an enrollee has died or cannot be located after reasonable effort by the HMO or CMP, the HMO or CMP must make the refund in accordance with State law.

(f) Reduction by HCFA. If the HMO or CMP does not make refund in accordance with paragraphs (b) through (d) of this section by the end of the contract period following the contract period during which an amount was determined to be due an enrollee, HCFA reduces its payment to the HMO or CMP by the amounts incorrectly collected or otherwise due, and arranges for those amounts to be paid to the Medicare enrollee.


§ 417.458 Recoupment of uncollected deductible and coinsurance amounts.

An HMO or CMP agrees not to recoup deductible and coinsurance amounts for which Medicare enrollees were liable in a previous contract period except in the following circumstances:

(a) The HMO or CMP failed to collect the deductible and coinsurance amounts during the contract period in which they were due because of—

(1) Underestimation of the actuarial value of the deductible and coinsurance amounts; or

(2) A billing error.

(b) The HMO or CMP has identified the amounts and obtained advance HCFA approval of the recoupment and the method and timing of recoupment.

(c) The HMO or CMP collects these amounts no later than the end of the contract period following the contract period during which they were found to be due.


§ 417.460 Disenrollment of beneficiaries by an HMO or CMP.

(a) General rule. Except as provided in paragraphs (b) through (i) of this section, an HMO or CMP may not—

(1) Disenroll a Medicare beneficiary; or
(2) Orally or in writing, or by any action or inaction, request or encourage a Medicare enrollee to disenroll.

(b) Bases for disenrollment: Overview.
(1) Optional disenrollment. Generally, an HMO or CMP may disenroll a Medicare enrollee if he or she—
(i) Fails to pay the required premiums or other charges;
(ii) Commits fraud or permits abuse of his or her enrollment card; or
(iii) Behaves in a manner that seriously impairs the HMO's or CMP's ability to furnish health care services to the particular enrollee or to other enrollees.

(2) Required disenrollment. Generally, an HMO or CMP must disenroll a Medicare enrollee if he or she—
(i) Moves out of the HMO's or CMP's geographic area;
(ii) Fails to convert to the risk provisions of the HMO's or CMP's Medicare contract;
(iii) Loses entitlement to Medicare Part B benefits; or
(iv) Dies.

(3) Related provisions. Specific requirements, limitations, and exceptions are set forth in paragraphs (c) through (i) of this section.

(c) Failure to pay premiums or other charges. (1) Basic rule. Except as specified in paragraph (c)(2) of this section, an HMO or CMP may disenroll a Medicare enrollee if he or she—
(i) Can demonstrate to HCFA that it made reasonable efforts to collect the unpaid amount;
(ii) Gives the enrollee written notice of disenrollment, including an explanation of the enrollee's right to a hearing under the grievance procedures and
(iii) Sends the notice of disenrollment to the enrollee before it notifies HCFA.
(2) Exception. If the enrollee fails to pay the premium for optional supplemental benefits (that is, a package of benefits that an enrollee is not required to accept), but pays the basic premium and other charges, the HMO or CMP may discontinue the optional benefits but may not disenroll the beneficiary.

(d) Enrollee commits fraud or permits abuse of the enrollment card. (1) Basis for disenrollment. An HMO or CMP may disenroll a Medicare beneficiary if the beneficiary—
(i) Knowingly provides, on the application form, fraudulent information that materially affects the beneficiary's eligibility to enroll in the HMO or CMP; or
(ii) Intentionally permits others to use his or her enrollment card to obtain services from the HMO or CMP.
(2) Notice requirement. If disenrollment is for either of the reasons specified in paragraph (d)(1) of this section, the HMO or CMP must give the beneficiary a written notice of termination of enrolment.
(i) The notice must be mailed to the enrollee before submission of the disenrollment notice to HCFA.
(ii) The notice must include an explanation of the enrollee's right to have the disenrollment heard under the grievance procedures established in accordance with §417.436.

(3) Report to the Inspector General. The HMO or CMP must report to the Office of the Inspector General of the Department any disenrollment based on fraud or abuse by the enrollee.

(4) Disenrollment for cause. (1) Basis for disenrollment. An HMO or CMP may disenroll a Medicare enrollee for cause if the enrollee's behavior is disruptive, unruly, abusive, or uncooperative to the extent that his or her continuing enrollment in the HMO or CMP seriously impairs the HMO's or CMP's ability to furnish services to either the particular enrollee or other enrollees.
(2) Effort to resolve the problem. The HMO or CMP must make a serious effort to resolve the problem presented by the enrollee, including the use (or attempted use) of internal grievance procedures.
(3) Consideration of extenuating circumstances. The HMO or CMP must ascertain that the enrollee's behavior is not related to the use of medical services or to mental illness.
(4) Documentation. The HMO or CMP must document the problems, efforts, and medical conditions as described in
paragraphs (e)(1) through (e)(3) of this section.

(5) HCFA review of an HMO's or CMP's proposed disenrollment for cause. (i) HCFA decides on the basis of review of the documentation submitted by the HMO or CMP, whether disenrollment requirements have been met.

(ii) HCFA makes this decision within 20 working days after receipt of the documentation material, and notifies the HMO or CMP within 5 working days after making its decision.

(6) Effective date of disenrollment. If HCFA permits an HMO or CMP to disenroll an enrollee for cause, the disenrollment takes effect on the first day of the calendar month after the month in which the HMO or CMP gives the enrollee a written notice of disenrollment that meets the requirements set forth in paragraphs (d)(2)(i) and (d)(2)(ii) of this section.

(f) Enrollee moves out of the HMO's or CMP's geographic area. (1) Basic rules. (i) Disenrollment. Except as provided in paragraph (f)(2) of this section, an HMO or CMP must disenroll a Medicare enrollee for cause, the disenrollment takes effect on the first day of the calendar month after the month in which the HMO or CMP gives the enrollee a written notice of disenrollment that meets the requirements set forth in paragraphs (d)(2)(i) and (d)(2)(ii) of this section.

(ii) Notice requirement. The HMO or CMP must comply with the notice requirements set forth in paragraph (d)(2) of this section.

(iii) Effect on geographic area. Failure to disenroll an enrollee who has moved out of the HMO's or CMP's geographic area does not expand that area to encompass the location of the enrollee's new residence.

(2) Exception. An HMO or CMP may retain a Medicare enrollee who is absent from its geographic area for an extended period, but who remains within the United States as defined in §400.200 of this chapter if the enrollee agrees. For purposes of this exception, the following provisions apply:

(i) An absence for an extended period means an uninterrupted absence from the HMO's or CMP's geographic area for more than 90 days but less than 1 year.

(ii) The HMO or CMP and the enrollee may mutually agree upon restrictions for obtaining services while the enrollee is absent for an extended period from the HMO's or CMP's geographic area. However, restrictions may not be imposed on the scope of services described in §417.440.

(iii) HMOs and CMPs that choose to exercise this exception must make the option available to all Medicare enrollees who are absent for an extended period from their geographic areas. However, HMOs and CMPs may limit this option to enrollees who go to a geographic area served by an affiliated HMO or CMP.

(iv) As used in this paragraph, “affiliated HMO or CMP” means an HMO or CMP that—

(A) Is under common ownership or control of the HMO or CMP that seeks to retain the absent enrollees; or

(B) Has in effect an agreement to furnish services to enrollees who are on an extended absence from the geographic area of the HMO or CMP that seeks to retain them.

(v) When the enrollee returns to the HMO's or CMP's geographic area (even temporarily), the restrictions of §417.448(a) (which limit payment for services not provided or arranged for by the HMO or CMP) apply again immediately.

(vi) If the enrollee fails to return to the HMO's or CMP's geographic area within 1 year from the date he or she left that area, the HMO or CMP must disenroll the beneficiary on the first day of the month following the anniversary of the date the enrollee left that area in accordance with paragraph (f)(1) of this section.

(g) Failure to convert to risk provisions of Medicare contract. (1) Basis for disenrollment. A risk HMO or CMP must disenroll a nonrisk Medicare enrollee who refuses to convert to the risk provisions of the Medicare contract after HCFA determines that all of the HMO's or CMP's nonrisk Medicare enrollees must convert.

(2) Advance notice requirement. At least 30 days before it gives HCFA notice of disenrollment, the HMO or CMP must give the enrollee written notice of the fact that failure to convert will result in disenrollment.
(h) Loss of entitlement to Medicare benefits. (1) Loss of entitlement to Part A benefits. If an enrollee loses entitlement to benefits under Part A of Medicare but remains entitled to benefits under Part B, the enrollee automatically continues as a Medicare enrollee of the HMO or CMP and is entitled to receive and have payment made for Part B services, beginning with the month immediately following the last month of his or her entitlement to Part A benefits.

(2) Loss of entitlement to Part B benefits. If a Medicare enrollee loses entitlement to Part B benefits, the HMO or CMP must disenroll him or her as a Medicare enrollee effective with the month following the last month of entitlement to Part B benefits. However, the HMO or CMP may continue to enroll the individual under its regular plan if the individual so chooses.

(i) Death of the enrollee. Disenrollment is effective with the month following the month of death.

[60 FR 45678, Sept. 1, 1995]

§ 417.464 End of HCFA's liability for payment: Disenrollment of beneficiaries and termination or default of contract.

(a) Effect of disenrollment: General rule. (1) HCFA's liability for monthly capitation payments to the HMO or CMP generally ends as of the first day of the month following the month in which disenrollment is effective, as shown on HCFA's records.

(2) Disenrollment is effective no earlier than the month immediately after, and no later than the third month after, the month in which HCFA receives the disenrollment notice in acceptable form.

(b) Effect of disenrollment: Special rules. (1) Fraud or abuse by the enrollee. If disenrollment is on the basis of fraud committed or abuse permitted by the enrollee, HCFA's liability ends as of the first day of the month in which disenrollment is effective.

(2) Loss of entitlement to Part B benefits. If disenrollment is on the basis of loss of entitlement to Part B benefits, HCFA's liability ends as of the first day of the month following the last month of Part B entitlement.

(3) Death of enrollee. If the enrollee dies, HCFA's liability ends as of the first day of the month following the month of death.

(4) Disenrollment at enrollee's request. If disenrollment is in response to the enrollee's request, HCFA's liability ends as of the first day of the month following the month of termination requested by the enrollee.

(c) Effect of termination or default of contract. (1) Termination of contract. If the contract between HCFA and the HMO or CMP is terminated by mutual consent or by unilateral action of either party, HCFA's liability for payments ends as of the first day of the month after the last month for which the contract is in effect.

[60 FR 45679, Sept. 1, 1995]
§ 417.470  Default of contract. If the HMO or CMP defaults on the contract before the end of the contract year because of bankruptcy or other reasons, HCFA—
(i) Determines the month in which its liability for payments ends; and
(ii) Notifies the HMO or CMP and all affected Medicare enrollees as soon as practicable.

[60 FR 45680, Sept. 1, 1995]

Subpart L—Medicare Contract Requirements

SOURCE: 50 FR 1346, Jan. 10, 1985, unless otherwise noted.

§ 417.470  Basis and scope.

(a) Basis. This subpart implements those portions of section 1857(e)(2) of the Act pertaining to cost sharing in enrollment-related costs and section 1876(c), (g), (h), and (l) of the Act that pertain to the contract between HCFA and an HMO or CMP for participation in the Medicare program.

(b) Scope. This subpart sets forth—
(1) Specific contract requirements; and
(2) Procedures for renewal, non-renewal, or termination of a contract.


§ 417.472  Basic contract requirements.

(a) Submittal of contract. An HMO or CMP that wishes to contract with HCFA to furnish services to Medicare beneficiaries must submit a signed contract that meets the requirements of this subpart and any other requirements established by HCFA.

(b) Agreement to comply with regulations and instructions. The contract must provide that the HMO or CMP agrees to comply with all the applicable requirements and conditions set forth in this subpart and in general instructions issued by HCFA.

(c) Other contract provisions. In addition to the requirements set forth in §§ 417.474 through 417.488, the contract must contain any other terms and conditions that HCFA requires to implement section 1876 of the Act.


(e) Compliance with civil rights laws. The HMO or CMP must comply with title VI of the Civil Rights Act of 1964 (regulations at 45 CFR part 80), section 504 of the Rehabilitation Act of 1973 (regulations at 45 CFR part 84), and the Age Discrimination Act of 1975 (regulations at 45 CFR part 91).

(f) Requirements for advance directives. The HMO or CMP must meet all the requirements for advance directives at §417.436(d).

(g) Authority to waive conflicting contract requirements. Under section 1876(i)(5) of the Act, HCFA is authorized to administer the terms of this subpart without regard to provisions of law or other regulations relating to the making, performance, amendment, or modification of contracts of the United States if it determines that those provisions are inconsistent with the efficient and effective administration of the Medicare program.

(h) Collection of fees from risk HMOs and CMPs. (1) The rules set forth in §422.10 of this chapter for M+C plans also apply to collection of fees from risk HMOs and CMPs.

(2) In applying the part 422 rules, references to “M+C organizations” or “M+C plans” must be read as references to “risk HMOs and CMPs”.


§ 417.474  Effective date and term of contract.

(a) Effective date. The contract must specify its effective date, which may not be earlier than the date it is signed by both HCFA and the HMO or CMP.

(b) Term. The contract must specify the duration of its term as follows:
(1) For the initial term, at least 12 months, but no more than 23 months.
(2) For any subsequent term, 12 months.

[60 FR 45680, Sept. 1, 1995]
§ 417.476 Waived conditions.

If HCFA waives any of the qualifying conditions required under subpart J of this part, the contract must specify the following information for each waived condition:

(a) The specific terms of the waiver.
(b) The expiration date of the waiver.
(c) Any other information required by HCFA.

[60 FR 45680, Sept. 1, 1995]

§ 417.478 Requirements of other laws and regulations.

The contract must provide that the HMO or CMP agrees to comply with—

(a) The requirements for PRO review of services furnished to Medicare enrollees as set forth in subchapter D of this chapter;
(b) Sections 1318(a) and (c) of the PHS Act, which pertain to disclosure of certain financial information;
(c) Section 1301(c)(8) of the PHS Act, which relates to liability arrangements to protect enrollees of the HMO or CMP; and
(d) The reporting requirements in §417.126(a), which pertain to the monitoring of an HMO’s or CMP’s continued compliance.


§ 417.479 Requirements for physician incentive plans.

(a) The contract must specify that an HMO or CMP may operate a physician incentive plan only if—

(1) No specific payment is made directly or indirectly under the plan to a physician or physician group as an inducement to reduce or limit medically necessary services furnished to an individual enrollee; and
(2) The stop-loss protection, enrollee survey, and disclosure requirements of this section are met.

(b) Applicability. The requirements in this section apply only to physician incentive plans that base compensation (in whole or in part) on the use or cost of services furnished to Medicare beneficiaries or Medicaid recipients.

(c) Definitions. For purposes of this section:

Bonus means a payment an HMO or CMP makes to a physician or physician group beyond any salary, fee-for-service payments, capitation, or returned withhold.

Capitation means a set dollar payment per patient per unit of time (usually per month) that an organization pays a physician or physician group to cover a specified set of services and administrative costs without regard to the actual number of services provided. The services covered may include the physician’s own services, referral services, or all medical services.

Payments means any amounts the HMO or CMP pays physicians or physician groups for services they furnish directly, plus amounts paid for administration and amounts paid (in whole or in part) based on use and costs of referral services (such as withhold amounts, bonuses based on referral levels, and any other compensation to the physician or physician group to influence the use of referral services). Bonuses and other compensation that are not based on referral levels (such as bonuses based solely on quality of care furnished, patient satisfaction, and participation on committees) are not considered payments for purposes of this section.

Physician group means a partnership, association, corporation, individual practice association, or other group that distributes income from the practice among members. An individual practice association is a physician group only if it is composed of individual physicians and has no subcontracts with physician groups.

Physician incentive plan means any compensation arrangement between an HMO or CMP and a physician or physician group that may directly or indirectly have the effect of reducing or limiting services furnished to Medicare beneficiaries or Medicaid recipients enrolled in the HMO or CMP.

Referral services means any specialty, inpatient, outpatient, or laboratory
services that a physician or physician group orders or arranges, but does not furnish directly.

Risk threshold means the maximum risk, if the risk is based on referral services, to which a physician or physician group may be exposed under a physician incentive plan without being at substantial financial risk.

Withhold means a percentage of payments or set dollar amounts that an HMO or CMP deducts from a physician's service fee, capitation, or salary payment, and that may or may not be returned to the physician, depending on specific predetermined factors.

(d) Prohibited physician payments. No specific payment of any kind may be made directly or indirectly under the incentive plan to a physician or physician group as an inducement to reduce or limit covered medically necessary services covered under the HMO's or CMP's contract furnished to an individual enrollee. Indirect payments include offerings of monetary value (such as stock options or waivers of debt) measured in the present or future.

(e) General rule: Determination of substantial financial risk. Substantial financial risk occurs when the incentive arrangements place the physician or physician group at risk for amounts beyond the risk threshold, if the risk is based on the use or costs of referral services. Amounts at risk based solely on factors other than a physician's or physician group's referral levels do not contribute to the determination of substantial financial risk. The risk threshold is 25 percent.

(f) Arrangements that cause substantial financial risk. For purposes of this paragraph, potential payments mean the maximum anticipated total payments (based on the most recent year's utilization and experience and any current or anticipated factors that may affect payment amounts) that could be received if use or costs of referral services were low enough. The following physician incentive plans cause substantial financial risk if risk is based (in whole or in part) on use or costs of referral services and the patient panel size is not greater than 25,000 patients:

   (1) Withholds greater than 25 percent of potential payments.
   (2) Withholds less than 25 percent of potential payments if the physician or physician group is potentially liable for amounts exceeding 25 percent of potential payments.
   (3) Bonuses that are greater than 33 percent of potential payments minus the bonus.
   (4) Withholds plus bonuses if the withholds plus bonuses equal more than 25 percent of potential payments. The threshold bonus percentage for a particular withhold percentage may be calculated using the formula:

   \[
   \text{Withhold} = 0.75 \times \text{Bonus \%} + 25\%.
   \]

   (5) Capitation, arrangements, if—

   (i) The difference between the maximum potential payments and the minimum potential payments is more than 25 percent of the maximum potential payments; or
   (ii) The maximum and minimum potential payments are not clearly explained in the physician's or physician group's contract.

   (6) Any other incentive arrangements that have the potential to hold a physician or physician group liable for more than 25 percent of potential payments.

(g) Requirements for physician incentive plans that place physicians at substantial financial risk. HMOs and CMPs that operate incentive plans that place physicians at substantial financial risk must do the following:

   (1) Conduct enrollee surveys. These surveys must—

   (i) Include either all current Medicare/Medicaid enrollees in the HMO or CMP and those who have disenrolled (other than because of loss of eligibility in Medicaid or relocation outside the HMO's or CMP's service area) in the past 12 months, or a sample of these enrollees and disenrollees;
   (ii) Be designed, implemented, and analyzed in accordance with commonly accepted principles of survey design and statistical analysis;
   (iii) Address enrollees/disenrollees satisfaction with the quality of the services provided and their degree of access to the services; and
   (iv) Be conducted no later than 1 year after the effective date of the Medicare contract and at least annually thereafter.
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(2) Ensure that all physicians and physician groups at substantial financial risk have either aggregate or per-patient stop-loss protection in accordance with the following requirements:

(i) If aggregate stop-loss protection is provided, it must cover 90 percent of the costs of referral services (beyond allocated amounts) that exceed 25 percent of potential payments.

(ii) If the stop-loss protection provided is based on a per-patient limit, the stop-loss limit per patient must be determined based on the size of the patient panel and may be a single combined limit or consist of separate limits for professional services and institutional services. In determining patient panel size, the patients may be pooled in accordance with paragraph (h)(3)(v) of this section. Stop-loss protection must cover 90 percent of the costs of referral services that exceed the per patient limit. The per-patient stop-loss limit is as follows:

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(h) Disclosure requirements for organizations with physician incentive plans—

(1) Disclosure to HCFA. Each HMO or CMP must provide to HCFA information concerning its physician incentive plans as required or requested. The disclosure must contain the following information in detail sufficient to enable HCFA to determine whether the incentive plan complies with the requirements specified in this section:

(i) Whether services not furnished by the physician or physician group are covered by the incentive plan. If only the services furnished by the physician or physician group are covered by the incentive plan, disclosure of other aspects of the plan need not be made.

(ii) The type of incentive arrangement; for example, withhold, bonus, capitation.

(iii) If the incentive plan involves a withhold or bonus, the percent of the withhold or bonus.

(iv) Proof that the physician or physician group has adequate stop-loss protection, including the amount and type of stop-loss protection.

(v) The panel size and, if patients are pooled, the method used. Pooling is permitted only if: it is otherwise consistent with the relevant contracts governing the compensation arrangements for the physician or physician group; the physician or physician group is at risk for referral services with respect to each of the categories of patients being pooled; the terms of the compensation arrangements permit the physician or physician group to spread the risk across the categories of patients being pooled; the distribution of payments to physicians from the risk pool is not calculated separately by patient category; and the terms of the risk borne by the physician or physician group are comparable for all categories of patients being pooled. If these conditions are met, the physician or physician group may use either or both of the following methods to pool patients:

(A) Pooling any combination of commercial, Medicare, or Medicaid patients enrolled in a specific HMO or CMP in the calculation of the panel size.

(B) Pooling together, by a physician group that contracts with more than one HMO, CMP, health insuring organization (as defined in §434.2 of this chapter), or prepaid health plan (as defined in §434.2 of this chapter) the patients of each of those entities.

(vi) In the case of capitated physicians or physician groups, capitation payments paid to primary care physicians for the most recent year broken down by percent for primary care services, referral services to specialists,
§ 417.480  Maintenance of records: Cost HMOs and CMPs.

A reasonable cost contract must provide that the HMO or CMP agrees to maintain books, records, documents, and other evidence of accounting procedures and practices that—

(a) Are sufficient to—

(1) Ensure an audit trail; and

(2) Properly reflect all direct and indirect costs claimed to have been incurred under the contract; and

(b) Include at least records of the following:

(2) Intermediate entities. An HMO or CMP that contracts with an entity (other than a physician group) for the provision of services to Medicare beneficiaries must do the following:

(i) Disclose to HCFA any incentive plan between the entity and a physician or physician group that bases compensation to the physician or physician group on the use or cost of services furnished to Medicare beneficiaries or Medicaid recipients. The disclosure must include the information required to be disclosed under paragraphs (h)(1)(i) through (h)(1)(vii) of this section and be made at the times specified in paragraph (h)(2) of this section.

(ii) If the physician incentive plan puts a physician or physician group at substantial financial risk for the cost of services the physician or physician group does not furnish—

(A) Meet the stop-loss protection requirements of this subpart; and

(B) Conduct enrollee surveys as specified in paragraph (g)(1) of this section.

(3) For purposes of paragraph (i)(2) of this section, an entity includes, but is not limited to, an individual practice association that contracts with one or more physician groups and a physician hospital organization.

(j) Sanctions against the HMO or CMP. HCFA may apply intermediate sanctions, or the Office of Inspector General may apply civil money penalties described at §417.500, if HCFA determines that an HMO or CMP fails to comply with the requirements of this section.

(1) Ownership, HMO or CMP, and operation of the HMO’s or CMP’s financial, medical, and other recordkeeping systems.

(2) Financial statements for the current contract period and three prior periods.

(3) Federal income tax or information returns for the current contract period and three prior periods.

(4) Asset acquisition, lease, sale, or other action.

(5) Agreements, contracts, and subcontracts.

(6) Franchise, marketing, and management agreements.

(7) Schedules of charges for the HMO’s or CMP’s fee-for-service patients.

(8) Matters pertaining to costs of operations.

(9) Amounts of income received by source and payment.

(10) Cash flow statements.

(11) Any financial reports filed with other Federal programs or State authorities.

§ 417.481 Maintenance of records: Risk HMOs and CMPs.

A risk contract must provide that the HMO or CMP agrees to maintain and make available to HCFA upon request, books, records, documents, and other evidence of accounting procedures and practices that—

(a) Are sufficient to—

(1) Establish component rates of the ACR for determining additional and supplementary benefits; and

(2) Determine the rates utilized in setting premiums for State insurance agency purposes; and

(b) Include at least any records or financial reports filed with other Federal programs or State authorities.

§ 417.482 Access to facilities and records.

The contract must provide that the HMO or CMP agrees to the following:

(a) HHS may evaluate, through inspection or other means, the quality, appropriateness, and timeliness of services furnished under the contract to its Medicare enrollees.

(b) HHS may evaluate, through inspection or other means, the facilities of the HMO or CMP when there is reasonable evidence of some need for that inspection.

(c) HHS, the Comptroller General, or their designees may audit or inspect any books and records of the HMO or CMP or its transferee that pertain to any aspect of services performed, reconciliation of benefit liabilities, and determination of amounts payable under the contract.

(d) HHS may evaluate, through inspection or other means, the enrollment and disenrollment records for the current contract period and three prior periods, when there is reasonable evidence of some need for that inspection.

(e) In the case of a reasonable cost HMO or CMP to make available for the purposes specified in paragraphs (a), (b), (c), and (d) of this section, its premises, physical facilities, and equipment, its records relating to its Medicare enrollees, the records specified in §417.480 and any additional relevant information that HCFA may require.

(f) That the right to inspect, evaluate, and audit, will extend through three years from the date of the final settlement for any contract period unless—

(1) HCFA determines there is a special need to retain a particular record or group of records for a longer period and notifies the HMO or CMP at least 30 days before the normal disposition date;

(2) There has been a termination, dispute, fraud, or similar fault by the HMO or CMP, in which case the retention may be extended to three years from the date of any resulting final settlement; or

(3) HCFA determines that there is a reasonable possibility of fraud, in which case it may reopen a final settlement at any time.

§ 417.484 Requirement applicable to related entities.

(a) Definition. As used in this section, related entity means any entity that is
§ 417.486 Disclosure of information and confidentiality.

The contract must provide that the HMO or CMP agrees to the following:

(a) To submit to HCFA—

(1) All financial information required under subpart O of this part and for final settlement; and

(2) Any other information necessary for the administration or evaluation of the Medicare program.

(b) To comply with the requirements set forth in part 420, subpart C, of this chapter pertaining to the disclosure of ownership and control information.

(c) To comply with the requirements of the Privacy Act, as implemented by 45 CFR part 5b and subpart B of part 401 of this chapter, with respect to any system of records developed in performing carrier or intermediary functions under §§ 417.532 and 417.533.

(d) To meet the confidentiality requirements of §482.24(b)(3) of this chapter for medical records and for all other enrollee information that is—

(1) Contained in its records or obtained from HCFA or other sources; and

(2) Not covered under paragraph (c) of this section.


§ 417.488 Notice of termination and of available alternatives: Risk contract.

A risk contract must provide that the HMO or CMP agrees to give notice as follows if the contract is terminated:

(a) At least 60 days before the effective date of termination, to give its Medicare enrollees a written notice that—

(1) Specifies the termination date; and

(2) Describes the alternatives available for obtaining Medicare services after termination.

(b) To pay the cost of the written notices.

[60 FR 45680, Sept. 1, 1995]

§ 417.490 Renewal of contract.

A contract with an HMO or CMP is renewed automatically for the next 12-month period unless HCFA or the HMO or CMP decides not to renew, in accordance with §417.492.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38082, July 15, 1993]

§ 417.492 Nonrenewal of contract.

(a) Nonrenewal by the HMO or CMP.

(1) If an HMO or CMP does not intend to renew its contract, it must—

(i) Give written notice to HCFA at least 90 days before the end of the current contract period;

(ii) Notify each Medicare enrollee by mail at least 60 days before the end of the contract period; and

(iii) Notify the general public at least 30 days before the end of the contract period, by publishing a notice in one or more newspapers of general circulation in each community or county located in the HMO’s or CMP’s geographic area.

(2) HCFA may accept a nonrenewal notice submitted less than 90 days before the end of a contract period if—
§ 417.494 Modification or termination of contract.

(a) Modification or termination by mutual consent. (1) HCFA and an HMO or CMP may modify or terminate a contract at any time by written mutual consent.

(2) If the contract is modified, the HMO or CMP must notify its Medicare enrollees of any changes that HCFA determines are appropriate for notification.

(3) If the contract is terminated, the HMO or CMP must notify its Medicare enrollees, and HCFA notifies the general public, at least 30 days before the termination date.

(b) Termination by HCFA. (1) HCFA may terminate a contract for any of the following reasons:

(i) The HMO or CMP has failed substantially to carry out the terms of the contract.

(ii) The HMO or CMP is carrying out the contract in a manner that is inconsistent with the effective and efficient implementation of section 1876 of the Act.

(iii) The HMO or CMP has failed substantially to comply with the composition of enrollment requirements specified in §417.413(d).

(iv) HCFA determines that the HMO or CMP no longer meets the requirements of section 1876 of the Act and this subpart for being an HMO or CMP.

(2) If HCFA decides to terminate a contract, it sends a written notice informing the HMO or CMP of its right to appeal the termination in accordance with subpart R of this part.

(3) An HMO or CMP with a risk contract must notify its Medicare enrollees of the termination as described in §417.488.

(4) HCFA notifies the HMO’s or CMP’s Medicare enrollees and the general public of the termination at least 30 days before the effective date of termination.

(c) Termination by the HMO or CMP. The HMO or CMP may terminate the contract if HCFA has failed substantially to carry out the terms of the contract.

(1) The HMO or CMP must notify HCFA at least 90 days before the effective date of the termination and must include in its notice the reasons for the termination.

(2) The HMO or CMP must notify its Medicare enrollees of the termination at least 60 days before the termination date. Risk HMOs or CMPs must also provide a written description of alternatives available for obtaining Medicare services after termination of the contract. The HMO or CMP is responsible for the cost of these notices.

(3) The HMO or CMP must notify the general public of the termination at least 30 days before the termination date.

(4) The contract is terminated effective 60 days after the HMO or CMP mails the notice to Medicare enrollees as required in paragraph (c)(2) of this section.

(5) HCFA’s liability for payment ends as of the first day of the month after
§ 417.500 Sanctions against HMOs and CMPs.

(a) Basis for imposition of sanctions. HCFA may impose the intermediate sanctions specified in paragraph (d) of this section, as an alternative to termination of contract, if HCFA determines that an HMO or CMP does one or more of the following:

(1) Fails substantially to provide the medically necessary services required to be provided to a Medicare enrollee and the failure adversely affects (or has a substantial likelihood of adversely affecting) the enrollee.

(2) Requires Medicare enrollees to pay amounts in excess of premiums permitted.

(3) Acts, in violation of the provisions of subpart K of this part, to expel or to refuse to reenroll an individual.

(4) Engages in any practice that could reasonably be expected to have the effect of denying or discouraging enrollment (except as permitted by subpart K of this part) by eligible individuals whose medical conditions or histories indicate a need for substantial future medical services.

(5) Misrepresents or falsifies information that it furnishes under this part to HCFA, an individual, or to any other entity.

(6) Fails to comply with the requirements of section 1876(g)(6)(A) of the Act relating to the prompt payment of claims.

(7) Fails to meet the requirement in section 1876(f)(1) of the Act that not more than 50 percent of the organization's enrollment be Medicare beneficiaries and Medicaid recipients.

(8) Has a Medicare risk contract and—

(i) Employs or contracts with any entity for the provision of those services (directly or indirectly) through an excluded individual or entity.

(ii) Employs or contracts with any entity for the provision of those services (directly or indirectly) through an excluded individual or entity.

(b) Notice of sanction and opportunity to respond. (1) Notice of sanction. Before imposing the intermediate sanctions specified in paragraph (d) of this section, HCFA—

(i) Sends a written notice to the HMO or CMP stating the nature and basis of the proposed sanction; and

(ii) Sends the OIG a copy of the notice (other than a notice regarding the restriction on Medicare and Medicaid enrollees as described in paragraph (a)(7) of this section), once the sanction has been confirmed following the notice period or the reconsideration.

(2) Opportunity to respond. HCFA allows the HMO or CMP 15 days from receipt of the notice to provide evidence that it has not committed an act or failed to comply with a requirement described in paragraph (a) of this section, as applicable. HCFA may allow a 15-day addition to the original 15 days upon receipt of a written request from the HMO or CMP. To be approved, the request must provide a credible explanation of why additional time is necessary and be received by HCFA before the end of the 15-day period following the date of receipt of the sanction notice. HCFA does not grant an extension if it determines that the HMO's or CMP's conduct poses a threat to an enrollee's health and safety.

(c) Informal reconsideration. If, consistent with paragraph (b)(2) of this section, the HMO or CMP submits a timely response to HCFA's notice of sanction, HCFA conducts an informal reconsideration that:

(1) Consists of a review of the evidence by a HCFA official who did not participate in the initial decision to impose a sanction; and

(2) Gives the HMO or CMP a concise written decision setting forth the factual and legal basis for the decision that affirms or rescinds the original determination.

(d) Specific sanctions. If HCFA determines that an HMO or CMP has acted
or failed to act as specified in paragraph (a) of this section and affirms this determination in accordance with paragraph (c) of this section, HCFA may—

(1) Require the HMO or CMP to suspend acceptance of applications for enrollment made by Medicare beneficiaries during the sanction period;

(2) Suspend payments to the HMO or CMP for Medicare beneficiaries enrolled during the sanction period; and

(3) Require the HMO or CMP to suspend all marketing activities to Medicare enrollees.

(e) Effective date and duration of sanctions—(1) Effective date. Except as provided in paragraph (e)(2) of this section, a sanction is effective 15 days after the date that the organization is notified of the decision to impose the sanction or, if the HMO or CMP timely seeks reconsideration under paragraph (c) of this section, on the date specified in the notice of HCFA’s reconsidered determination.

(2) Exception. If HCFA determines that the HMO’s or CMP’s conduct poses a serious threat to an enrollee’s health and safety, HCFA may make the sanction effective on a date before issuance of HCFA’s reconsidered determination.

(3) Duration of sanction. The sanction remains in effect until HCFA notifies the HMO or CMP that HCFA is satisfied that the basis for imposing the sanction has been corrected and is not likely to recur.

(f) Termination by HCFA. In addition to or as an alternative to the sanctions described in paragraph (d) of this section, HCFA may decline to renew a HMO’s or CMP’s contract in accordance with §417.492(b), or terminate the contract in accordance with §417.494(b).

(g) Civil money penalties. If HCFA determines that a HMO or CMP has committed an act or failed to comply with a requirement described in paragraph (a) of this section (with the exception of the requirement to limit the percentage of Medicare and Medicaid enrollees described in paragraph (a)(7) of this section), HCFA notifies the OIG of that determination. HCFA also conveys to the OIG information when it reverses or terminates a sanction imposed under this subpart. In accordance with the provisions of 42 CFR part 1003, the OIG may impose civil money penalties on the HMO or CMP in addition to or in place of the sanctions that HCFA may impose under paragraph (d) of this section.


Subpart M—Change of Ownership and Leasing of Facilities: Effect on Medicare Contract

§ 417.520 Effect on HMO and CMP contracts.

(a) The provisions set forth in subpart L of part 422 of this chapter also apply to Medicare contracts with HMOs and CMPs under section 1876 of the Act.

(b) In applying these provisions, references to “M+C organizations” must be read as references to “HMOs and CMPs”.

(c) In §422.550, reference to “subpart K of this part” must be read as reference to “subpart L of part 417 of this chapter”.

(d) In §422.553, reference to “subpart K of this part” must be read as reference to “subpart J of part 417 of this chapter”.

[63 FR 35067, June 26, 1998]

Subpart N—Medicare Payment to HMOs and CMPs: General Rules

§ 417.524 Payment to HMOs or CMPs: General.

(a) Basic rule. The payments that HCFA makes to an HMO or CMP under this subpart and subparts O and P of this part for furnishing covered Medicare services are in place of any payment that HCFA would otherwise make to a beneficiary or the HMO or CMP under sections 1814(b) and 1833(a) of the Act.

(b) Basis of payment. (1) HCFA pays the HMOs or CMPs on either a reasonable cost basis or a risk basis depending on the type of contract the HMO or CMP has with HCFA.

(2) In certain cases a risk HMO or CMP also receives payments on a reasonable cost basis for certain Medicare enrollees who retain nonrisk status, as
§ 417.526 Payment for covered services.

Subpart O of this part set forth the principles that HCFA follows in determining Medicare payment to an HMO or CMP that has a reasonable cost contract. Subpart P of this part describes the per capita method of Medicare payment to HMOs or CMPs that contract on a risk basis.


§ 417.528 Payment when Medicare is not primary payer.

(a) Limits on payments and charges. (1) HCFA may not pay for services to the extent that Medicare is not the primary payer under section 1862(b) of the Act and part 411 of this chapter.

(2) The circumstances under which an HMO or CMP may charge, or authorize a provider to charge, for covered Medicare services for which Medicare is not the primary payer are stated in paragraphs (b) and (c) of this section.

(b) Charge to other insurers or the enrollee. If a Medicare enrollee receives from an HMO or CMP covered services that are also covered under State or Federal worker's compensation, automobile medical, or any no-fault insurance, or any liability insurance policy or plan, including a self-insured plan, the HMO or CMP may charge, or authorize a provider that furnished the service to charge—

(1) The insurance carrier, employer, or other entity that is liable to pay for these services; or

(2) The Medicare enrollee, to the extent that he or she has been paid by the carrier, employer, or other entity.

(c) Charge to group health plans (GHPs) or large group health plans (LGHPs). An HMO or CMP may charge a GHP or LGHP for covered services it furnished to a Medicare enrollee and may charge the Medicare enrollee to the extent that he or she has been paid by the GHP or LGHP for these covered services if—

(1) The Medicare enrollee is covered under the plan; and

(2) Under section 1862(b) of the Act, HCFA is precluded from paying for the covered services.

(d) Responsibilities of HMO or CMP. An HMO or CMP must—

(1) Identify payers that are primary to Medicare under section 1862(b) of the Act;

(2) Determine the amounts payable by these payers; and

(3) Coordinate the benefits of its Medicare enrollees with these payers.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38080, July 15, 1993; 60 FR 46229, Sept. 6, 1995]

Subpart O—Medicare Payment: Cost Basis

SOURCE: 50 FR 1346, Jan. 10, 1985, unless otherwise noted.

§ 417.530 Basis and scope.

This subpart sets forth the principles that HCFA follows to determine the amount it pays for services furnished by a cost HMO or CMP to its Medicare enrollees. These principles are based on sections 1861(v) and 1876 of the Act and are, for the most part, the same as those set forth—

(a) In part 412 of this chapter, for paying the costs of inpatient hospital services which, for cost HMOs and CMPs, are considered "reasonable" only if they do not exceed the amounts allowed under the prospective payment system; and

(b) In part 413 of this chapter, for the costs of all other covered services.

[60 FR 46230, Sept. 6, 1995]

§ 417.531 Hospice care services.

(a) If a Medicare enrollee of an HMO or CMP with a reasonable cost contract makes an election under §418.24 of this chapter to receive hospice care services, payment for these services is made to the hospice that furnishes the services in accordance with part 418 of this chapter.

(b) While the enrollee's hospice election is in effect, HCFA pays the HMO or CMP on a reasonable cost basis for only the following covered Medicare services furnished to the Medicare enrollee:
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(1) Services of the enrollee's attending physician if the physician is an employee or contractor of the HMO or CMP and is not employed by or under contract to the enrollee's hospice.

(2) Services not related to the treatment of the terminal condition for which hospice care was elected or a condition related to the terminal condition.

§ 417.532 General considerations.

(a) Conditions and criteria for payment.

(1) The costs incurred by the HMO or CMP to furnish services covered by Medicare are reimbursable if they are—

(i) Proper and necessary;

(ii) Reasonable in amount; and

(iii) Except as provided in §417.550, appropriately apportioned among the HMO's or CMP's Medicare enrollees, other enrollees, and nonenrolled patients.

(2) In determining fair and equitable payment for the HMOs or CMPs, HCFA generally applies the cost payment principles set forth in §413.5 of this chapter.

(3) In judging whether costs are reasonable, HCFA applies the weighted average of the AAPCCs of each class of the HMO's or CMP's Medicare enrollees (as defined in §417.582) for the HMO's or CMP's geographic area as an absolute limitation on the total amount payable.

(b) Method and amount of payment to the HMO or CMP

(1) HCFA makes interim per capita payments each month for each Medicare enrollee, equivalent to the interim per capita cost rate determined in accordance with §417.570.

(2) HCFA adjusts the interim per capita rate as necessary during the contract period and makes final adjustments at the end of the contract period.

(3) In determining the amount due the HMO or CMP, HCFA deducts from the reasonable cost actually incurred by the HMO or CMP for covered services furnished to its Medicare enrollees, an amount equal to the actuarial value of the applicable Medicare Part A and Part B deductible and coinsurance amounts that would have applied to the covered services for which payment is being made if these enrollees had not enrolled in the HMO or CMP or another HMO or CMP.

(c) Election by HMO or CMP. An HMO or CMP must elect, on an individual provider basis, one of the following methods for payment for hospital and SNF services it furnishes to Medicare enrollees:

(1) Direct payment by HCFA.

(2) Direct payment by the HMO or CMP.

(d) Notice of election. The election must be made in writing before the beginning of the contract period and is binding for that period.

(e) Payment by HMO or CMP. If the HMO or CMP elects to pay providers directly, as provided in paragraph (c) of this section, it must—

(1) Determine the eligibility of its Medicare enrollees to receive covered services through the HMO or CMP;

(2) Make proper coverage decisions and appropriate payments, in accordance with §§421.100 and 421.200 of this chapter, for the services furnished to its Medicare enrollees;

(3) Ensure that providers maintain and furnish appropriate documentation of physician certification and recertification, to the extent required under subpart B of part 424 of this chapter;

(4) Carry out any other procedures required by HCFA.

(f) Review of HMO's or CMP's bill processing capabilities. If the HMO or CMP elects to pay providers directly, HCFA determines whether the HMO or CMP has the experience and capability to carry out the responsibilities specified in paragraph (e) of this section in an efficient and effective manner.

(g) Direct payment by HCFA. (1) If the HMO or CMP elects to have HCFA pay for provider services, HCFA pays each provider on a reasonable cost basis or under the PPS system, whichever is appropriate for the particular provider under part 412 or part 413 of this chapter.

(2) In computing the Medicare payment to the HMO or CMP, HCFA deducts these payments and any other payments made by the Medicare intermediary or carrier on behalf of the HMO or CMP (such as payment for
§ 417.533 Part B carrier responsibilities.

In paying for Part B services furnished to its enrollees by suppliers, the HMO or CMP must—

(a) Determine the eligibility of individuals to receive those services through the HMO or CMP;

(b) Make proper coverage decisions and appropriate payment as authorized under §421.200 of this chapter for the services for which its Medicare enrollees are eligible; and

(c) Carry out any other procedures that HCFA may require.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38082, July 15, 1993]

§ 417.534 Allowable costs.

(a) Definition—Allowable costs means the direct and indirect costs, including normal standby costs incurred by the HMO or CMP, that are proper and necessary for efficient delivery of needed health care services. They include the costs of furnishing services to the HMO’s or CMP’s Medicare enrollees, other enrollees, and nonenrolled patients, which are typical “provider” costs, and costs (such as marketing, enrollment, membership, and operation of the HMO or CMP) that are peculiar to health care prepayment organizations.

(b) Basic rules. (1) The allowability of an HMO’s or CMP’s costs for furnishing services is generally determined in accordance with principles applicable to provider costs, as set forth in §417.536.

(2) The allowability of other costs is determined in accordance with principles set forth in §§417.538 through 417.550.

(3) Costs for covered services for which Medicare is not the primary payor, as described in §417.528, are not allowable.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38082, July 15, 1993]

§ 417.536 Cost payment principles.

(a) Applicability. Unless otherwise specified in this subpart, the principles set forth in parts 412 and 413 of this chapter are applicable to the costs incurred by an HMO or CMP or by providers and other facilities owned or operated by the HMO or CMP or related to it by common ownership or control.

The most common examples of these costs are set forth in this section.

(b) Depreciation. An appropriate allowance for depreciation on buildings and equipment is an allowable cost, in accordance with §§413.134, 413.144, and 413.149 of this chapter.

(c) Interest expense. Necessary and proper interest on both current and capital indebtedness is an allowable cost, in accordance with §413.153 of this chapter.

(d) Cost of educational activities. An appropriate part of the net cost of approved educational activities of a provider or other health care facility owned or operated by an HMO or CMP is an allowable cost in accordance with §413.85 of this chapter.

(e) Compensation of owners. An appropriate amount of compensation for services of owners is an allowable cost, if the services are actually performed and are necessary, as specified in §413.102 of this chapter.

(f) Bad debts. (1) In accordance with §413.80 of this chapter, bad debts are deductions from revenue and may be included as allowable costs only if—

(i) They are attributable to Medicare deductible and coinsurance amounts for which the Medicare enrollee is liable; and

(ii) The HMO or CMP has made a reasonable, but unsuccessful, effort to collect those amounts.

(2) If all or part of the deductible and coinsurance amounts is payable through a monthly premium or other periodic payment, the amount allowed as a bad debt may not exceed three
§ 417.538 Enrollment and marketing costs.

(a) Principle. Costs incurred by an HMO or CMP in performing the enrollment and marketing activities described in subpart k of this part are allowable.

(b) Included costs. Allowable enrollment and marketing costs are those necessary and proper costs incurred in offering the HMO's or CMP's plan to potential enrollees in accordance with this part. Those costs include selling, advertising, promotional, and other marketing costs and may not exceed an amount that would be incurred by a

§ 417.540 Prudent and cost-conscious management.
(c) Application. Enrollment and marketing costs are allowable, whether incurred directly by HMO or CMP staff or under contract with marketing specialists or other outside consultants.
(d) Limitation on payment. The relatively higher costs that an HMO or CMP is likely to incur in initially offering its plan to Medicare beneficiaries are taken into account in determining whether enrollment and marketing costs are reasonable in amount. However, if those costs exceed amounts that would be paid by prudent management, the excess is not allowable.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38082, July 15, 1993; 60 FR 46230, Sept. 6, 1995]

§ 417.540 Enrollment costs.
(a) Principle. Enrollment costs are allowable if incurred in maintaining and servicing subscriber contracts for pre-payment enrollees.
(b) Kind of costs included. Enrollment costs include, but are not limited to, reasonable costs incurred in connection with maintaining statistical, financial, and other data on enrollees.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38082, July 15, 1993]

§ 417.542 Reinsurance costs.
Reinsurance costs are not allowable.

§ 417.544 Physicians’ services furnished directly by the HMO or CMP.
(a) Principles. (1) Compensation paid by an HMO or CMP to physicians is an allowable cost to the extent that it is commensurate with the compensation paid for similar services performed by similar physicians practicing in the same or a similar locality.
(2) Physician compensation may take various forms, but the aggregate compensation allowable must be reasonable in relation to the services personally furnished.
(3) If aggregate physician compensation costs exceed what is normally incurred, the excess is not a reasonable cost.
(b) Application. (1) In determining the allowability of the costs of physicians’ services, the cost of personal services (for example, expenses attributable to salaries, wages, incentive payments, fringe benefits) must be distinguished from the cost of nonpersonal services (for example, expenses attributable to facilities, equipment, support personnel, supplies).
(2) To be allowable, compensation must be reasonable in relation to the personal services furnished.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38082, July 15, 1993; 60 FR 46230, Sept. 6, 1995]

§ 417.546 Physicians’ services and other Part B supplier services furnished under arrangements.
General principle. The amount paid by an HMO or CMP for physicians’ services and other Part B supplier services furnished under arrangements is an allowable cost to the extent it is reasonable. Costs are considered reasonable if they—
(a) Do not exceed those that a prudent and cost-conscious buyer would incur to purchase those services; and
(b) Are comparable to costs incurred for similar services furnished by similar physicians or other suppliers in the same or a similar geographic area.


§ 417.548 Provider services through arrangements.
(a) Principle. The cost incurred by an HMO or CMP for covered services furnished under arrangement with a provider is allowable to the extent that it would be allowable and payable under parts 412 and 413 of this chapter, unless the HMO or CMP petitions HCFA and demonstrates to HFCA’s satisfaction that payment in excess of the amount authorized under parts 412 and 413 of this chapter is justified on the basis of advantages gained by the HMO or CMP.
(b) Application. An advantage gained must represent a real and tangible benefit received by the HMO or CMP for the excess cost incurred, and any excess payment is subject to other applicable requirements of parts 405, 412 and 413 of this chapter, including tests of reasonableness.
(c) Example. In the case of an arrangement an HMO or CMP has with a provider that is located outside the HMO’s or CMP’s geographic area and that is not related to the HMO or CMP by common ownership or control, payment of the provider’s charges to the HMO or CMP (rather than the payment amounts determined under part 412 or part 413 of this chapter) may be justified in exchange for the advantages of not having to incur the administrative costs of determining the provider’s reasonable cost and of making a more timely final settlement with the HMO or CMP. However, repayment of the provider’s charges would be acceptable only if—

(1) The provider furnishes services to the HMO’s or CMP’s enrollees infrequently;

(2) The charges represent an insignificant portion of total Medicare reimbursement to the HMO or CMP; and

(3) The charges do not exceed the customary charges by the provider to its other patients for similar services.


§ 417.550 Special Medicare program requirements.

(a) Principle. HCFA pays the full reasonable cost incurred by an HMO or CMP for activities that are solely for Medicare purposes and unique to Medicare contracts under section 1876 of the Act.

(b) Application. HCFA pays the full reasonable cost of the following activities:

(1) Reporting increases and decreases in the number of Medicare enrollees.

(2) Obtaining independent certification of the HMO’s or CMP’s cost report to the extent that it is for Medicare purposes.

(3) Reporting special data that HCFA requires solely for program planning and evaluation.

(c) Prior approval requirement. The costs specified in paragraph (b) of this section must be separately budgeted and approved by HCFA before the contract period begins.

(d) Limit on full payment. Full payment is limited to the costs specified in paragraph (b) of this section. All other administrative costs must be apportioned in accordance with §417.552.

[60 FR 46230, Sept. 6, 1995]

§ 417.552 Cost apportionment: General provisions.

(a) Basic rule. The HMO or CMP must apportion its total allowable direct and indirect costs among its Medicare enrollees, its other enrollees, and its non-enrolled patients—

(1) In accordance with this subpart; and

(2) Using methods approved by HCFA.

(b) Purpose of apportionment. The purpose of apportionment is to ensure that—

(1) The cost of services furnished to Medicare enrollees is not borne by other enrollees and nonenrolled patients; and

(2) The cost of the services furnished to other enrollees and nonenrolled patients is not borne by Medicare.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38082, July 15, 1993; 60 FR 46230, Sept. 6, 1995]

§ 417.554 Apportionment: Provider services furnished directly by the HMO or CMP.

The Medicare share of the cost of covered services furnished to Medicare enrollees by providers that are owned or operated by the HMO or CMP or are related to the HMO or CMP by common ownership or control must be determined in accordance with the apportionment methods set forth in part 412, §§413.24, 413.55, and 415.55 of this chapter.


§ 417.556 Apportionment: Provider services furnished by the HMO or CMP through arrangements with others.

The Medicare share of the cost of covered services furnished to Medicare enrollees through arrangements with providers other than those specified in §417.554 must be determined as follows:

(a) The Medicare share must be based on the cost the HMO or CMP pays the provider under their arrangement, to the extent that such cost is reasonable and...
within the limits established by §§ 417.534 through 417.548.

(b) Except as specified in paragraph (c) of this section, apportionment must be on the same approved basis that is used by the provider for Medicare beneficiaries who are not Medicare enrollees of the HMO or CMP, subject to the conditions and limitations set forth in §417.548.

(c) If, because of the special nature or terms of the HMO’s or CMP’s arrangement with the provider, apportionment on the basis specified in paragraph (b) of this section would result in Medicare’s bearing the costs of furnishing services to individuals other than the HMO’s or CMP’s Medicare enrollees, apportionment must be on another basis that is approved by HCFA and that will ensure that Medicare does not pay any of the cost of furnishing services to individuals who are not Medicare enrollees of the HMO or CMP.

(d) If the HMO or CMP elects to have providers reimbursed by the HMO’s or CMP’s Medicare intermediary, the Medicare share is the amount the intermediary paid the provider.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38082, July 15, 1993]

§ 417.558 Emergency, urgently needed, and out-of-area services for which the HMO or CMP accepts responsibility.

(a) Source of payment. Either HCFA or the HMO or CMP may pay a provider for emergency or urgently needed services or other covered out-of-area services for which the HMO or CMP accepts responsibility.

(b) Limits on payment. If the HMO or CMP pays, the payment amount may not exceed the amount that is allowable under part 412 or part 413 of this chapter.

(c) Exception to limit on payment. Payment in excess of the limit imposed by paragraph (b) of this section is allowable only if the HMO or CMP demonstrates to HCFA’s satisfaction that it is justified on the basis of advantages gained by the HMO or CMP, as set forth in §417.548.

[60 FR 46231, Sept. 6, 1995]

§ 417.560 Apportionment: Part B physician and supplier services.

(a) Medical services furnished directly by the HMO or CMP. The total allowable cost of Part B physician and supplier services furnished by employees or partners of the HMO or CMP or by a related entity of the HMO or CMP must be apportioned on the basis of the ratio of covered Part B services furnished to Medicare enrollees to total services furnished to all the HMO’s or CMP’s enrollees and nonenrolled patients. The HMO or CMP must use a method for reporting costs that is approved by HCFA. HCFA bases its approval on a finding that the method—

(1) Results in an accurate and equitable allocation of allowable costs; and

(2) Is justifiable from an administrative and cost efficiency standpoint.

(b) Medical services furnished under arrangements made by the HMO or CMP. When the HMO or CMP pays for Part B physician and supplier services on some basis other than fee-for-service, the reasonable cost the HMO or CMP pays under its financial arrangement with the physician or supplier must be apportioned between Medicare enrollees and others based on the ratio of covered services furnished to Medicare enrollees to the total services furnished to all enrollees and nonenrolled patients. If apportionment on this basis would result in Medicare bearing the cost of furnishing services to individuals who are not Medicare enrollees, the Medicare share must be determined on another basis (approved by HCFA) to ensure that Medicare pays only for services furnished to Medicare enrollees.

(c) Medical services furnished under an arrangement that provides for the HMO or CMP to pay on a fee-for-service basis. The Medicare share of the cost of Part B physician and supplier services furnished to Medicare enrollees under arrangements, and paid for by the HMO or CMP on a fee-for-service basis, is determined by multiplying the total amount for all such services by the ratio of charges for covered services furnished to Medicare enrollees to the total charges for all such services.

(d) Emergency services, urgently needed services, and other covered medical services for which the HMO or CMP assumes
§ 417.564 Apportionment and allocation of administrative and general costs.

(a) Costs not directly associated with providing medical care. Enrollment, marketing, and other administrative and general costs that benefit the total enrollment of the HMO or CMP and are not directly associated with furnishing medical care must be apportioned on the basis of a ratio of Medicare enrollees to the total HMO or CMP enrollment.

(b) Costs significantly related to providing medical services. (1) The following administrative and general costs, which bear a significant relationship to the services furnished, are not apportioned to Medicare directly; they must be allocated or distributed to the HMO or CMP components and then apportioned to Medicare in accordance with §§ 417.552 through 417.560:

(i) Facility costs.
(ii) Interest expense.
(iii) Medical record costs.
(iv) Centralized purchasing costs.
(v) Accounting and data processing costs.

(ii) Other administrative and general costs that are not included in paragraph (a) of this section.

(2) The allocation or distribution process must be as follows:

(i) If a separate entity or department of an HMO or CMP performs administrative functions the benefit of which cannot be quantitatively measured (such as facility costs), the total allowable costs of this entity or department must be allocated or distributed to the components of the HMO or CMP on the basis of a ratio of total incurred and distributed costs per component to the total incurred and distributed costs for all components.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38082, July 15, 1993; 60 FR 34888, July 5, 1995]

§ 417.566 Other methods of allocation and apportionment.

(a) Justification. A method of apportionment or allocation of costs, other than the methods prescribed in this subpart may be used if it results in a more accurate and equitable apportionment of allowable costs and is justifiable from an administrative and cost standpoint.

(b) Required approval. (1) An HMO or CMP that desires to use an alternative method must submit a written request for HCFA approval at least 90 days before the beginning of the period for which the different method is to be used.

(2) If HCFA approves use of a different method, the HMO or CMP may not revert to another method without first obtaining HCFA’s approval.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38082, July 15, 1993]

§ 417.568 Adequate financial records, statistical data, and cost finding.

(a) Maintenance of records. (1) An HMO or CMP must maintain sufficient financial records and statistical data for proper determination of costs payable by HCFA for covered services the HMO or CMP furnished to its Medicare enrollees either directly or under arrangements with others. These include accurate and sufficient detail of incurred costs and enrollment data.

(2) Unless otherwise provided for in this subpart, the HMO or CMP must follow standardized definitions and accounting, statistics, and reporting practices that are widely accepted in the health care industry.

(b) Provision of data. (1) The HMO or CMP must provide adequate cost and statistical data, based on its financial...
and statistical records, that can be verified by qualified auditors.

(2) The cost data must be based on an approved method of cost finding and, except as provided in paragraph (b)(3) of this section, on the accrual method of accounting.

(3) For governmental institutions that use a cash basis of accounting, cost data developed on this basis is acceptable. However, only depreciation on capital assets, rather than the expenditure for the capital asset, is allowable.

(c) Provider services furnished directly by the HMO or CMP. If the HMO or CMP furnishes provider services directly, the provider is subject to the cost-finding and cost-reporting requirements set forth in parts 412 and 413 of this chapter. The provider must use an approved cost-finding method described in §413.24 of this chapter to determine the actual cost of these covered services.

(d) Supplier services furnished directly by the HMO or CMP. If the HMO or CMP furnishes Part B physician and supplier services directly, it must furnish statistics that indicate the frequency and type of service provided, in the form and detail prescribed by HCFA.

(e) Part B physician and supplier services furnished through arrangement. If the HMO or CMP furnishes Part B physician and supplier services under arrangements with others, it must furnish to HCFA statistical, financial, and other information with respect to those services in the form and detail prescribed by HCFA.

§ 417.572 Budget and enrollment forecast and interim reports.

(a) Annual submittal. The HMO or CMP must submit an annual operating budget and enrollment forecast, in the form and detail required by HCFA, at least 90 days before the beginning of each contract period. The forecast

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must be based on financial and statistical data and records that can be verified if HCFA requires a detailed review of supporting records. The data and records include, but are not limited to, all ledgers, books, records, and original evidence of costs, and statistical data used in the determination of reasonable cost.

(b) Effect of failure to submit on time. If the HMO or CMP does not submit the budget and enrollment forecast on time, HCFA may—

(1) Establish an interim per capita rate of payment on the basis of the best available data and adjust payments on the basis of that rate until the required reports are submitted and a new interim per capita rate can be established; or

(2) If there is not enough data on which to base an interim per capita rate, inform the HMO or CMP that interim payments will not be made until the required reports are submitted.

(c) Interim cost reports. (1) An HMO or CMP must submit interim cost reports on a quarterly basis in the form and detail prescribed by HCFA. These interim cost reports must be submitted no later than 60 days after the close of each quarter of the contract period.

(2) HCFA may reduce the frequency of the reports required under paragraph (c)(1) of this section if HCFA determines that, on the basis of the HMO’s or CMP’s reporting experience, there is good cause to do so.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38082, July 15, 1993]

§ 417.574 Interim settlement.

(a) Determination. Within 30 days following the receipt of the HMO’s or CMP’s final interim cost and enrollment reports, HCFA will make an interim determination of the estimated amount payable to the HMO or CMP for the reasonable cost of covered services furnished to its Medicare enrollees, determined in accordance with subpart O of this part and including—

(i) The per capita costs incurred in furnishing covered services to its Medicare enrollees, determined in accordance with subpart O of this part and including—

(A) The costs incurred by entities related to the HMO or CMP by common ownership or control; and

(B) For reports for cost-reporting periods that begin on or after January 1, 1996, the costs of hospital and SNF services paid by Medicare’s intermediaries under the option provided by §417.532(d).

(ii) The HMO’s or CMP’s methods of apportioning cost among Medicare enrollees, and nonenrolled patients, in accordance with the payment procedures specified in this subpart (as, applicable, in parts 412 and 413 of this chapter); and

(iii) Any other information required by HCFA.

(b) Payment. Any difference between the total amount of interim payments and the amount found payable on the basis of the interim determination under paragraph (a) of this section, must be paid by the HMO or CMP or will be paid by HCFA, whichever is appropriate, no later than 30 days after HCFA’s determination.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38082, July 15, 1993]

§ 417.576 Final settlement.

(a) General rule. Final settlement and payment of amounts due the HMO or CMP or the appropriate Medicare trust funds are made following the HMO’s or CMP’s submission and HCFA’s review of an independently certified cost report and supporting documents described in paragraph (b) of this section.

(b) Certified cost report as basis for final settlement—(1) Timing of cost report. The HMO or CMP must submit to HCFA an independently certified cost report and supporting documents, in the form and detail required by HCFA, no later than 180 days after the end of each contract period, unless HCFA extends the period for good cause shown by the HMO or CMP.

(2) Content of cost report. The cost report and supporting documents must include the following:

(i) The per capita costs incurred in furnishing covered services to its Medicare enrollees, determined in accordance with subpart O of this part and including—

(A) The costs incurred by entities related to the HMO or CMP by common ownership or control; and

(B) For reports for cost-reporting periods that begin on or after January 1, 1996, the costs of hospital and SNF services paid by Medicare’s intermediaries under the option provided by §417.532(d).

(ii) The HMO’s or CMP’s methods of apportioning cost among Medicare enrollees, and nonenrolled patients, in accordance with the payment procedures specified in this subpart (as, applicable, in parts 412 and 413 of this chapter); and

(iii) Any other information required by HCFA.

(3) Failure to report required financial information. If the HMO or CMP fails to
submit the required cost report and supporting documents within 180 days (or an extended period approved by HCFA under paragraph (b)(1) of this section), HCFA may—

(i) Consider the failure to report as evidence of likely overpayment; and

(ii) Initiate recovery of amounts previously paid, or reduce interim payments, or both.

(c) Final determination and adjustment. (1) After receipt of acceptable reports as specified in paragraph (b) of this section, HCFA determines the total payment due the HMO or CMP for furnishing covered services to its Medicare enrollees (which is subject to the audit provisions of this subpart) and makes a retroactive adjustment to bring interim payments into agreement with the payable amount due the HMO or CMP.

(2) A final settlement may be made with the HMO or CMP even though a provider that is not owned or operated by the HMO or CMP or related to the HMO or CMP by common ownership or control and that provides services to the HMO’s or CMP’s Medicare enrollees has not had a final settlement with HCFA under parts 412 and 413 of this chapter for services furnished to Medicare beneficiaries who are not enrolled in the HMO or CMP. In this situation—

(i) HCFA must be satisfied that the costs of covered services furnished to the HMO’s or CMP’s Medicare enrollees, as shown in the reports specified in paragraph (b) of this section, are reasonable and that the interest of the Medicare program would best be served by not delaying final settlement with the HMO or CMP until there is a final settlement with the provider for services furnished to Medicare beneficiaries who are not enrolled in the HMO or CMP; and

(ii) Prompt settlement with the HMO or CMP would be in the best interest of the Medicare program if, for instance, the provider’s costs represent an insignificant portion of total payment due to the HMO or CMP; or if HCFA is satisfied that the provider’s costs, as shown in the reports specified in paragraph (b) of this section, will not be modified, to any significant extent, by the final settlement with the provider under parts 412 and 413 of this chapter.

(d) Notice of amount of payment. The notice of amount of Medicare payment—

(1) Explains HCFA’s determination regarding total Medicare payment due the HMO or CMP for the contract period covered by the financial information specified in paragraph (b) of this section;

(2) Relates this determination to the HMO’s or CMP’s claimed total payable cost for that period;

(3) Explains the amounts and reasons, by appropriate reference to law, regulations, and Medicare program policy and procedures, if the determined amounts differ from the HMO’s or CMP’s claim; and

(4) Informs the HMO or CMP of its right to a hearing in accordance with subpart R of part 405 of this chapter.

(e) Basis for retroactive adjustment. (1) HCFA’s determination (as contained in the notice of amount of Medicare payment) constitutes the basis for making retroactive adjustments to any Medicare payment made to the HMO or CMP during the period to which the determination applies.

(2) Further payments to the HMO or CMP may be withheld or offset in order to recover, or to aid in the recovery of, any overpayment identified in the determination as having been made to the HMO or CMP, even if the HMO or CMP requests a hearing under subpart R of part 405 of this chapter.

(3) Any withholding continues until the earliest of the following occurs:

(i) The overpayment is liquidated.

(ii) The HMO or CMP enters into an agreement with HCFA to refund the overpaid amount.

(iii) HCFA, on the basis of subsequently acquired information, determines that there was no overpayment.

(iv) The decision of a hearing specified in paragraph (d)(4) of this section is that there was no overpayment.


Subpart P—Medicare Payment: Risk Basis

Source: 50 FR 1346, Jan. 10, 1985, unless otherwise noted.
§ 417.580 Basis and scope.

(a) Basis. This subpart implements those portions of section 1876 (a), (e), and (g) of the Act that pertain to the amount HCFA pays an organization for its Medicare enrollees who are enrolled on a risk basis.

(b) Scope. This subpart sets forth—

(1) Method of payment;

(2) Procedures for determining the HMO’s or CMP’s payment rate; and

(3) Procedures for determining the additional benefits (and their value) the HMO or CMP must provide to its Medicare enrollees.

§ 417.582 Definitions.

As used in this subpart—

AAPCC stands for adjusted average per capita cost.

ACR stands for adjusted community rate.

Actuarial factors means factors such as the age, sex, and disability level distribution of the population and any other relevant factors that HCFA determines have a significant effect on the level of utilization and cost of health services.

APCRP stands for average of per capita rates of payment.

Class of Medicare enrollees means a group of Medicare enrollees of an HMO or CMP that HCFA constructs on the basis of actuarial factors.

Similar area means an area similar to the HMO’s or CMP’s geographic area but free from special characteristics that would distort the determination of the AAPCC.

U.S. per capita incurred cost means the average per capita cost, including intermediary or carrier administrative costs, incurred by Medicare, as determined on an accrual basis, for covered services furnished to Medicare beneficiaries nationwide during the most recent period for which HCFA has complete data.

§ 417.584 Payment to HMOs or CMPs with risk contracts.

Except in the circumstances specified in §417.440(d) for inpatient hospital care, and as provided in §417.585 for hospice care, HCFA makes payment for covered services only to the HMO or CMP.

(a) Principle of payment. HCFA makes monthly advance payments equivalent to the HMO’s or CMP’s per capita rate of payment for each beneficiary who is registered in HCFA records as a Medicare enrollee of the HMO or CMP.

(b) Determination of rate. (1) The annual per capita rate of payment for each class of Medicare enrollees is equal to 95 percent of the AAPCC (as determined under the provisions of §417.588) for that class of Medicare enrollees.

(2) HCFA furnishes each HMO or CMP with its per capita rate of payment for each class of Medicare enrollees not later than 90 days before the beginning of the HMO’s or CMP’s contract period.

(c) Adjustments to payments. If the actual number of Medicare enrollees differs from the estimated number on which the amount of advance monthly payment was based, HCFA adjusts subsequent monthly payments to take account of the difference.

(d) Reduction of payments. If an HMO or CMP requests a reduction in its monthly payment in accordance with §417.592(b)(2), HCFA reduces the amount of payment by the appropriate amount.

(e) Determination of rate for calendar year 1998. For calendar year 1998, HMOs or CMPs with risk contracts will be paid in accordance with §417.592(b)(2), HCFA reduces the amount of payment by the appropriate amount.

§ 417.585 Special rules: Hospice care.

(a) No payment is made to an HMO or CMP on behalf of a Medicare enrollee who has elected hospice care under §418.24 of this chapter except for the portion of the payment applicable to the additional benefits described in §417.592. This no-payment rule is effective from the first day of the month.
§ 417.588 Computation of adjusted average per capita cost (AAPCC).

(a) Basic data. In computing the AAPCC, HCFA uses the U.S. per capita incurred cost and adjusts it by the factors specified in paragraph (c) of this section to establish an AAPCC for each class of Medicare enrollees.

(b) Advance notice to the HMO or CMP. Before the beginning of a contract period, HCFA informs the HMO or CMP of the specific adjustment factors it will use in computing the AAPCC.

(c) Adjustment factors. (1) Geographic. HCFA makes an adjustment to reflect the relative level of Medicare expenditures for beneficiaries who reside in the HMO’s or CMP’s geographic area (or a similar area). This adjustment is based on reimbursement for Medicare covered services and uses the most accurate and timely data that pertain to the HMO’s or CMP’s geographic area and that is available to HCFA when it makes the determination.

(2) Enrollment. HCFA makes a further adjustment to remove the cost effect of all area Medicare beneficiaries who are enrolled in the HMO or CMP or another HMO or CMP.

(3) Age, sex, and disability status. HCFA makes adjustments to reflect the age and sex distribution and the disability status of the HMO’s or CMP’s enrollees based on Medicare program experience and available data that indicate cost differences that result from those factors.

(4) Other relevant factors. If accurate data are available and appropriate, HCFA makes adjustments to reflect welfare and institutional status and other relevant factors.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38083, July 15, 1993; 60 FR 46232, Sept. 6, 1995]

§ 417.590 Computation of the average of the per capita rates of payment.

(a) Computation by the HMO or CMP. As indicated in § 417.584(b), before an HMO’s or CMP’s contract period begins, HCFA determines a per capita rate of payment for each class of the HMO’s or CMP’s Medicare enrollees. In order to determine the additional benefits required under § 417.592, weighted averages of those per capita rates must be computed separately for enrollees entitled to Part A and Part B, and for enrollees entitled only to Part B. Except as provided in paragraph (b) of this section, the HMO or CMP must make the computations.

(b) Computation by HCFA. If the HMO or CMP claims to have insufficient enrollment experience to make the computations required by paragraph (a) of this section, and HCFA agrees with the claim, HCFA makes the computations, using the best available information, which may include the enrollment experience of other risk HMOs and CMPs.

[58 FR 38075, July 15, 1993]

§ 417.592 Additional benefits requirement.

(a) General rules. (1) An HMO or CMP that has an APCRP (as determined under § 417.590) greater than its ACR (as determined under § 417.594) must elect one of the options specified in paragraph (b) of this section.
§ 417.594 Computation of adjusted community rate (ACR).

(a) Basic rule. Each HMO or CMP must compute its basic rate as follows:

(1) Compute an initial rate in accordance with paragraph (b) of this section.

(2) Adjust and reduce the initial rate in accordance with paragraphs (c) and (d) of this section.

(b) Computation of initial rates. (1) The HMO or CMP must compute its initial rate using either of the following systems:

(i) A community rating system as defined in §417.104(b); or

(ii) A system, approved by HCFA, under which the HMO or CMP develops an aggregate premium for all its enrollees and weights the aggregate by the size of the various enrolled groups that compose its enrollment.

(For purposes of this section, enrolled groups are defined as employee groups or other bodies of subscribers that enroll in the HMO or CMP through payment of premiums.)

(2) Regardless of which method the HMO or CMP uses—

(i) The initial rate must be equal to the premium it would charge its non-Medicare enrollees for the Medicare-covered services;

(ii) The HMO or CMP must compute the rates separately for enrollees entitled to both Part A and Part B benefits and for those entitled only to Part B;

(iii) The HMO or CMP must identify and take into account anticipated revenue from health insurance payers for those services for which Medicare is not the primary payer as provided in §417.528.

(c) Special rules: Additional benefits option. (1) The HMO or CMP must determine additional benefits separately for enrollees entitled to both Part A and Part B benefits and those entitled only to Part B.

(2) The HMO or CMP may elect to provide additional benefits in any of the following forms—

(i) A reduction in the HMO’s or CMP’s premium or in other charges it imposes in the form of deductibles or coinsurance.

(ii) Health benefits in addition to the required Part A and Part B covered services.

(iii) A combination of reduced charges and additional benefits.

(d) Notification to HCFA. (1) The HMO or CMP must give HCFA notice of its ACR and its weighted APCRP at least 45 days before its contract period begins.

(2) An HMO or CMP that elects the option of providing additional benefits must include in its submittal—

(i) A description of the additional benefits it will provide to its Medicare enrollees; and

(ii) Supporting evidence to show that the selected benefits meet the requirements of paragraph (a)(2) of this section with respect to dollar value equivalence.

[60 FR 46232, Sept. 6, 1995]
(iii) Other medical services (for example, X-ray and laboratory services).
(iv) Home health services.
(v) Out-of-plan claims for emergency services.
(vi) Skilled nursing care services.
(vii) Ambulance services.
(viii) Other Medicare covered services.
(ix) General and administrative.
(x) Noncovered Medicare services (for example, eyeglasses).
(xi) Services for which Medicare is the secondary payer.
(xii) Enrollee liabilities (for example, deductibles, coinsurance, or copayments) for covered services.

(4) An HMO or CMP that does not usually separate its premium components as described in paragraph (b)(3) of this section may calculate its initial rate with the methods it uses for its other enrolled groups if the HMO or CMP provides HCFA with the documentation necessary to support any adjustments the HMO or CMP makes to the initial rate in accordance with paragraph (e) of this section.

(5) The initial rate calculation must not carry forward any losses experienced by the HMO or CMP during prior contract periods. The HMO or CMP must submit supporting documentation to assure HCFA that rates do not include past losses but only premiums for the price of additional benefits and services of the upcoming contract period.

(c) Adjustment of initial rates. (1) Purpose of adjustment. The purpose of adjustment is to reflect the utilization characteristics of Medicare enrollees.

(2) Adjustment by the HMO or CMP. The HMO or CMP may adjust the rate for a particular service using more than one of the following factors if they do not duplicate each other:

(i) Unit of service. If the HMO or CMP purchases or identifies services on a unit of service basis and the unit of service is defined the same for all enrollees, the HMO or CMP may make an adjustment in its initial rate to reflect the number of units of services furnished to its Medicare enrollees in comparison to those furnished to other enrollees.

(ii) Complexity or intensity of services. The HMO or CMP may make an adjustment to reflect the differences in the complexity or intensity of services furnished to its Medicare enrollees if the calculation of its initial rate includes the elements of this adjustment.

(3) Support documentation. All adjustments made by the HMO or CMP must be accompanied by adequate supporting data. If an HMO or CMP does not have sufficient enrollment experience to develop this data, it may, during its initial contract period, use documented statistics from a nationally recognized statistical source.

(4) Adjustment by HCFA. If the HMO or CMP does not have adequate data to adjust the initial rate calculated under paragraph (b) of this section to reflect the utilization characteristics of its Medicare enrollees, HCFA will, at the HMO's or CMP's request, adjust the initial rate. HCFA adjusts the rate on the basis of differences in the utilization characteristics of—

(i) Medicare and non-Medicare enrollees in other HMOs or CMPs; or

(ii) Medicare beneficiaries (in the HMO's or CMP's area, or State, or the United States) who are eligible to enroll in an HMO or CMP and other individuals in that same area, or State, or the United States.

(d) Reduction of adjusted rates. The HMO or CMP or HCFA further reduces the adjusted rates by the actuarial value of applicable Medicare deductibles and coinsurance.

(e) HCFA review. (1) Submission of data. The HMO or CMP must submit its ACR and the methodology used to compute it for HCFA review and approval, and must include adequate supporting data.

(2) Appeals procedures. (i) If HCFA determines that an HMO's or CMP's ACR computation is not acceptable, the HMO or CMP may, within 30 days after receipt of notice of the determination, file with HCFA a request for a hearing.

(ii) The request must state why the HMO or CMP believes the determination is incorrect, and include any supporting evidence the HMO or CMP considers pertinent.

(iii) A hearing officer designated by HCFA conducts the hearing in accordance with the hearing procedures set
§ 417.596 Establishment of a benefit stabilization fund.

(a) General. If an HMO or CMP is required to provide its Medicare enrollees with additional benefits as described in § 417.592, the organization may request that HCFA withhold a part of its monthly per capita payment in a benefit stabilization fund. The fund will be used to prevent excessive fluctuation in the provision of those additional benefits in subsequent contract periods.

(b) Notification to HCFA. An HMO's or CMP's request to have monies withheld in a benefit stabilization fund must be made when the HMO or CMP notifies HCFA under § 417.592(d) of its ACR and its APCRP in preparation for its next contract period.

(c) Limitations on the amounts withheld—(1) Limit per contract period. Except as provided in paragraph (c)(3) of this section, HCFA does not withhold in a benefit stabilization fund more than 15 percent of the difference between an HMO's or CMP's ACR and its APCRP for a given contract period.

(2) Cumulative limit. If HCFA has established a benefit stabilization fund for an HMO or CMP, it does not approve a request for withholding made by that HMO or CMP for a subsequent contract period that would cause the total value of the benefit stabilization fund to exceed 25 percent of the difference between an HMO's or CMP's ACR and its APCRP for a given contract period.

(3) Exception. HCFA may grant an exception to the limit described in paragraph (c)(3) of this section if an HMO or CMP can demonstrate to HCFA's satisfaction that the value of the additional benefits it provides to its Medicare enrollees fluctuates substantially in excess of 15 percent from one contract period to another.

(d) Financial management of benefit stabilization funds. (1) The amounts withheld by HCFA to establish and maintain a benefit stabilization fund are in the custody of the Federal Health Insurance Trust Fund and the Federal Supplementary Medical Insurance Trust Fund.

(2) The amounts withheld in a benefit stabilization fund are accounted for by HCFA in accounts in which interest does not accrue to the HMO or CMP.

§ 417.597 Withdrawal from a benefit stabilization fund.

(a) Notification to HCFA. An HMO's or CMP's request to make a withdrawal from its benefit stabilization fund for use during a contract period must be made when the HMO or CMP notifies HCFA of its ACR and its APCRP for that contract period. In making its request, the HMO or CMP must—

(1) Indicate how it intends to use the withdrawn amounts;

(2) Justify the need for the withdrawal in terms of stabilizing the additional benefits it provides to Medicare enrollees;

(3) Document the HMO's or CMP's experience with fluctuations of revenue requirements relative to the additional benefits it provides to Medicare enrollees; and

(4) Document its experience during the contract period previous to the one for which it requests withdrawal to ensure that the HMO or CMP will not be using the withdrawn amounts to refinance losses suffered during that previous contract period.

(b) Criteria for HCFA approval. HCFA approves a request for a withdrawal from a benefit stabilization fund for use during the next contract period only if—

(1) The HMO's or CMP's average of its per capita rates of payment for the next contract period is less than that of the previous contract period;

(2) The HMO's or CMP's ACR for the next contract period is significantly higher than that of the previous contract period; or

(3) The HMO's or CMP's revenue requirements for the next contract period for providing the additional benefits it provided during the previous contract period is significantly higher
than the requirements for that previous period and the ACR for the next contract period results in an additional benefits package that is less in total value than that of the previous contract period.

(c) Basis for denial. HCFA does not approve a request for a withdrawal from a benefit stabilization fund if the withdrawal would allow the HMO or CMP to—

(1) Offer without charge the supplemental services it provides to its Medicare enrollees under the provisions of §417.440(b)(2) or (b)(3); or

(2) Refinance prior contract period losses or to avoid losses in the upcoming contract period.

(d) Form of payment. Payment of monies withdrawn from a benefit stabilization fund is made, in equal parts, as an additional amount to the monthly advance payment made to the HMO or CMP under §417.594 during the period of the contract.

[58 FR 38075, July 15, 1993, as amended at 60 FR 46233, Sept. 6, 1995]

§417.598 Annual enrollment reconciliation.

HCFA’s payment to an HMO or CMP may be subject to an enrollment reconciliation at least annually. HCFA conducts this reconciliation as necessary to ensure that the payments made do not exceed or fall short of the appropriate per capita rate of payment for each Medicare enrollee of the HMO or CMP during the contract period. The HMO or CMP must submit any information or reports required by HCFA to conduct the reconciliation.

[50 FR 38075, July 15, 1993, as amended at 58 FR 36080, July 15, 1993; 60 FR 46233, Sept. 6, 1995]

Subpart Q—Beneficiary Appeals

§417.600 Basis and scope.

(a) Statutory basis. (1) Section 1869 of the Act provides the right to a hearing and to judicial review for any individual dissatisfied with a determination regarding his or her Medicare benefits.

(2) Section 1876 of the Act provides for Medicare payments to HMOs and CMPs that contract with HCFA to enroll Medicare beneficiaries and furnish Medicare-covered health care services to them. Section 1876(c)(5) provides that—

(i) An HMO or CMP must establish grievance and appeals procedures; and

(ii) Medicare enrollees dissatisfied because they do not receive health care services to which they believe they are entitled, at no greater cost than they believe they are required to pay, have the following appeal rights:

(A) The right to an ALJ hearing if the amount in controversy is $100 or more.

(B) The right to judicial review of the hearing decision if the amount in controversy is $1000 or more.

(iii) The Medicare enrollee and the HMO or CMP are parties to the hearing and to the judicial review.

(b) Scope. This subpart sets forth—

(1) The appeals procedures, as required by section 1876(c)(5)(B) of the Act for Medicare enrollees who are dissatisfied with an “organization determination” as defined in §417.606;

(2) The applicability of grievance procedures established by the HMO or CMP under section 1876(c)(5)(A) of the Act and §417.604(a) for complaints that do not involve an organization determination;

(3) The responsibility of the HMO or CMP—

(i) To develop and maintain procedures; and

(ii) To ensure all Medicare enrollees have a complete written explanation of their grievance and appeal rights, the availability of expedited reviews, the steps to follow, and the time limits for each procedure; and

(4) The special rules that apply when a beneficiary requests immediate PRO review of a determination that he or she no longer needs inpatient hospital care.


§417.602 Definitions.

As used in this subpart, unless the context indicates otherwise—ALJ stands for administrative law judge.
§ 417.604 General provisions.

(a) Responsibilities of the HMO or CMP.

(1) The HMO or CMP must establish and maintain—

(i) Appeals procedures that meet the requirements of this subpart for issues that involve organization determinations; and

(ii) Grievance procedures for dealing with issues that do not involve organization determinations.

(2) The HMO or CMP must ensure that all enrollees receive written information about the grievance and appeals procedures that are available to them.

(b) Limits on applicability of this subpart.

(1) If an enrollee requests immediate PRO review (as provided in § 417.605) of a determination of noncoverage of inpatient hospital care—

(i) The enrollee is not entitled to subsequent review of that issue under this subpart; and

(ii) The PRO review decision is subject to the appeals procedures set forth in part 473 of this chapter.

(2) Any determination regarding services that were furnished by the HMO or CMP, either directly or under arrangement, for which the enrollee has no further liability for payment are not subject to appeal.

(3) Services included in an optional supplemental plan under (§ 417.440(b)(2)) are subject only to a grievance procedure.

(4) Physicians and other individuals who furnish services under arrangement with an HMO or CMP have no right of appeal under this subpart, except as provided in §§ 417.609(c)(4) and 417.617(c)(4), which allow physicians and other health professionals to act on behalf of an enrollee in time-sensitive situations when an organization determination or reconsideration is being requested.

(c) Applicability of other regulations. Unless otherwise provided in this subpart, regulations at 20 CFR, part 404, subparts J and R, (pertaining respectively to conduct of hearings and representation of parties under title II of the Act) are applicable under this subpart.

§ 417.605 Immediate PRO review of a determination of noncoverage of inpatient hospital care.

(a) Right to review. A Medicare enrollee who disagrees with a determination made by an HMO, CMP, or a hospital that inpatient care is no longer necessary may remain in the hospital and may (directly or through his or her authorized representative) request immediate PRO review of the determination.

(b) Procedures. For the immediate PRO review process, the following rules apply:

(1) The enrollee or authorized representative must submit the request for immediate review—

(i) To the PRO that has an agreement with the hospital under § 466.78 of this chapter;

(ii) In writing or by telephone; and

(iii) By noon of the first working day after receipt of the written notice of the determination that the hospital stay is no longer necessary.

(2) On the date it receives the enrollee's request, the PRO must notify the HMO or CMP that a request for immediate review has been filed.

(3) The HMO or CMP must supply any information that the PRO requires to conduct its review and must make it available, by phone or in writing, by the close of business of the first full working day immediately following the day the enrollee submits the request.

(4) In response to a request from the HMO or CMP, the hospital must submit medical records and other pertinent information to the PRO by close of business of the first full working day immediately following the day the HMO or CMP makes its request.

(5) The PRO must solicit the views of the enrollee who requested the immediate PRO review (or the enrollee's representative).

(6) The PRO must make a determination and notify the enrollee, the hospital, and the HMO or CMP by close of business of the first working day after
§ 417.606 Organization determinations.

(a) Actions that are organization determinations. An organization determination is any determination made by an HMO or CMP with respect to any of the following:

(1) Payment for emergency or urgently needed services.

(2) Any other health services furnished by a provider or supplier other than the HMO or CMP that the enrollee believes—

(i) Are covered under Medicare; and

(ii) Should have been furnished, arranged for, or reimbursed by the HMO or CMP.

(3) The HMO's or CMP's refusal to provide services that the enrollee believes should be furnished or arranged for by the HMO or CMP and the enrollee has not received the services outside the HMO or CMP.

(4) Discontinuation of a service (such as a skilled nursing facility discharge), if the enrollee disagrees with the determination that the service is no longer medically necessary.

(b) Actions that are not organization determinations. The following are not organization determinations for purposes of this subpart:

(1) A determination regarding services that were furnished by the HMO or CMP, either directly or under arrangement, for which the enrollee has no further obligation for payment.

(2) A determination regarding services included in an optional supplemental plan (see §417.440(b)(2)).

(c) Relation to grievances. A determination that is not an organization determination is subject only to a grievance procedure under §417.436(a)(2).


§ 417.608 Notice of adverse organization determination.

(a) If an HMO or CMP makes an organization determination that is partially or fully adverse to the enrollee, it must notify the enrollee of the determination—

(1) Within 60 days of receiving the enrollee's request for payment for services; or

(2) As specified in §417.609(c)(3) for expedited organization determinations.

(b) The notice must—

(1) State the specific reasons for the determination; and

(2) Inform the enrollee of his or her right to a reconsideration, including the right to and conditions for obtaining an expedited reconsidered determination.

(c) The failure to provide the enrollee with timely notification of an adverse organization determination as specified in paragraph (a) of this section or in §417.609(b) (concerning time frames for expediting certain organization determinations) constitutes an adverse organization determination and may be appealed.


§ 417.609 Expediting certain organization determinations.

(a) An enrollee, or an authorized representative of the enrollee, may request that an organization determination as defined in §§417.606(a)(3) and (a)(4) be expedited. The request may be made orally to the HMO or CMP.

(b) The HMO or CMP must maintain procedures for expediting organization determinations when, upon request from an enrollee or authorized representative of the enrollee, the organization decides that making the determination according to the procedures
§ 417.616 Request for reconsideration.

(a) Method and place for filing a request. A request for reconsideration must be made in writing and filed with—

(1) The HMO or CMP that made the organization determination;

(2) An SSA office; or

(3) In the case of a qualified railroad retirement beneficiary, an RRB office.

(b) Time for filing a request. Except as provided in paragraph (c) of this section, the request for reconsideration must be filed within 60 days from the date of the notice of the organization determination.

(c) Extension of time to file a request.

(1) Rule. If good cause is shown, the HMO or CMP that made the organization determination may extend the time for filing the request for reconsideration.

§ 417.614 Right to reconsideration.

Any party who is dissatisfied with an organization determination or with one that has been reopened and revised may request reconsideration of the determination in accordance with the procedures of § 417.616, concerning a request for reconsideration, or § 417.617, concerning certain expedited reconsiderations.

§417.617 Expiring certain reconsiderations.

(a) An enrollee, or an authorized representative of the enrollee, may request that a reconsideration be expedited. The request may be made orally to the HMO or CMP.

(b) The HMO or CMP must maintain procedures for expediting reconsiderations when, upon request from an enrollee or an authorized representative of the enrollee, the organization decides that the longer time frames permitted in §417.620(c) could seriously jeopardize the life or health of the enrollee or the enrollee’s ability to regain maximum function.

(c) The procedures must comply with the requirements for reconsidered determinations set forth in §§417.614 through 417.626 and include the following items:

(1) Receipt of oral requests, followed by written documentation of the oral requests.

(2) Prompt decision-making regarding whether the request will be expedited or handled within the standard time frame of §417.620(c), including notification of the enrollee if the request is not expedited.

(3) Notification of the enrollee, and the physician as appropriate, as expeditiously as the enrollee’s health condition requires, but within 72 hours of the request. An extension of up to 10 working days is permitted if requested by the enrollee or if the HMO or CMP finds that additional information is necessary and the delay is in the interest of the enrollee.

(i) Notification must comply with §417.624(b), concerning the content of a notice of a reconsidered determination.

(ii) If the initial notification is not in writing, written confirmation must be mailed to the enrollee within 2 working days.

(iii) In cases for which the HMO or CMP must receive medical information from a physician or provider not affiliated with the HMO or CMP, the time standard begins with receipt of the information.

(d) Granting the request of a physician, regardless of whether the physician is affiliated with the organization or not, to expedite the request.


§417.618 Opportunity to submit evidence.

The HMO or CMP must provide the parties to the reconsideration reasonable opportunity to present evidence and allegations of fact or law, related to the issue in dispute, in person as well as in writing. In the case of an expedited reconsideration, the opportunity to present evidence is limited by the short time frames for making decisions, and the organization must inform the enrollee, or the authorized representative of the enrollee, of the conditions for submitting the evidence.


§417.620 Responsibility for reconsiderations; time limits.

(a) If the HMO or CMP can make a reconsidered determination that is completely favorable to the enrollee, the HMO or CMP issues the reconsidered determination.

(b) If the HMO or CMP recommends partial or complete affirmation of its
adverse determination, the HMO or CMP must prepare a written explanation and send the entire case to HFCA. HCFA makes the reconsidered determination.

(c) The HMO or CMP must issue the reconsidered determination to the enrollee, or submit the explanation and file to HCFA within 60 calendar days from the date of receipt of the request for reconsideration. In the case of an expedited reconsideration, the HMO or CMP must issue the reconsidered determination as specified in §417.617(c)(3) or submit the explanation and file to HCFA within 24 hours of its determination, the expiration of the 72-hour review period, or the expiration of the extension.

(d) For good cause shown, HCFA may allow extensions to the time limit set forth in paragraph (c) of this section.

(e) Failure by the HMO or CMP to provide the enrollee with a reconsidered determination within the time limits described in paragraph (c) of this section or to obtain a good cause extension described in paragraph (d) of this section constitutes an adverse determination, and the HMO or CMP must submit the file to HCFA.

(f) If the HMO or CMP refers the matter to HCFA, it must concurrently notify the beneficiary of that action.

§417.622 Reconsidered determination.

A reconsidered determination is a new determination that—

(a) Is based on a review of the organization determination, the evidence and findings upon which it was based, and any other evidence submitted by the parties or obtained by HCFA or the HMO or CMP; and

(b) Is made by a person or persons who were not involved in making the organization determination.

§417.624 Notice of reconsidered determination.

(a) Responsibility for notice. The entity that makes the reconsidered determination is responsible for mailing notice to the parties and, if that entity is not HCFA, for sending a copy to HCFA.

(b) Content of notice. The notice must—

(1) State the specific reasons for the reconsidered determination;

(2) Inform the party of his or her right to a hearing if the amount in controversy is $100 or more; and

(3) Describe the procedures that the party must follow to obtain a hearing.

[50 FR 1346, Jan. 10, 1985]

§417.626 Effect of reconsidered determination.

A reconsidered determination is binding on all parties unless a request for a hearing is filed in accordance with the provisions of §417.632, or unless it is revised in accordance with §417.638.


§417.630 Right to a hearing.

If the amount remaining in controversy is $100 or more, any party to the reconsideration who is dissatisfied with the reconsidered determination has a right to a hearing. The amount remaining in controversy, which can include any combination of Part A and Part B services, is computed in accordance with §405.740 of this chapter for Part A services and §405.820(b) of this chapter for Part B services. If the basis for the appeal is the refusal of services, the projected value of those services is used in computing the amount remaining in controversy.

[59 FR 59942, Nov. 21, 1994]

§417.632 Request for hearing.

(a) Method and place for filing a request. A request for a hearing must be made in writing and filed at one of the places specified in §417.616(a).

(b) Time for filing a request. Except when the time is extended by an ALJ as provided in 20 CFR 404.933(c), a request for a hearing must be filed within 60 days of the date of the notice of reconsidered determination.

(c) Parties to a hearing. (1) The parties to a hearing must be the parties to the reconsideration and any other person or entity whose rights with respect to the reconsideration may be affected by the hearing, as determined by the ALJ.
§ 417.634

(2) The HMO or CMP must be made a party to the hearing but does not have a right to request a hearing.

(d) ALJ action when the amount in controversy is less than $100. (1) If the request plainly shows that the amount in controversy is less than $100, the ALJ dismisses the request.

(2) If, after a hearing is initiated, the ALJ finds that the amount in controversy is less than $100, he or she discontinues the hearing and does not rule on the substantive issues raised in the appeal.


§ 417.635

Departmental Appeals Board (DAB) review.

Any party to the hearing, including the HMO or CMP, who is dissatisfied with the hearing decision, may request the DAB to review the ALJ’s decision or dismissal. Regulations beginning at 20 CFR 404.967 regarding SSA Appeals Council Review are applicable to DAB review for matters addressed by this subpart.


§ 417.636

Court review.

(a) Review of ALJ’s decision. A party or the HMO or CMP may request judicial review of an ALJ’s decision if—

(1) The Departmental Appeals Board denied the party’s or the HMO’s or CMP’s request for review; and

(2) The amount in controversy is $1,000 or more.

(b) Review of Departmental Appeals Board decision. A party or the HMO or CMP may request judicial review of the Departmental Appeals Board decision if—

(1) It is the final decision of HCFA; and

(2) The amount in controversy is $1,000 or more.

(c) Request for review. The civil action must be filed in a district court of the United States in accordance with section 205(g) of the Act (see 20 CFR 422.210 for a description of the procedures to follow in requesting judicial review).


§ 417.638

Reopening determinations and decisions.

An organization, reconsidered, or revised determination made by an HMO, CMP, or HCFA, or a decision or revised decision of an ALJ or the Departmental Appeals Board, may be reopened in accordance with the provisions of §405.750 of this chapter.

[59 FR 59942, Nov. 21, 1994, as amended at 61 FR 32348, June 24, 1996]

Subpart R—Medicare Contract Appeals

SOURCE: 50 FR 1346, Jan. 10, 1985, unless otherwise noted.

§ 417.640

Determinations subject to appeal.

This subpart establishes the procedures for making and reviewing the following initial determinations:

(a) A determination that an HMO or CMP is not qualified to enter into a contract with HCFA under section 1876 of the Act.

(b) A determination that an HMO or CMP is qualified only for a reasonable cost contract.

(c) A determination to terminate, or to refuse to renew, a contract with an HMO or CMP because—

(1) The HMO or CMP has failed substantially to carry out the terms of the contract;

(2) The HMO or CMP is carrying out the contract in a manner that is inconsistent with the efficient and effective administration of section 1876 of the Act;

(3) The HMO or CMP no longer meets the applicable conditions necessary to qualify as an HMO or CMP under section 1876 of the Act and this subpart; or
(4) The HMO or CMP has failed to comply with the composition of enrollment requirements specified in §417.413(d).


§417.642 Administrative actions that are not initial determinations.
Administrative actions that are not initial determinations under this subpart include, but are not limited to, HCFA's refusal to renew a contract with an HMO or CMP when the refusal is not based on the causes specified in §417.640(c).

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38080, July 15, 1993]

§417.644 Notice of initial determination.
(a) When HCFA makes an initial determination, it gives the HMO or CMP written notice.

(b) The notice specifies—
(1) The reasons for the determination; and
(2) The HMO's or CMP's right to request reconsideration.

(c) HCFA mails the notice to the HMO or CMP at least 90 days before the end of the contract period, or in the case of termination, at least 90 days before the effective date of the termination.

[50 FR 1346, Jan. 10, 1985, as amended at 58 FR 38083, July 15, 1993; 60 FR 46234, Sept. 6, 1995]

§417.646 Effect of initial determination.
The initial determination is final and binding on all parties unless—
(a) It is reconsidered in accordance with §§417.648 through 417.658; or
(b) In the case of an initial determination described in §417.640(c), a request for a hearing is filed; or
(c) It is revised as a result of a reopening under §417.692.

§417.648 Reconsideration: Applicability.
(a) Reconsideration is the first step for appealing an organization determination specified in §417.414(a) or (b).

(b) HCFA reconsiders either of the specified determinations if the HMO or CMP files a written request in accordance with §417.650.

[60 FR 46234, Sept. 6, 1995]

§417.650 Request for reconsideration.
(a) Method and place for filing a request. A request for reconsideration must be made in writing and filed with any HCFA office.

(b) Time for filing a request. Except as provided in paragraph (c) of this section, the request for reconsideration must be filed within 60 days from the date of the notice of the initial determination.

(c) Extension of time to file a request. HCFA may, in response to a party's written petition showing good cause, accept a request for reconsideration after the expiration of the 60 day period.

(d) Proper party to file a request. Only an authorized official of the entity that was a party to an initial determination may file the request for reconsideration.

(e) Withdrawal of a request. A request for reconsideration may be withdrawn by the party who filed the request at any time before the notice of the reconsidered determination is mailed. The request for withdrawal must be in writing and filed with HCFA. If HCFA approves, the request for reconsideration is withdrawn.

§417.652 Opportunity to submit evidence.
HCFA provides the parties to the reconsideration reasonable opportunity to present as evidence any documents or written statements that are relevant and material to the matters at issue.

[50 FR 1346, Jan. 10, 1985, as amended at 60 FR 46234, Sept. 6, 1995]

§417.654 Reconsidered determination.
A reconsidered determination is a new determination that—
(a) Is based on a review of the initial determination, the evidence and findings upon which that was based, and any other written evidence submitted before notice of the reconsidered determination is mailed, including facts relating to the status of the entity subsequent to the initial determination; and
(b) Time for filing a request. Except as provided in paragraph (c) of this section, a request for a hearing must be filed within 60 days after the date of receipt of the notice of initial or reconsidered determination.

(c) Extension of time to file a request. If good cause is shown, the 60-day period to request a hearing may be extended by HCFA.

(d) Parties to a hearing. The parties to a hearing must be—

(1) The parties described in §417.660;

(2) At the discretion of the hearing officer, any interested parties who make a showing that their rights may be prejudiced by the decision to be rendered at the hearing; and

(3) HCFA.

§ 417.664 Postponement of effective date of initial determination.

When a request for a hearing with respect to an initial determination is filed timely—

(a) The effective date of the initial determination to terminate a contract with an HMO or CMP will be postponed until a hearing decision is reached; and

(b) The current contract will be extended at the end of the contract period (in the case of a determination not to renew) only—

(1) If HCFA finds that an extension of the contract will be consistent with the purpose of section 1876 of the Act; and

(2) For such period as HCFA and the HMO or CMP agree.

§ 417.666 Designation of hearing officer.

HCFA designates a hearing officer to conduct the hearing. The hearing officer need not be an ALJ.

§ 417.668 Disqualification of hearing officer.

(a) A hearing officer may not conduct a hearing in a case in which he or she is prejudiced or partial to any party or
§ 417.682 Discovery.

(a) Prehearing discovery is permitted upon timely request of a party.

(b) A request is timely if it is made before the beginning of the hearing.

§ 417.680 Witnesses.

(a) The hearing officer may examine the witnesses.

(b) The parties or their representatives are permitted to examine their witnesses and cross-examine witnesses of other parties.

§ 417.678 Evidence.

The hearing officer rules on the admissibility of evidence and may admit evidence that would be inadmissible under rules applicable to court procedures.

§ 417.672 Appointment of representatives.

A party may appoint as its representative at the hearing anyone not disqualified or suspended from acting as a representative before HCFA or otherwise prohibited by law.

§ 417.674 Authority of representatives.

(a) A representative appointed and qualified in accordance with § 417.672 may, on behalf of the represented party—

(1) Give or accept any notice or request pertinent to the proceedings set forth in this subpart;

(2) Present evidence and allegations as to facts and law in any proceedings affecting that party; and

(3) Obtain information to the same extent as the party.

(b) A notice or request sent to the representative has the same force and effect as if it had been sent to the party.

§ 417.676 Conduct of hearing.

(a) The hearing is open to the parties and to the public.

(b) The hearing officer inquires fully into all the matters at issue and receives in evidence the testimony of witnesses and any documents that are relevant and material.

(c) The hearing officer provides the parties an opportunity to enter any objection to the inclusion of any document.

(d) The hearing officer decides the order in which the evidence and the arguments of the parties are presented and the conduct of the hearing.

§ 417.670 Time and place of hearing.

(a) The hearing officer fixes a time and place for the hearing and sends written notice to the parties. The notice also informs the parties of the general and specific issues to be resolved and information about the hearing procedure.

(b) The hearing officer may, on his or her own motion, or at the request of a party, change the time and place for the hearing. The hearing officer may adjourn or postpone the hearing.

(c) The hearing officer will give the parties reasonable notice of any change in the time or place of hearing, or of adjournment or postponement.

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(a) The hearing officer fixes a time and place for the hearing and sends written notice to the parties. The notice also informs the parties of the general and specific issues to be resolved and information about the hearing procedure.

(b) The hearing officer may, on his or her own motion, or at the request of a party, change the time and place for the hearing. The hearing officer may adjourn or postpone the hearing.

(c) The hearing officer will give the parties reasonable notice of any change in the time or place of hearing, or of adjournment or postponement.

§ 417.672 Appointment of representatives.

A party may appoint as its representative at the hearing anyone not disqualified or suspended from acting as a representative before HCFA or otherwise prohibited by law.

§ 417.674 Authority of representatives.

(a) A representative appointed and qualified in accordance with § 417.672 may, on behalf of the represented party—

(1) Give or accept any notice or request pertinent to the proceedings set forth in this subpart;

(2) Present evidence and allegations as to facts and law in any proceedings affecting that party; and

(3) Obtain information to the same extent as the party.

(b) A notice or request sent to the representative has the same force and effect as if it had been sent to the party.
§ 417.684 Prehearing.

The hearing officer may schedule a prehearing conference if he or she believes that a conference would more clearly define the issues.

§ 417.686 Record of hearing.

(a) A complete record of the proceedings at the hearing is made and transcribed and made available to all parties upon request.

(b) The record may not be closed until a hearing decision has been issued.

§ 417.688 Authority of hearing officer.

In exercising his or her authority, the hearing officer must comply with the provisions of title XVIII and related provisions of the Act, the regulations issued by HCFA, and general instructions issued by HCFA in implementing that Act.

§ 417.690 Notice and effect of hearing decision.

(a) As soon as practical after the close of the hearing, the hearing officer issues a written decision that—

(1) Is based upon the evidence of record; and

(2) Contains separately numbered findings of fact and conclusions of law.

(b) The hearing officer provides a copy of the hearing decision to each party.

(c) The hearing decision is final and binding unless it is reopened and revised in accordance with §417.692.

§ 417.692 Reopening of initial or reconsidered determination or decision of a hearing officer.

(a) Initial or reconsidered determination. An initial or reconsidered determination may be reopened and revised by HCFA upon its own motion within one year of the date of the notice of determination.

(b) Decision of hearing officer. A decision of a hearing officer that is unfavorable to any party and is otherwise final may be reopened and revised by the hearing officer upon the officer's own motion within one year of the notice of the hearing decision. It may be reopened and revised by another hearing officer designated by HCFA if the hearing officer who issued the decision is unavailable.

(c) Notices. (1) The notice of reopening and of any revisions following the reopening is mailed to the parties.

(2) The notice of revision specifies the reasons for revisions.

§ 417.694 Effect of revised determination.

The revision of an initial or reconsidered determination is binding unless a party files a written request for hearing of the revised determination in accordance with §417.662.

Subparts S–T [Reserved]

Subpart U—Health Care Prepayment Plans

Source: 50 FR 1375, Jan. 10, 1985, unless otherwise noted.

§ 417.800 Payment to HCPPs: Definitions and basic rules.

(a) Definitions. As used in this subpart, unless the context indicates otherwise—

Covered Part B services means physicians' services, diagnostic X-ray tests, laboratory, other diagnostic tests, and any additional medical and other health services, that the HCPP furnishes to its Medicare enrollees.

Health care prepayment plan (HCPP) means an organization that meets the following conditions:

(1) Effective January 1, 1999, (or on the effective date of the HCPP agreement in the case of a 1998 applicant) either—
§ 417.801 Agreements between HCFA and health care prepayment plans.

(a) General requirement. In order to participate and receive payment under the Medicare program as an HCPP as defined in §417.800, an organization of the reasonable costs it incurs, as specified in subpart O of this part, for the covered Part B services furnished to its Medicare enrollees.

(ii) Deductions. In determining the amount due an HCPP for covered Part B services furnished to its Medicare enrollees, HCFA deducts, from the reasonable cost actually incurred by the HCPP, the following:

(A) The actuarial value of the Part B deductible.

(B) An amount equal to 20 percent of the cost incurred for any service that is subject to the Medicare coinsurance.

(d) Covered services not reimbursed to an HCPP. (1) Services reimbursed under Part A are not reimbursable to an HCPP. HCFA makes payment for these services directly to the hospital, or other provider of services, on a reasonable cost basis through the provider's Medicare fiscal intermediary (for more details, see parts 412 and 413 of this chapter).

(2) Covered Part B services furnished by a provider of services to an HCPP's Medicare enrollees are not payable to the HCPP. HCFA makes payment for these services to the provider on behalf of the Medicare enrollee through the provider's Medicare fiscal intermediary. This requirement does not affect Medicare payment to the HCPP for physicians' services furnished to its Medicare enrollees for which the physicians are compensated by the HCPP.

(e) Payment for services to nonenrollees. HCFA makes payment to an HCPP for covered Part B services furnished by the HCPP to a Medicare beneficiary who is not enrolled in the HCPP, if the beneficiary assigns his rights to payment in accordance with §424.55 of this chapter. Payment is made on a reasonable charge basis through the HCPP's Medicare carrier.

§ 417.802

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must enter into a written agreement with HCFA.

(2) An existing group practice prepayment plan (GPPP) that continues as an HCPP under this subpart U must have entered into a written agreement with HCFA within 60 days of January 31, 1983.

(b) Terms. The agreement must provide that the HCPP agrees to—

(1) Maintain compliance with the requirements for participation and reimbursement on a reasonable cost basis of HCPPs as specified in §417.800;

(2) Not charge the Medicare enrollee or any other person for items or services for which that enrollee is entitled to have payment made under the provisions of this part, except for any deductible or coinsurance amounts for which the enrollee is liable;

(3) Refund, as promptly as possible, any money incorrectly collected as charges or premiums, or in any other way from Medicare enrollees in the HCPP in accordance with the requirements specified in §417.456;

(4) Not impose any limitations on the acceptance of Medicare enrollees or beneficiaries for care and treatment that it does not impose on all other individuals;

(5) Meet the advance directives requirements specified in §417.436(d) of this part;

(6) Establish administrative review procedures in accordance with §§417.830 through 417.840 for Medicare enrollees who are dissatisfied with denied services or claims; and

(7) Consider any additional requirements that HCFA finds necessary or desirable for efficient and effective program administration.

(c) Duration of agreement. Except for the term of the initial agreement, the agreement is for a term of one year and may be renewed annually by mutual consent. The term of the initial agreement is set by HCFA.

(d) Termination or nonrenewal of agreement by HCFA. (1) HCFA may terminate or not renew an agreement if it determines that—

(i) The HCPP no longer meets the requirements for participation and reimbursement as an HCPP as specified in §417.800;

(ii) The HCPP is not in substantial compliance with the provisions of the agreement, applicable HCFA regulations, or applicable provisions of the Medicare law; or

(iii) The HCPP undergoes a change in ownership as specified in subpart M of this part.

(2) HCFA will give notice of termination or nonrenewal to the HCPP at least 90 days before the effective date stated in the notice.

(e) Termination or nonrenewal of agreement by HCPP. (1) If an HCPP does not wish to renew its agreement at the end of the term, it must give written notice to HCFA at least 90 days before the end of the term of the agreement. If an HCPP wishes to terminate its agreement before the end of the term, it must file a written notice with HCFA stating the intended effective date of termination.

(2) HCFA may approve the termination date proposed by the HCPP, or set a different date no later than 6 months after that date. HCFA makes this decision based on a finding that termination on a specific date would not—

(i) Unduly disrupt the furnishing of services to the community serviced by the HCPP; or

(ii) Otherwise interfere with the efficient administration of the Medicare program.

§ 417.802 Allowable costs.

(a) General rule. The costs that are considered allowable for HCPP reimbursement are the same as those for reasonable cost HMOs and CMPs specified in subpart O of this part, except those in §§417.531, 417.532 (a)(3) and (c) through (g), 417.536 (l) and (m), 417.546, 417.548, and 417.550(b)(2).

(b) Physicians' services and other Part B supplier services furnished under arrangements—(1) Principle. The amount paid by an HCPP for physicians' services and other Part B supplier services furnished under arrangements is an allowable cost to the extent it is reasonable.
(2) Application: Payment on other than a fee-for-service basis. If the HCPP pays for physicians’ services and other Part B supplier services on other than a fee-for-service basis—

(i) Except as specified in paragraph (b)(2)(ii) of this section, the costs incurred by the HCPP may be considered reasonable if they—

(A) Do not exceed those that a prudent and cost-conscious buyer would incur to purchase those services; and

(B) Are comparable to costs incurred for similar services furnished by similar physicians and other suppliers in the same or a similar locality.

(ii)(A) If a physician group to whom the HCPP makes payment compensates its physicians on a fee-for-service basis, the HCPP’s payment to the group may not exceed the reasonable charges for those services, as defined in subpart E of part 405 of this chapter.

(B) Payment in excess of the limits specified in paragraph (b)(2)(ii)(A) of this section is allowable if the group has procedures under which members of the group accept effective incentives, such as risk-sharing, designed to avoid unnecessary or unduly costly utilization of health services. In such cases, the amount paid by an HCPP is considered reasonable if it meets the conditions specified in paragraph (b)(2)(i) of this section.

§ 417.804 Cost apportionment.

(a) The HCPP follows the cost apportionment principles specified in §§ 417.552 through 417.566, except for provisions on provider costs and provisions on departmental apportionment.

(b) The HCPP may use a method for reporting costs that is approved by HCFA. HCFA bases its approval on a finding that the method—

(1) Results in an accurate and equitable allocation of allowable costs; and

(2) Is justifiable from an administrative and cost efficiency standpoint.

§ 417.806 Financial records, statistical data, and cost finding.

(a) The principles specified in § 417.568 apply to HCPPs, except those in paragraph (c) of that section.

(b) The HCPP may use a method for reporting costs that is approved by HCFA. HCFA bases its approval on a finding that the method—

(1) Results in an accurate and equitable allocation of allowable costs; and

(2) Is justifiable from an administrative and cost efficiency standpoint.

(c) An HCPP must permit the Department and the Comptroller General to audit or inspect any books and records of the HCPP and of any related organization that pertain to the determination of amounts payable for covered Part B services furnished its Medicare enrollees. For purposes of this requirement, the principles specified in § 417.486 apply to HCPPs.

§ 417.808 Interim per capita payments.

The HCPP follows the principles specified in §§ 417.570 and 417.572 on interim per capita payments, except for the following:

(a) When applying these principles to HCPPs, the term “reporting period” should be used instead of the term “contract period” contained in that section.

(b) An HCPP must submit to HCFA an annual operating budget and enrollment forecast, in the form and detail specified by HCFA, at least 60 days before the beginning of each reporting period. A reporting period must be 12 consecutive months, except that the HCPP’s initial reporting period for participating in Medicare may be as short as 6 months or as long as 18 months.

(c) An HCPP must submit to HCFA an interim cost report and enrollment data applicable to the first 6-month period of the HCPP’s reporting period in the form and detail specified by HCFA. The interim cost report must be submitted not later than 45 days after the close of the first 6-month period of the HCPP’s reporting period.

(d) In lieu of an interim payment based on the actual monthly enrollment in an HCPP, HCFA and the HCPP may agree to a uniform monthly interim reimbursement rate for a reporting period. This interim rate is based on the HCPP’s budget and enrollment forecast, if HCFA is satisfied that the rate is consistent with efficiency and economy, and will not result in excessive adjustment at the end of the reporting period.

§ 417.810 Final settlement.

(a) General requirement. HCFA and an HCPP must make a final settlement, and payment of amounts due either to the HCPP or to HCFA, following the submission and review of the HCPP’s annual cost report and the supporting documents specified in paragraph (b) of this section.

(b) Annual cost report as basis for final settlement—(1) Form and due date. An HCPP must submit to HCFA a cost report and supporting documents in the form and detail specified by HCFA, no later than 120 days following the close of a reporting period.

(2) Contents. The report must include—

(i) The HCPP’s per capita incurred costs of providing covered Part B services to its Medicare enrollees during the reporting period, including any costs incurred by another organization related to the HCPP by common ownership or control;

(ii) The HCPP’s methods of apportioning costs among its Medicare enrollees, enrollees who are not Medicare beneficiaries, and other nonenrollees, including Medicare beneficiaries receiving health care services on a fee-for-service or other basis; and

(iii) Information on enrollment and other data as specified by HCFA.

(3) Extension of time to submit cost report. HCFA may grant an HCPP an extension of time to submit a cost report for good cause shown.

(4) Failure to report required financial information. If an HCPP does not submit the required cost report and supporting documents within the time specified in paragraph (b)(1) of this section, and has not requested and received an extension of time for good cause shown, HCFA may—

(i) Regard the failure to report this information as evidence of likely overpayment and reduce or suspend interim payments to the HCPP; and

(ii) Determine that amounts previously paid are overpayments, and make appropriate recovery.

(c) Determination of final settlement. Following the HCPP’s submission of the reports specified in paragraph (b) of this section in acceptable form, HCFA makes a determination of the total reimbursement due the HCPP for the reporting period and the difference, if any, between this amount and the total interim payments made to the HCPP. HCFA sends to the HCPP a notice of the amount of reimbursement by the Medicare program. This notice—

(1) Explains HCFA’s determination of total reimbursement due the HCPP for the reporting period; and

(2) Informs the HCPP of its right to have the determination reviewed at a hearing as provided in part 405, subpart R of this chapter.

(d) Payment of amounts due. (1) Within 30 days of HCFA’s determination, HCFA or the HCPP, as appropriate,
will make payment of any difference between the total amount due and the total interim payments made to the HCPP by HCFA.

(2) If the HCPP does not pay HCFA within 30 days of HCFA’s determination of any amounts the HCPP owes HCFA, HCFA may offset further payments to the HCPP to recover, or to aid in the recovery of, any overpayment identified in its determination.

(3) Any offset of payments HCFA makes under paragraph (d)(2) of this section will remain in effect even if the HCPP has requested a hearing on the determination under the provisions of part 405, subpart R of this chapter.

(e) Tentative settlement. (1) If a final settlement cannot be made within 90 days after the HCPP submits the report specified in paragraph (b) of this section, HCFA will make an interim settlement by estimating the amount payable to the HCPP.

(2) HCFA or the HCPP will make payment within 30 days of HCFA’s determination under the tentative settlement of any estimated amounts due.

(3) The tentative settlement is subject to adjustment at the time of a final settlement.

§ 417.834 Responsibility for establishing administrative review procedures.

The HCPP is responsible for establishing and maintaining the administrative review procedures that are specified in §§ 417.830 through 417.840.

§ 417.836 Written description of administrative review procedures.

Each HCPP is responsible for ensuring that all Medicare enrollees are informed in writing of the administrative review procedures that are available to them.

§ 417.838 Organization determinations.

(a) Actions that are organization determinations. For purposes of §§ 417.830 through 417.840, an organization determination is a refusal to furnish or arrange for services, or reimburse the party for services provided to the beneficiary, on the grounds that the services are not covered by Medicare.

(b) Actions that are not organization determinations. The following are not organization determinations for purposes of §§ 417.830 through 417.840:

(1) A determination regarding services that were furnished by the HCPP, either directly or under arrangement, for which the enrollee has no further obligation for payment.

(2) A determination regarding services that are not covered under the HCPP’s agreement with HCFA.
§ 417.840 Administrative review procedures.

The HCPP must apply §§ 417.608 through 417.638 to organization determinations that affect its Medicare enrollees, and to reconsideration, hearings, Departmental Appeals Board review, and judicial review of those organization determinations.

[59 FR 59943, Nov. 21, 1994, as amended at 61 FR 32348, June 24, 1996]

Subpart V—Administration of Outstanding Loans and Loan Guarantees

§ 417.910 Applicability.

The regulations in this subpart apply, as appropriate, to public and private entities that have loans or loan guarantees that—
(a) Were awarded to them before October 1986 under section 1304 or section 1305 of the PHS Act; and
(b) Are still outstanding.

[59 FR 49842, Sept. 30, 1994]

§ 417.911 Definitions.

As used in this subpart—

Any 12-month period means the 12-month period beginning on the first day of any month.

Expansion of services means—(1) The addition of any health service not previously provided by or through the HMO, that requires an increase in the facilities, equipment, or health professionals of the HMO; or
(2) The improvement or upgrading of existing facilities or equipment, or an increase in the number of categories of health professionals, of the HMO so that the HMO could provide directly services that it previously provided through contract or referral or which it could not previously provide with its existing facilities or equipment.

First 60 months of operation or expansion means the 60-month period beginning on the first day of the month during which the HMO first provided services to enrollees, or in the case of significant expansion, first provided services in accordance with its expansion plan.

Health system agency means an entity that has been designated in accordance with section 1515 of the PHS Act; and the term State health planning and development agency means an agency that has been designated in accordance with section 1521 of the PHS Act.

Initial costs of operation means any cost incurred in the first 60 months of an operation or expansion that met any of the following requirements:
(1) Under generally accepted accounting principles or under accounting practices prescribed or permitted by State regulatory authority, was not a capital cost.
(2) Was required by State regulatory authority to meet reserves or tangible net equity requirements.
(3) Was for a payment made to reduce balance sheet liabilities existing at the beginning of the 60-month period, but only if—(i) The payment had been approved in writing by the Secretary; and
(ii) The total of these payments did not exceed 20 percent of the amount of the loan.
(4) Was for a small capital expenditure, but only if—(i) The cost had been approved in writing by the Secretary; and
(ii) The total of these costs did not exceed $200,000 in any 12-month period, and $400,000 during the first 60 months of operation or expansion.

Nonprofit as applied to a private entity, means a private agency, institution, or organization, no part of the net earnings of which inures, or may lawfully inure, to the benefit of any private shareholder or individual.

Significant expansion means—(1) A planned substantial increase in the enrollment of the HMO, that requires an increase in the number of health professionals serving enrollees of the HMO or an expansion of the physical capacity of the HMO's total health facilities; or
(2) A planned expansion of the service area beyond the current service area, that would be made possible by the addition of health service delivery facilities and health professionals to serve enrollees at a new site or sites in areas previously without service sites.

Small capital expenditure means expenditures for—(1) Equipment as defined in 45 CFR 74.132; or
(2) Alterations and renovations required to change the interior arrangements or other physical characteristics of an existing facility or installed equipment, so that it may be more effectively used for its currently designated purpose, or adapted to a changed use.


§ 417.920 Planning and initial development.

(a) Under section 1304 of the PHS Act, grants and loan guarantees were awarded for projects for planning and initial development of HMOs.

(b) Planning projects included projects for any of the following:

(1) Establishment of an HMO.

(2) Significant expansion of the HMO’s enrollment or geographic area.

(c) Initial development projects included projects for any of the following:

(1) Establishment of an HMO.

(2) Significant expansion of the HMO’s enrollment or geographic area.

(3) Expansion of the range or amount of services furnished by the HMO.

[58 FR 38076, July 15, 1993]

§ 417.930 Initial costs of operation.

Under section 1305 of the PHS, loans and loan guarantees were awarded for initial costs of operation of HMOs.

[58 FR 38077, July 15, 1993]

§ 417.931 [Reserved]

§ 417.934 Reserve requirement.

(a) Timing. Unless the Secretary approved a longer period, an entity that received a loan or loan guarantee under section 1305 of the PHS Act was required to establish a restricted reserve account on the earlier of the following:

(1) When the HMO’s revenues and costs of operation reached the break-even point.

(2) At the end of the 60-month period following the Secretary’s endorsement of the loan or loan guarantee.

(b) Purpose and amount of reserve. The reserve had to be constituted so as to accumulate, no later than 12 years after endorsement of the loan or loan guarantee, an amount equal to 1 year’s principal and interest.

[59 FR 49842, Sept. 30, 1994]

§ 417.937 Loan and loan guarantee provisions.

(a) Disbursement of loan proceeds. The principal amount of any loan made or guaranteed by the Secretary under this subpart was disbursed to the entity in accordance with an agreement entered into between the parties to the loan and approved by the Secretary.

(b) Length and maturity of loans. The principal amount of each loan or loan guarantee, together with interest thereon, is repayable over a period of 22 years, beginning on the date of endorsement of the loan, or loan guarantee by the Secretary. The Secretary could approve a shorter repayment period if he or she determined that a repayment period of less than 22 years is more appropriate to an entity’s total financial plan.

(c) Repayment. The principal amount of each loan or loan guarantee, together with interest thereon is repayable in accordance with a repayment schedule that is agreed upon by the parties to the loan or loan guarantee and approved by the Secretary before or at the time of endorsement of the loan. Unless otherwise specifically authorized by the Secretary, each loan made or guaranteed by the Secretary is repayable in substantially level combined installments of principal and interest to be paid at intervals not less frequently than annually, sufficient in amount to amortize the loan through the final year of the life of the loan. Principal repayment during the first 60 months of operation could be deferred with payment of interest only during that period. The Secretary could set rates of interest for each disbursement at a rate comparable to the rate of interest prevailing on the date of disbursement for marketable obligations of the United States of comparable maturities, adjusted to provide for appropriate administrative charges.

[59 FR 49842, Sept. 30, 1994]
§ 417.940 Civil action to enforce compliance with assurances.

The provisions of §417.163(g) apply to entities that have outstanding loans or loan guarantees administered under this subpart.

[59 FR 49843, Sept. 30, 1994]

PART 418—HOSPICE CARE

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AUTHORITY: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh).

SOURCE: 48 FR 56026, Dec. 16, 1983, unless otherwise noted.
Health Care Financing Administration, HHS  § 418.21

Subpart A—General Provision and Definitions

§ 418.1 Statutory basis.
This part implements section 1861(dd) of the Social Security Act. Section 1861(dd) specifies services covered as hospice care and the conditions that a hospice program must meet in order to participate in the Medicare program. The following sections of the Act are also pertinent:
(a) Sections 1812(a)(4) and (d) of the Act specify eligibility requirements for the individual and the benefit periods.
(b) Section 1813(a)(4) of the Act specifies coinsurance amounts.
(c) Sections 1814(a)(7) and 1814(i) of the Act contain conditions and limitations on coverage of, and payment for, hospice care.
(d) Sections 1862(a)(1), (6) and (9) of the Act establish limits on hospice coverage.


§ 418.2 Scope of part.
Subpart A of this part sets forth the statutory basis and scope and defines terms used in this part. Subpart B specifies the eligibility requirements and the benefit periods. Subpart C specifies conditions of participation for hospices. Subpart D describes the covered services and specifies the limits on services covered as hospice care. Subpart E specifies the reimbursement methods and procedures. Subpart F specifies coinsurance amounts applicable to hospice care.

§ 418.3 Definitions.
For purposes of this part—
Attending physician means a physician who—
(a) Is a doctor of medicine or osteopathy; and
(b) Is identified by the individual, at the time he or she elects to receive hospice care, as having the most significant role in the determination and delivery of the individual’s medical care.

Bereavement counseling means counseling services provided to the individual’s family after the individual’s death.

Cap period means the twelve-month period ending October 31 used in the application of the cap on overall hospice reimbursement specified in § 418.309.
Employee means an employee (defined by section 210(j) of the Act) of the hospice or, if the hospice is a subdivision of an agency or organization, an employee of the agency or organization who is appropriately trained and assigned to the hospice unit. “Employee” also refers to a volunteer under the jurisdiction of the hospice.
Hospice means a public agency or private organization or subdivision of either of these that—is primarily engaged in providing care to terminally ill individuals.
Physician means physician as defined in § 410.20 of this chapter.
Representative means an individual who has been authorized under State law to terminate medical care or to elect or revoke the election of hospice care on behalf of a terminally ill individual who is mentally or physically incapacitated.
Social worker means a person who has at least a bachelor’s degree from a school accredited or approved by the Council on Social Work Education.
Terminally ill means that the individual has a medical prognosis that his or her life expectancy is 6 months or less if the illness runs its normal course.


Subpart B—Eligibility, Election and Duration of Benefits

§ 418.20 Eligibility requirements.
In order to be eligible to elect hospice care under Medicare, an individual must be—
(a) Entitled to Part A of Medicare; and
(b) Certified as being terminally ill in accordance with § 418.22.

§ 418.21 Duration of hospice care coverage—Election periods.
(a) Subject to the conditions set forth in this part, an individual may elect to receive hospice care during one
§ 418.22 Certification of terminal illness.

(a) Timing of certification—(1) General rule. The hospice must obtain written certification of terminal illness for each of the periods listed in § 418.21, even if a single election continues in effect for two, three, or four periods, as provided in § 418.24(c).

(2) Basic requirement. Except as provided in paragraph (a)(3) of this section, the hospice must obtain the written certification no later than two calendar days after the period begins.

(3) Exception. For the initial 90-day period, if the hospice cannot obtain the written certifications within two calendar days, it must obtain oral certifications within two calendar days, and written certifications no later than eight calendar days after the period begins.

(b) Content of certification. The certification must specify that the individual's prognosis is for a life expectancy of 6 months or less if the terminal illness runs its normal course.

(c) Sources of certification. (1) For the initial 90-day period, the hospice must obtain written certification statements (and oral certification statements if required under paragraph (a)(3) of this section) from—

(i) The medical director of the hospice or the physician member of the hospice interdisciplinary group; and

(ii) The individual's attending physician if the individual has an attending physician.

(2) For subsequent periods, the only requirement is certification by one of the physicians listed in paragraph (c)(1)(i) of this section.

(d) Maintenance of records. Hospice staff must—

(1) Make an appropriate entry in the patient's medical record as soon as they receive an oral certification; and

(2) File written certifications in the medical record.


§ 418.24 Election of hospice care.

(a) Filing an election statement. An individual who meets the eligibility requirement of § 418.20 may file an election statement with a particular hospice. If the individual is physically or mentally incapacitated, his or her representative (as defined in § 418.3) may file the election statement.

(b) Content of election statement. The election statement must include the following:

(1) Identification of the particular hospice that will provide care to the individual.

(2) The individual's or representative's acknowledgement that he or she has been given a full understanding of the palliative rather than curative nature of hospice care, as it relates to the individual's terminal illness.

(3) Acknowledgement that certain Medicare services, as set forth in paragraph (d) of this section, are waived by the election.

(4) The effective date of the election, which may be the first day of hospice care or a later date, but may be no earlier than the date of the election statement.

(5) The signature of the individual or representative.

(c) Duration of election. An election to receive hospice care will be considered to continue through the initial election period and through the subsequent election periods without a break in care as long as the individual—

(1) Remains in the care of a hospice; and

(2) Does not revoke the election under the provisions of § 418.28.

(d) Waiver of other benefits. For the duration of an election of hospice care, an individual waives all rights to Medicare payments for the following services:

(1) Hospice care provided by a hospice other than the hospice designated by the individual (unless provided under
arrangements made by the designated hospice).

(2) Any Medicare services that are related to the treatment of the terminal condition for which hospice care was elected or a related condition or that are equivalent to hospice care except for services—

(i) Provided by the designated hospice;

(ii) Provided by another hospice under arrangements made by the designated hospice; and

(iii) Provided by the individual’s attending physician if that physician is not an employee of the designated hospice or receiving compensation from the hospice for those services.

(e) Re-election of hospice benefits. If an election has been revoked in accordance with §418.28, the individual (or his or her representative if the individual is mentally or physically incapacitated) may at any time file an election, in accordance with this section, for any other election period that is still available to the individual.

§418.28 Revoking the election of hospice care.

(a) An individual or representative may revoke the individual’s election of hospice care at any time during an election period.

(b) To revoke the election of hospice care, the individual or representative must file a statement with the hospice that includes the following information:

(1) A signed statement that the individual or representative revokes the individual’s election for Medicare coverage of hospice care for the remainder of that election period.

(2) The date that the revocation is to be effective. (An individual or representative may not designate an effective date earlier than the date that the revocation is made).

(c) An individual, upon revocation of the election of Medicare coverage of hospice care for a particular election period—

(1) Is no longer covered under Medicare for hospice care;

(2) Resumes Medicare coverage of the benefits waived under §418.24(e)(2); and

(3) May at any time elect to receive hospice coverage for any other hospice election periods that he or she is eligible to receive.

§418.30 Change of the designated hospice.

(a) An individual or representative may change, once in each election period, the designation of the particular hospice from which hospice care will be received.

(b) The change of the designated hospice is not a revocation of the election for the period in which it is made.

(c) To change the designation of hospice programs, the individual or representative must file, with the hospice from which care has been received and with the newly designated hospice, a statement that includes the following information:

(1) The name of the hospice from which the individual has received care and the name of the hospice from which he or she plans to receive care.

(2) The date the change is to be effective.

Subpart C—Conditions of Participation—General Provisions and Administration

§418.50 Condition of participation—General provisions.

(a) Standard: Compliance. A hospice must maintain compliance with the conditions of this subpart and subparts D and E of this part.

(b) Standard: Required services. A hospice must be primarily engaged in providing the care and services described in §418.202, must provide bereavement counseling and must—

(1) Make nursing services, physician services, and drugs and biologicals routinely available on a 24-hour basis;

(2) Make all other covered services available on a 24-hour basis to the extent necessary to meet the needs of individuals for care that is reasonable and necessary for the palliation and management of terminal illness and related conditions; and

(3) Provide these services in a manner consistent with accepted standards of practice.

(c) Standard: Disclosure of information. The hospice must meet the disclosure
§ 418.52 Condition of participation—Governing body.

A hospice must have a governing body that assumes full legal responsibility for determining, implementing and monitoring policies governing the hospice's total operation. The governing body must designate an individual who is responsible for the day to day management of the hospice program. The governing body must also ensure that all services provided are consistent with accepted standards of practice.

§ 418.54 Condition of participation—Medical director.

The medical director must be a hospice employee who is a doctor of medicine or osteopathy who assumes overall responsibility for the medical component of the hospice's patient care program.

§ 418.56 Condition of participation—Professional management.

Subject to the conditions of participation pertaining to services in §§ 418.80 and 418.90, a hospice may arrange for another individual or entity to furnish services to the hospice's patients. If services are provided under arrangement, the hospice must meet the following standards:

(a) Standard: Continuity of care. The hospice program assures the continuity of patient/family care in home, outpatient, and inpatient settings.

(b) Standard: Written agreement. The hospice has a legally binding written agreement for the provision of arranged services. The agreement includes at least the following:

(1) Identification of the services to be provided.

(2) A stipulation that services may be provided only with the express authorization of the hospice.

(3) The manner in which the contracted services are coordinated, supervised, and evaluated by the hospice.

(4) The delineation of the role(s) of the hospice and the contractor in the admission process, patient/familyassessment, and the interdisciplinary group care conferences.

(5) Requirements for documenting that services are furnished in accordance with the agreement.

(6) The qualifications of the personnel providing the services.

(c) Standard: Professional management responsibility. The hospice retains professional management responsibility for those services and ensures that they are furnished in a safe and effective manner by persons meeting the qualifications of this part, and in accordance with the patient's plan of care and the other requirements of this part.

(d) Standard: Financial responsibility. The hospice retains responsibility for payment for services.

(e) Standard: Inpatient care. The hospice ensures that inpatient care is furnished only in a facility which meets the requirements in § 418.98 and its arrangement for inpatient care is described in a legally binding written agreement that meets the requirements of paragraph (b) and that also specifies, at a minimum—

(1) That the hospice furnishes to the inpatient provider a copy of the patient's plan of care and specifies the inpatient services to be furnished;

(2) That the inpatient provider has established policies consistent with those of the hospice and agrees to abide by the patient care protocols established by the hospice for its patients;

(3) That the medical record includes a record of all inpatient services and events and that a copy of the discharge summary and, if requested, a copy of the medical record are provided to the hospice;

(4) The party responsible for the implementation of the provisions of the agreement; and

(5) That the hospice retains responsibility for appropriate hospice care training of the personnel who provide the care under the agreement.

§ 418.58 Condition of participation—Plan of care.

A written plan of care must be established and maintained for each individual admitted to a hospice program,
§ 418.70 Condition of participation—Volunteers.

The hospice in accordance with the numerical standards, specified in paragraph (e) of this section, uses volunteers, in defined roles, under the supervision of a designated hospice employee.

(a) Standard: Training. The hospice must provide appropriate orientation for the quality assurance program must—

(a) Implement and report on activities and mechanisms for monitoring the quality of patient care;

(b) Identify and resolve problems; and

(c) Make suggestions for improving patient care.

§ 418.68 Condition of participation—Interdisciplinary group.

A hospice may not discontinue or diminish care provided to a Medicare beneficiary because of the beneficiary’s inability to pay for that care.

§ 418.62 Condition of participation—Informed consent.

A hospice must demonstrate respect for an individual’s rights by ensuring that an informed consent form that specifies the type of care and services that may be provided as hospice care during the course of the illness has been obtained for every individual, either from the individual or representative as defined in §418.3.

§ 418.64 Condition of participation—Inservice training.

A hospice must provide an ongoing program for the training of its employees.

§ 418.66 Condition of participation—Quality assurance.

A hospice must conduct an ongoing, comprehensive, integrated, self-assessment of the quality and appropriateness of care provided, including inpatient care, home care and care provided under arrangements. The findings are used by the hospice to correct identified problems and to revise hospice policies if necessary. Those responsible

and the care provided to an individual must be in accordance with the plan.

(a) Standard: Establishment of plan. The plan must be established by the attending physician, the medical director or physician designee and interdisciplinary group prior to providing care.

(b) Standard: Review of plan. The plan must be reviewed and updated, at intervals specified in the plan, by the attending physician, the medical director or physician designee and interdisciplinary group. These reviews must be documented.

(c) Standard: Content of plan. The plan must include an assessment of the individual’s needs and identification of the services including the management of discomfort and symptom relief. It must state in detail the scope and frequency of services needed to meet the patient’s and family’s needs.

§ 418.60 Condition of participation—Continuation of care.

A hospice must demonstrate respect for an individual’s rights by ensuring that an informed consent form that specifies the type of care and services that may be provided as hospice care during the course of the illness has been obtained for every individual, either from the individual or representative as defined in §418.3.

§ 418.62 Condition of participation—Informed consent.

A hospice must demonstrate respect for an individual’s rights by ensuring that an informed consent form that specifies the type of care and services that may be provided as hospice care during the course of the illness has been obtained for every individual, either from the individual or representative as defined in §418.3.

§ 418.64 Condition of participation—Inservice training.

A hospice must provide an ongoing program for the training of its employees.

§ 418.66 Condition of participation—Quality assurance.

A hospice must conduct an ongoing, comprehensive, integrated, self-assessment of the quality and appropriateness of care provided, including inpatient care, home care and care provided under arrangements. The findings are used by the hospice to correct identified problems and to revise hospice policies if necessary. Those responsible for the quality assurance program must—

(a) Implement and report on activities and mechanisms for monitoring the quality of patient care;

(b) Identify and resolve problems; and

(c) Make suggestions for improving patient care.

§ 418.68 Condition of participation—Interdisciplinary group.

A hospice must designate an interdisciplinary group or groups composed of individuals who provide or supervise the care and services offered by the hospice.

(a) Standard: Composition of group. The hospice must have an interdisciplinary group or groups that include at least the following individuals who are employees of the hospice:

(1) A doctor of medicine or osteopathy.

(2) A registered nurse.

(3) A social worker.

(4) A pastoral or other counselor.

(b) Standard: Role of group. The interdisciplinary group is responsible for—

(1) Participation in the establishment of the plan of care;

(2) Provision or supervision of hospice care and services;

(3) Periodic review and updating of the plan of care for each individual receiving hospice care; and

(4) Establishment of policies governing the day-to-day provision of hospice care and services.

(c) If a hospice has more than one interdisciplinary group, it must designate in advance the group it chooses to execute the functions described in paragraph (b)(4) of this section.

(d) Standard: Coordinator. The hospice must designate a registered nurse to coordinate the implementation of the plan of care for each patient.

§ 418.70 Condition of participation—Volunteers.

The hospice in accordance with the numerical standards, specified in paragraph (e) of this section, uses volunteers, in defined roles, under the supervision of a designated hospice employee.

(a) Standard: Training. The hospice must provide appropriate orientation
and training that is consistent with acceptable standards of hospice practice.

(b) Standard: Role. Volunteers must be used in administrative or direct patient care roles.

(c) Standard: Recruiting and retaining. The hospice must document active and ongoing efforts to recruit and retain volunteers.

(d) Standard: Cost saving. The hospice must document the cost savings achieved through the use of volunteers. Documentation must include—

1. The identification of necessary positions which are occupied by volunteers;
2. The work time spent by volunteers occupying those positions; and
3. Estimates of the dollar costs which the hospice would have incurred if paid employees occupied the positions identified in paragraph (d)(1) for the amount of time specified in paragraph (d)(2).

(e) Standard: Level of activity. A hospice must document and maintain a volunteer staff sufficient to provide administrative or direct patient care in an amount that, at a minimum, equals 5 percent of the total patient care hours of all paid hospice employees and contract staff. The hospice must document a continuing level of volunteer activity. Expansion of care and services achieved through the use of volunteers, including the type of services and the time worked, must be recorded.

(f) Standard: Availability of clergy. The hospice must make reasonable efforts to arrange for visits of clergy and other members of religious organizations in the community to patients who request such visits and must advise patients of this opportunity.

§ 418.72 Condition of participation—Licensure.

The hospice and all hospice employees must be licensed in accordance with applicable Federal, State and local laws and regulations.

(a) Standard: Licensure of program. If State or local law provides for licensing of hospices, the hospice must be licensed.

(b) Standard: Licensure of employees. Employees who provide services must be licensed, certified or registered in accordance with applicable Federal or State laws.

§ 418.74 Condition of participation—Central clinical records.

In accordance with accepted principles of practice, the hospice must establish and maintain a clinical record for every individual receiving care and services. The record must be complete, promptly and accurately documented, readily accessible and systematically organized to facilitate retrieval.

(a) Standard: Content. Each clinical record is a comprehensive compilation of information. Entries are made for all services provided. Entries are made and signed by the person providing the services. The record includes all services whether furnished directly or under arrangements made by the hospice. Each individual's record contains—

1. The initial and subsequent assessments;
2. The plan of care;
3. Identification data;
4. Consent and authorization and election forms;
5. Pertinent medical history; and
6. Complete documentation of all services and events (including evaluations, treatments, progress notes, etc.).

(b) Standard: Protection of information. The hospice must safeguard the clinical record against loss, destruction and unauthorized use.

Subpart D—Conditions of Participation: Core Services

§ 418.80 Condition of participation—Furnishing of core services.

Except as permitted in §418.83, a hospice must ensure that substantially all the core services described in this subpart are routinely provided directly by hospice employees. A hospice may use contracted staff if necessary to supplement hospice employees in order to meet the needs of patients during periods of peak patient loads or under extraordinary circumstances. If contracting is used, the hospice must maintain professional, financial, and administrative responsibility for the services and must assure that the
§ 418.82 Condition of participation—Nursing services.

The hospice must provide nursing care and services by or under the supervision of a registered nurse.

(a) Nursing services must be directed and staffed to assure that the nursing needs of patients are met.

(b) Patient care responsibilities of nursing personnel must be specified.

(c) Services must be provided in accordance with recognized standards of practice.

§ 418.83 Nursing services—Waiver of requirement that substantially all nursing services be routinely provided directly by a hospice.

(a) HCFA may approve a waiver of the requirement in §418.80 for nursing services provided by a hospice which is located in a non-urbanized area. The location of a hospice that operates in several areas is considered to be the location of its central office. The hospice must provide evidence that it was operational on or before January 1, 1983, and that it made a good faith effort to hire a sufficient number of nurses to provide services directly. HCFA bases its decision as to whether to approve a waiver application on the following:

1. The current Bureau of the Census designations for determining non-urbanized areas.

2. Evidence that a hospice was operational on or before January 1, 1983 including:
   (i) Proof that the organization was established to provide hospice services on or before January 1, 1983;
   (ii) Evidence that hospice-type services were furnished to patients on or before January 1, 1983; and
   (iii) Evidence that the hospice care was a discrete activity rather than an aspect of another type of provider’s patient care program on or before January 1, 1983.

3. Evidence that a hospice made a good faith effort to hire nurses, including:
   (i) Copies of advertisements in local newspapers that demonstrate recruitment efforts;
   (ii) Job descriptions for nurse employees;
   (iii) Evidence that salary and benefits are competitive for the area; and
   (iv) Evidence of any other recruiting activities (e.g., recruiting efforts at health fairs and contacts with nurses at other providers in the area);

(b) Any waiver request is deemed to be granted unless it is denied within 60 days after it is received.

(c) Waivers will remain effective for one year at a time.

(d) HCFA may approve a maximum of two one-year extensions for each initial waiver. If a hospice wishes to receive a one-year extension, the hospice must submit a certification to HCFA, prior to the expiration of the waiver period, that the employment market for nurses has not changed significantly since the time the initial waiver was granted.

§ 418.84 Condition of participation—Medical social services.

Medical social services must be provided by a qualified social worker, under the direction of a physician.

§ 418.86 Condition of participation—Physician services.

In addition to palliation and management of terminal illness and related conditions, physician employees of the hospice, including the physician member(s) of the interdisciplinary group, must also meet the general medical needs of the patients to the extent that these needs are not met by the attending physician.

§ 418.88 Condition of participation—Counseling services.

Counseling services must be available to both the individual and the family. Counseling includes bereavement counseling, provided after the patient’s death as well as dietary, spiritual and any other counseling services for the individual and family provided while the individual is enrolled in the hospice.

(a) Standard: Bereavement counseling. There must be an organized program
§418.90  Condition of participation—Furnishing of other services.

A hospice must ensure that the services described in this subpart are provided directly by hospice employees or under arrangements made by the hospice as specified in §418.56.

§418.92  Condition of participation—Physical therapy, occupational therapy, and speech-language pathology.

(a) Physical therapy services, occupational therapy services, and speech-language pathology services must be available, and when provided, offered in a manner consistent with accepted standards of practice.

(b) (1) If the hospice engages in laboratory testing outside of the context of assisting an individual in self-administering a test with an appliance that has been cleared for that purpose by the FDA, such testing must be in compliance with all applicable requirements of part 493 of this chapter.

(2) If the hospice chooses to refer specimens for laboratory testing to another laboratory, the referral laboratory must be certified in the appropriate specialties and subspecialties of services in accordance with the applicable requirements of part 493 of this chapter.

[57 FR 7135, Feb. 28, 1992]

§418.94  Condition of participation—Home health aide and homemaker services.

Home health aide and homemaker services must be available and adequate in frequency to meet the needs of the patients. A home health aide is a person who meets the training, attitude and skill requirements specified in §484.36 of this chapter.

(a) Standard: Supervision. A registered nurse must visit the home site at least every two weeks when aide services are being provided, and the visit must include an assessment of the aide services.

(b) Standard: Duties. Written instructions for patient care are prepared by a registered nurse. Duties include, but may not be limited to, the duties specified in §484.36(c) of this chapter.


§418.96  Condition of participation—Medical supplies.

Medical supplies and appliances including drugs and biologicals, must be provided as needed for the palliation and management of the terminal illness and related conditions.

(a) Standard: Administration. All drugs and biologicals must be administered in accordance with accepted standards of practice.

(b) Standard: Controlled drugs in the patient’s home. The hospice must have a policy for the disposal of controlled drugs maintained in the patient’s home when those drugs are no longer needed by the patient.

(c) Standard: Administration of drugs and biologicals. Drugs and biologicals are administered only by the following individuals:

(1) A licensed nurse or physician.

(2) An employee who has completed a State-approved training program in medication administration.

(3) The patient if his or her attending physician has approved.

(4) Any other individual in accordance with applicable State and local
laws. The persons, and each drug and biological they are authorized to administer, must be specified in the patient's plan of care.

§ 418.98 Condition of participation—Short term inpatient care.

Inpatient care must be available for pain control, symptom management and respite purposes, and must be provided in a participating Medicare or Medicaid facility.

(a) Standard: Inpatient care for symptom control. Inpatient care for pain control and symptom management must be provided in one of the following:

(1) A hospice that meets the condition of participation for providing inpatient care directly as specified in § 418.100.

(2) A hospital or an SNF that also meets the standards specified in § 418.100 (a) and (e) regarding 24-hour nursing service and patient areas.

(b) Standard: Inpatient care for respite purposes. Inpatient care for respite purposes must be provided by one of the following:

(1) A provider specified in paragraph (a) of this section.

(2) An ICF that also meets the standards specified in § 418.100 (a) and (e) regarding 24-hour nursing service and patient areas.

(c) Standard: Inpatient care limitation. The total number of inpatient days used by Medicare beneficiaries who elected hospice coverage in any 12-month period preceding a certification survey in a particular hospice may not exceed 20 percent of the total number of hospice days for this group of beneficiaries.

(d) Standard: Exemption from limitation. Until October 1, 1986, any hospice that began operation before January 1, 1975 is not subject to the limitation specified in paragraph (c).


§ 418.100 Condition of participation—Hospices that provide inpatient care directly.

A hospice that provides inpatient care directly must comply with all of the following standards.

(a) Standard: Twenty-four-hour nursing services. (1) The facility provides 24-hour nursing services which are sufficient to meet total nursing needs and which are in accordance with the patient plan of care. Each patient receives treatments, medications, and diet as prescribed, and is kept comfortable, clean, well-groomed, and protected from accident, injury, and infection.

(2) Each shift must include a registered nurse who provides direct patient care.

(b) Standard: Disaster preparedness. The hospice has an acceptable written plan, periodically rehearsed with staff, with procedures to be followed in the event of an internal or external disaster and for the care of casualties (patients and personnel) arising from such disasters.

(c) Standard: Health and safety laws. The hospice must meet all Federal, State, and local laws, regulations, and codes pertaining to health and safety, such as provisions regulating—

(1) Construction, maintenance, and equipment for the hospice;

(2) Sanitation;

(3) Communicable and reportable diseases; and

(4) Post mortem procedures.

(d) Standard: Fire protection. (1) Except as provided in paragraphs (d) (2) and (3) of this section, the hospice must meet the provisions of the 1985 edition of the Life Safety Code of the National Fire Protection Association (which is incorporated by reference) that are applicable to hospices.

(2) In consideration of a recommendation by the State survey agency, HCFA may waive, for periods deemed appropriate, specific provisions of the Life Safety Code which, if rigidly applied would result in unreasonable hardship for the hospice, but only if the waiver would not adversely affect the health and safety of the patients.

(3) Any hospice that, on May 9, 1988, complies with the requirements of the 1981 edition of the Life Safety Code, with or without waivers, will be considered to be in compliance with this standard, as long as the hospice continues to remain in compliance with that edition of the Life Safety Code.

1 See footnote to § 405.1134(a) of this chapter.
§418.100

(4) Any facility of two or more stories that is not of fire resistive construction and is participating on the basis of a waiver of construction type or height, may not house blind, non-ambulatory, or physically handicapped patients above the street-level floor unless the facility—

(i) Is one of the following construction types (as defined in the Life Safety Code):

(A) Type II (1, 1, 1)—protected non-combustible.

(B) Fully sprinklered Type II (0, 0, 0)—non-combustible.

(C) Fully sprinklered Type III (2, 1, 1)—protected ordinary.

(D) Fully sprinklered Type V (1, 1, 1)—protected wood frame; or

(ii) Achieves a passing score on the Fire Safety Evaluation System (FSES).

(e) Standard: Patient areas.

(1) The hospice must design and equip areas for the comfort and privacy of each patient and family members.

(2) The hospice must have—

(i) Physical space for private patient/family visiting;

(ii) Accommodations for family members to remain with the patient throughout the night;

(iii) Accommodations for family privacy after a patient’s death; and

(iv) Decor which is homelike in design and function.

(3) Patients must be permitted to receive visitors at any hour, including small children.

(f) Standard: Patient rooms and toilet facilities.

Patient rooms are designed and equipped for adequate nursing care and the comfort and privacy of patients.

(1) Each patient’s room must—

(i) Be equipped with or conveniently located near toilet and bathing facilities;

(ii) Be at or above grade level;

(iii) Contain a suitable bed for each patient and other appropriate furniture;

(iv) Have closet space that provides security and privacy for clothing and personal belongings;

(v) Contain no more than four beds;

(vi) Measure at least 100 square feet for a single patient room or 80 square feet for each patient for a multipatient room; and

(vii) Be equipped with a device for calling the staff member on duty.

(2) For an existing building, HCFA may waive the space and occupancy requirements of paragraphs (f)(1)(v) and (vi) of this section for as long as it is considered appropriate if it finds that—

(i) The requirements would result in unreasonable hardship on the hospice if strictly enforced; and

(ii) The waiver serves the particular needs of the patients and does not adversely affect their health and safety.

(g) Standard: Bathroom facilities. The hospice must—

(1) Provide an adequate supply of hot water at all times for patient use; and

(2) Have plumbing fixtures with control valves that automatically regulate the temperature of the hot water used by patients.

(h) Standard: Linen. The hospice has available at all times a quantity of linen essential for proper care and comfort of patients. Linens are handled, stored, processed, and transported in such a manner as to prevent the spread of infection.

(i) Standard: Isolation areas. The hospice must make provision for isolating patients with infectious diseases.

(j) Standard: Meal service, menu planning, and supervision.

(1) Serve at least three meals or their equivalent each day at regular times, with not more than 14 hours between a substantial evening meal and breakfast;

(2) Procure, store, prepare, distribute, and serve all food under sanitary conditions;

(3) Have a staff member trained or experienced in food management or nutrition who is responsible for—

(i) Planning menus that meet the nutritional needs of each patient, following the orders of the patient’s physician and, to the extent medically possible, the recommended dietary allowances of the Food and Nutrition Board of the National Research Council, National Academy of Sciences (Recommended Dietary Allowances (9th ed., 1981) is available from the Printing and Publications Office, National Academy of Sciences, Washington, DC 20418); and
(ii) Supervising the meal preparation and service to ensure that the menu plan is followed; and

(4) If the hospice has patients who require medically prescribed special diets, have the menus for those patients planned by a professionally qualified dietitian and supervise the preparation and serving of meals to ensure that the patient accepts the special diet.

(k) Standard: Pharmaceutical services. The hospice provides appropriate methods and procedures for the dispensing and administering of drugs and biologicals. Whether drugs and biologicals are obtained from community or institutional pharmacists or stocked by the facility, the facility is responsible for drugs and biologicals for its patients, insofar as they are covered under the program and for ensuring that pharmaceutical services are provided in accordance with accepted professional principles and appropriate Federal, State, and local laws. (See §405.1124(g), (h), and (i) of this chapter.)

(1) Licensed pharmacist. The hospice must—

(i) Employ a licensed pharmacist; or

(ii) Have a formal agreement with a licensed pharmacist to advise the hospice on ordering, storage, administration, disposal, and recordkeeping of drugs and biologicals.

(2) Orders for medications. (i) A physician must order all medications for the patient.

(ii) If the medication order is verbal—

(A) The physician must give it only to a licensed nurse, pharmacist, or another physician; and

(B) The individual receiving the order must record and sign it immediately and have the prescribing physician sign it in a manner consistent with good medical practice.

(3) Administering medications. Medications are administered only by one of the following individuals:

(i) A licensed nurse or physician.

(ii) An employee who has completed a State-approved training program in medication administration.

(iii) The patient if his or her attending physician has approved.

(4) Control and accountability. The pharmaceutical service has procedures for control and accountability of all drugs and biologicals throughout the facility. Drugs are dispensed in compliance with Federal and State laws. Records of receipt and disposition of all controlled drugs are maintained in sufficient detail to enable an accurate reconciliation. The pharmacist determines that drug records are in order and that an account of all controlled drugs is maintained and reconciled.

(5) Labeling of drugs and biologicals. The labeling of drugs and biologicals is based on currently accepted professional principles, and includes the appropriate accessory and cautionary instructions, as well as the expiration date when applicable.

(6) Storage. In accordance with State and Federal laws, all drugs and biologicals are stored in locked compartments under proper temperature controls and only authorized personnel have access to the keys. Separately locked compartments are provided for storage of controlled drugs listed in Schedule II of the Comprehensive Drug Abuse Prevention & Control Act of 1970 and other drugs subject to abuse, except under single unit package drug distribution systems in which the quantity stored is minimal and a missing dose can be readily detected. An emergency medication kit is kept readily available.

(7) Drug disposal. Controlled drugs no longer needed by the patient are disposed of in compliance with State requirements. In the absence of State requirements, the pharmacist and a registered nurse dispose of the drugs and prepare a record of the disposal.

§ 418.202 Covered services.

All services must be performed by appropriately qualified personnel, but it is the nature of the service, rather than the qualification of the person who provides it, that determines the coverage category of the service. The following services are covered hospice services:

(a) Nursing care provided by or under the supervision of a registered nurse.

(b) Medical social services provided by a social worker under the direction of a physician.

(c) Physicians' services performed by a physician as defined in §410.20 of this chapter except that the services of the hospice medical director or the physician member of the interdisciplinary group must be performed by a doctor of medicine or osteopathy.

(d) Counseling services provided to the terminally ill individual and the family members or other persons caring for the individual at home. Counseling, including dietary counseling, may be provided both for the purpose of training the individual's family or other caregiver to provide care, and for the purpose of helping the individual and those caring for him or her to adjust to the individual's approaching death.

(e) Short-term inpatient care provided in a participating hospice inpatient unit, or a participating hospital or SNF, that additionally meets the standards in §418.202(a) and (e) regarding staffing and patient areas. Services provided in an inpatient setting must conform to the written plan of care. Inpatient care may be required for procedures necessary for pain control or acute or chronic symptom management.

Inpatient care may also be furnished as a means of providing respite for the individual's family or other persons caring for the individual at home. Respite care must be furnished as specified in §418.108(b). Payment for inpatient care will be made at the rate appropriate to the level of care as specified in §418.302.

(f) Medical appliances and supplies, including drugs and biologicals. Only drugs as defined in section 1861(t) of the Act and which are used primarily for the relief of pain and symptom control related to the individual's terminal illness are covered. Appliances may include covered durable medical equipment as described in §410.38 of this chapter as well as other self-help and personal comfort items related to the palliation or management of the patient's terminal illness. Equipment is provided by the hospice for use in the patient's home while he or she is under hospice care. Medical supplies include those that are part of the written plan of care.

(g) Home health aide services furnished by qualified aides as designated in §418.94 and homemaker services. Home health aides may provide personal care services as defined in §409.45(b) of this chapter. Aides may perform household services to maintain a safe and sanitary environment in areas of the home used by the patient, such as changing bed linens or light cleaning and laundering essential to the comfort and cleanliness of the patient. Aide services must be provided under the general supervision of a registered nurse. Homemaker services may include assistance in maintenance of a safe and healthy environment and services to enable the individual to carry out the treatment plan.

(h) Physical therapy, occupational therapy and speech-language pathology services in addition to the services described in §409.33 (b) and (c) of this chapter provided for purposes of symptom control or to enable the patient to maintain activities of daily living and basic functional skills.

§ 418.204 Special coverage requirements.

(a) Periods of crisis. Nursing care may be covered on a continuous basis for as much as 24 hours a day during periods of crisis as necessary to maintain an individual at home. Either homemaker or home health aide services or both may be covered on a 24-hour continuous basis during periods of crisis but...
Health Care Financing Administration, HHS § 418.302

(b) Respite care. (1) Respite care is short-term inpatient care provided to the individual only when necessary to relieve the family members or other persons caring for the individual.

(2) Respite care may be provided only on an occasional basis and may not be reimbursed for more than five consecutive days at a time.

(d) Bereavement counseling. Bereavement counseling is a required hospice service but it is not reimbursable.


Subpart G—Payment for Hospice Care

§ 418.301 Basic rules.

(a) Medicare payment for covered hospice care is made in accordance with the method set forth in §418.302.

(b) Medicare reimbursement to a hospice in a cap period is limited to a cap amount specified in §418.309.


§ 418.302 Payment procedures for hospice care.

(a) HCFA establishes payment amounts for specific categories of covered hospice care.

(b) Payment amounts are determined within each of the following categories:

1. Routine home care day. A routine home care day is a day on which an individual who has elected to receive hospice care is at home and is not receiving continuous care as defined in paragraph (b)(2) of this section.

2. Continuous home care day. A continuous home care day is a day on which an individual who has elected to receive hospice care is not in an inpatient facility and receives hospice care consisting predominantly of nursing care on a continuous basis at home. Home health aide or homemaker services or both may also be provided on a continuous basis. Continuous home care is only furnished during brief periods of crisis as described in §418.204(a) and only as necessary to maintain the terminally ill patient at home.

3. Inpatient respite care day. An inpatient respite care day is a day on which the individual who has elected hospice care receives care in an approved facility on a short-term basis for respite.

4. General inpatient care day. A general inpatient care day is a day on which an individual who has elected hospice care receives general inpatient care in an inpatient facility for pain control or acute or chronic symptom management which cannot be managed in other settings.

(c) The payment amounts for the categories of hospice care are fixed payment rates that are established by HCFA in accordance with the procedures described in §418.306. Payment rates are determined for the following categories:

1. Routine home care.

2. Continuous home care.

3. Inpatient respite care.


(d) The intermediary reimburses the hospice at the appropriate payment amount for each day for which an eligible Medicare beneficiary is under the hospice's care.

(e) The intermediary makes payment according to the following procedures:

1. Payment is made to the hospice for each day during which the beneficiary is eligible and under the care of the hospice, regardless of the amount of services furnished on any given day.

2. Payment is made for only one of the categories of hospice care described in §418.302(b) for any particular day.

3. On any day on which the beneficiary is not an inpatient, the hospice is paid the routine home care rate, unless the patient receives continuous care as defined in paragraph (b)(2) of this section for a period of at least 8 hours. In that case, a portion of the continuous care day rate is paid in accordance with paragraph (e)(4) of this section.

4. The hospice payment on a continuous care day varies depending on the number of hours of continuous services provided. The continuous home care rate is divided by 24 to yield an hourly rate. The number of hours of continuous care provided during a continuous
home care day is then multiplied by the hourly rate to yield the continuous home care payment for that day. A minimum of 8 hours of care must be furnished on a particular day to qualify for the continuous home care rate.

(5) Subject to the limitations described in paragraph (f) of this section, on any day on which the beneficiary is an inpatient in an approved facility for inpatient care, the appropriate inpatient rate (general or respite) is paid depending on the category of care furnished. The inpatient rate (general or respite) is paid for the date of admission and all subsequent inpatient days, except the day on which the patient is discharged. For the day of discharge, the appropriate home care rate is paid unless the patient dies as an inpatient. In the case where the beneficiary is discharged deceased, the inpatient rate (general or respite) is paid for the discharge day. Payment for inpatient respite care is subject to the requirement that it may not be provided consecutively for more than 5 days at a time.

Payment for the sixth and any subsequent day of respite care is made at the routine home care rate.

(f) Payment for inpatient care is limited as follows: (1) The total payment to the hospice for inpatient care (general or respite) is subject to a limitation that total inpatient care days for Medicare patients not exceed 20 percent of the total days for which these patients had elected hospice care.

(2) At the end of a cap period, the intermediary calculates a limitation on payment for inpatient care to ensure that Medicare payment is not made for days of inpatient care in excess of 20 percent of the total number of days of hospice care furnished to Medicare patients.

(3) If the number of days of inpatient care furnished to Medicare patients is equal to or less than 20 percent of the total days of hospice care to Medicare patients, no adjustment is necessary. Overall payments to a hospice are subject to the cap amount specified in §418.309.

(4) If the number of days of inpatient care furnished to Medicare patients exceeds 20 percent of the total days of hospice care to Medicare patients, the total payment for inpatient care is determined in accordance with the procedures specified in paragraph (f)(5) of this section. That amount is compared to actual payments for inpatient care, and any excess reimbursement must be refunded by the hospice. Overall payments to the hospice are subject to the cap amount specified in §418.309.

(5) If a hospice exceeds the number of inpatient care days described in paragraph (f)(4), the total payment for inpatient care is determined as follows:

(i) Calculate the ratio of the maximum number of allowable inpatient days to the actual number of inpatient care days furnished by the hospice to Medicare patients.

(ii) Multiply this ratio by the total reimbursement for inpatient care made by the intermediary.

(iii) Multiply the number of actual inpatient days in excess of the limitation by the routine home care rate.

(iv) Add the amounts calculated in paragraphs (f)(5)(ii) and (iii) of this section.


§ 418.304 Payment for physician services.

(a) The following services performed by hospice physicians are included in the rates described in §418.302:

(1) General supervisory services of the medical director.

(2) Participation in the establishment of plans of care, supervision of care and services, periodic review and updating of plans of care, and establishment of governing policies by the physician member of the interdisciplinary group.

(b) For services not described in paragraph (a) of this section, a specified Medicare contractor pays the hospice an amount equivalent to 100 percent of the physician's reasonable charge for those physician services furnished by hospice employees or under arrangements with the hospice. Reimbursement for these physician services is included in the amount subject to the hospice payment limit described in §418.309. Services furnished voluntarily by physicians are not reimbursable.
§ 418.306 Determination of payment rates.

(a) Applicability. HCFA establishes payment rates for each of the categories of hospice care described in § 418.302(b). The rates are established using the methodology described in section 1814(i)(1)(C) of the Act.

(b) Payment rates. The payment rates for routine home care and other services included in hospice care are as follows:

1. The following rates, which are 120 percent of the rates in effect on September 30, 1989, are effective January 1, 1990 through September 30, 1990 and October 21, 1990 through December 31, 1990:

   - Routine home care: $75.80
   - Continuous home care:
     - Full rate for 24 hours: $422.40
     - Hourly rate: $18.43
   - Inpatient respite care: $78.40
   - General inpatient care: $337.20

2. Except for the period beginning October 21, 1990, through December 31, 1990, the payment rates for routine home care and other services included in hospice care for Federal fiscal years 1991, 1992, and 1993 and those that begin on or after October 1, 1997, are the payment rates in effect under this paragraph during the previous fiscal year increased by the market basket percentage increase as defined in section 1814(b)(3)(B)(iii) of the Act, otherwise applicable to discharges occurring in the fiscal year. The payment rates for the period beginning October 21, 1990, through December 31, 1990, are the same as those shown in paragraph (b)(1) of this section.

3. For Federal fiscal years 1994 through 1997, the payment rate is the payment rate in effect during the previous fiscal year increased by a factor equal to the market basket percentage increase minus—

   - (i) 2 percentage points in FY 1994;
   - (ii) 1.5 percentage points in FYs 1995 and 1996; and
   - (iii) 0.5 percentage points in FY 1997.

(c) Adjustment for wage differences. HCFA will issue annually, in the Federal Register, a hospice wage index based on the most current available HCFA hospital wage data, including any changes to the definitions of Metropolitan Statistical Areas. The payment rates established by HCFA are adjusted by the intermediary to reflect local differences in wages according to the revised wage index.

(d) Federal Register notices. HCFA publishes as a notice in the Federal Register any proposal to change the methodology for determining the payment rates.


§ 418.307 Periodic interim payments.

Subject to the provisions of § 413.64(h) of this chapter, a hospice may elect to receive periodic interim payments (PIP) effective with claims received on or after July 1, 1987. Payment is made biweekly under the PIP method unless the hospice requests a longer fixed interval (not to exceed one month) between payments. The biweekly interim payment amount is based on the total estimated Medicare payments for the reporting period (as described in §§ 418.302-418.306). Each payment is made 2 weeks after the end of a biweekly period of service as described in § 413.64(h)(5) of this chapter. Under certain circumstances that are described in § 413.64(g) of this chapter, a hospice that is not receiving PIP may request an accelerated payment.

[59 FR 36713, July 12, 1994]

§ 418.308 Limitation on the amount of hospice payments.

(a) Except as specified in paragraph (b) of this section, the total Medicare payment to a hospice for care furnished during a cap period is limited by the hospice cap amount specified in § 418.309.
§ 418.309 Hospice cap amount.

(a) The cap amount is $6,500 per year and is adjusted for inflation or deflation for cap years that end after October 1, 1984, by using the percentage change in the medical care expenditure category of the Consumer Price Index (CPI) for urban consumers that is published by the Bureau of Labor Statistics. This adjustment is made using the change in the CPI from March 1984 to the fifth month of the cap year. The cap year runs from November 1 of each year until October 31 of the following year.

(b) Each hospice's cap amount is calculated by the intermediary by multiplying the adjusted cap amount determined in paragraph (a) of this section by the number of Medicare beneficiaries who elected to receive hospice care from that hospice during the cap period. For purposes of this calculation, the number of Medicare beneficiaries includes—

(1) Those Medicare beneficiaries who have not previously been included in the calculation of any hospice cap and who have filed an election to receive hospice care, in accordance with §418.24, from the hospice during the period beginning on September 28 (35 days before the beginning of the cap period) and ending on September 27 (35 days before the end of the cap period).

(2) In the case in which a beneficiary has elected to receive care from more than one hospice, each hospice includes in its number of Medicare beneficiaries only that fraction which represents the portion of a patient's total stay in all hospices that was spent in that hospice. (The hospice can obtain this information by contacting the intermediary.)

(b) Until October 1, 1986, payment to a hospice that began operation before January 1, 1975 is not limited by the amount of the hospice cap specified in §418.309.

(c) The intermediary notifies the hospice of the determination of program reimbursement at the end of the cap year in accordance with procedures similar to those described in §405.1803 of this chapter.

(d) Payments made to a hospice during a cap period that exceed the cap amount are overpayments and must be refunded.

§ 418.310 Reporting and recordkeeping requirements.

Hospices must provide reports and keep records as the Secretary determines necessary to administer the program.

§ 418.311 Administrative appeals.

A hospice that believes its payments have not been properly determined in accordance with these regulations may request a review from the intermediary or the Provider Reimbursement Review Board (PRRB) if the amount in controversy is at least $1,000 or $10,000, respectively. In such a case, the procedure in 42 CFR part 405, subpart R, will be followed to the extent that it is applicable. The PRRB, subject to review by the Secretary under §405.1874 of this chapter, shall have the authority to determine the issues raised. The methods and standards for the calculation of the payment rates by HCFA are not subject to appeal.

Subpart H—Coinsurance

§ 418.400 Individual liability for coinsurance for hospice care.

An individual who has filed an election for hospice care in accordance with §418.24 is liable for the following coinsurance payments. Hospices may charge individuals the applicable coinsurance amounts.

(a) Drugs and biologicals. An individual is liable for a coinsurance payment for each palliative drug and biological prescription furnished by the hospice while the individual is not an inpatient. The amount of coinsurance for each prescription approximates 5 percent of the cost of the drug or biological to the hospice determined in accordance with the drug copayment schedule established by the hospice, except that the amount of coinsurance for each prescription may not exceed $5. The cost of the drug or biological may not exceed what a prudent buyer would pay in similar circumstances.
The drug copayment schedule must be reviewed for reasonableness and approved by the intermediary before it is used.

(b) Respite care. (1) The amount of co-insurance for each respite care day is equal to 5 percent of the payment made by HCFA for a respite care day.

2 The amount of the individual’s co-insurance liability for respite care during a hospice coinsurance period may not exceed the inpatient hospital deductible applicable for the year in which the hospice coinsurance period began.

3 The individual hospice coinsurance period—

(i) Begins on the first day an election filed in accordance with § 418.24 is in effect for the beneficiary; and

(ii) Ends with the close of the first period of 14 consecutive days on each of which an election is not in effect for the beneficiary.

§ 418.402 Individual liability for services that are not considered hospice care.

Medicare payment to the hospice discharges an individual’s liability for payment for all services, other than the hospice coinsurance amounts described in § 418.400, that are considered covered hospice care (as described in § 418.202). The individual is liable for the Medicare deductibles and coinsurance payments and for the difference between the reasonable and actual charge on unassigned claims on other covered services that are not considered hospice care. Examples of services not considered hospice care include: Services furnished before or after a hospice election period; services of the individual’s attending physician, if the attending physician is not an employee of or working under an arrangement with the hospice; or Medicare services received for the treatment of an illness or injury not related to the individual’s terminal condition.

§ 418.405 Effect of coinsurance liability on Medicare payment.

The Medicare payment rates established by HCFA in accordance with § 418.306 are not reduced when the individual is liable for coinsurance payments. Instead, when establishing the payment rates, HCFA offsets the estimated cost of services by an estimate of average coinsurance amounts hospices collect.

[56 FR 26919, June 12, 1991]

PART 419—PROSPECTIVE PAYMENT SYSTEM FOR HOSPITAL OUT- PATIENT DEPARTMENT SERVICES

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Subpart F—Limitations on Review

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419.70 Transitional adjustment to limit decline in payment.

AUTHORITY: Secs. 1102, 1833(t), and 1871 of the Social Security Act (42 U.S.C. 1302, 1395l(t), and 1395hh).

SOURCE: 65 FR 18542, Apr. 7, 2000, unless otherwise noted.

Subpart A—General Provisions

§ 419.1 Basis and scope.

(a) Basis. This part implements section 1833(t) of the Act by establishing a prospective payment system for services furnished on or after July 1, 2000 by hospital outpatient departments to Medicare beneficiaries who are registered on hospital records as outpatients.

(b) Scope. This subpart describes the basis of payment for outpatient hospital services under the prospective payment system. Subpart B sets forth the categories of hospitals and services that are subject to the outpatient hospital prospective payment system and those categories of hospitals and services that are excluded from the outpatient hospital prospective payment system. Subpart C sets forth the basic methodology by which prospective payment rates for hospital outpatient services are determined. Subpart D describes Medicare payment amounts, beneficiary copayment amounts, and methods of payment to hospitals under the hospital outpatient prospective payment system. Subpart E describes how the hospital outpatient prospective payment system may be updated. Subpart F describes limitations on administrative and judicial review. Subpart G describes the transitional payment adjustments that are made before 2004 to limit declines in payment for outpatient services.

§ 419.2 Basis of payment.

(a) Unit of payment. Under the hospital outpatient prospective payment system, predetermined amounts are paid for designated services furnished to Medicare beneficiaries. These services are identified by codes established under the Health Care Financing Administration Common Procedure Coding System (HCPCS). The prospective payment rate for each service or procedure for which payment is allowed under the hospital outpatient prospective payment system is determined according to the methodology described in subpart C of this part. The manner in which the Medicare payment amount and the beneficiary copayment amount for each service or procedure are determined is described in subpart D of this part.

(b) Determination of hospital outpatient prospective payment rates: Included costs. The prospective payment system establishes a national payment rate, standardized for geographic wage differences, that includes operating and capital-related costs that are directly related and integral to performing a procedure or furnishing a service on an outpatient basis. In general, these costs include, but are not limited to—

(1) Use of an operating suite, procedure room, or treatment room;
(2) Use of recovery room;
(3) Use of an observation bed;
(4) Anesthesia, certain drugs, biologicals, and other pharmaceuticals; medical and surgical supplies and equipment; surgical dressings; and devices used for external reduction of fractures and dislocations;
(5) Supplies and equipment for administering and monitoring anesthesia or sedation;
(6) Intraocular lenses (IOLs);
(7) Incidental services such as venipuncture;
(8) Capital-related costs;
(9) Implantable items used in connection with diagnostic X-ray tests, diagnostic laboratory tests, and other diagnostic tests;
(10) Durable medical equipment that is implantable;
(11) Implantable prosthetic devices (other than dental) which replace all or part of an internal body organ (including colostomy bags and supplies directly related to colostomy care), including replacement of these devices; and
(12) Costs incurred to procure donor tissue other than corneal tissue.

(c) Determination of hospital outpatient prospective payment rates: Excluded costs. The following costs are excluded from the hospital outpatient prospective payment rates:
(1) Medical education costs for approved nursing and allied health education programs.
(2) Corneal tissue acquisition costs incurred by hospitals that are paid for on a reasonable cost basis.
(3) Costs for services listed in §419.22.

Subpart B—Categories of Hospitals and Services Subject to and Excluded From the Hospital Outpatient Prospective Payment System

§419.20 Hospitals subject to the hospital outpatient prospective payment system.
(a) Applicability. The hospital outpatient prospective payment system is applicable to any hospital participating in the Medicare program, except those specified in paragraph (b) of this section, for services furnished on or after July 1, 2000.
(b) Hospitals excluded from the outpatient prospective payment system. (1) Those services furnished by Maryland hospitals that are paid under a cost containment waiver in accordance with section 1814(b)(3) of the Act are excluded from the hospital outpatient prospective payment system.
(2) Critical access hospitals (CAHs) are excluded from the hospital outpatient prospective payment system.

§419.21 Hospital outpatient services subject to the outpatient prospective payment system.
Except for services described in §419.22, effective for services furnished on or after July 1, 2000, payment is made under the hospital outpatient prospective payment system for the following:
(a) Medicare Part B services furnished to hospital outpatients designated by the Secretary under this part.
(b) Services designated by the Secretary that are covered under Medicare Part B when furnished to hospital inpatients who are either not entitled to benefits under Part A or who have exhausted their Part A benefits but are entitled to benefits under Part B of the program.
(c) Partial hospitalization services furnished by community mental health centers (CMHCs).
(d) The following medical and other health services furnished by a comprehensive outpatient rehabilitation facility (CORF) when they are provided outside the patient’s plan of care; or by a home health agency (HHA) to patients who are not under an HHA plan or treatment; or by a hospice program furnishing services to patients outside the hospice benefit:
(1) Antigens.
(2) Splints and casts.
(3) Pneumococcal vaccine, influenza vaccine, and hepatitis B vaccine.

§419.22 Hospital outpatient services excluded from payment under the hospital outpatient prospective payment system.
The following services are not paid for under the hospital outpatient prospective payment system:
(a) Physician services that meet the requirements of §415.102(a) of this chapter for payment on a fee schedule basis.
(b) Nurse practitioner and clinical nurse specialist services, as defined in section 1861(s)(2)(K)(ii) of the Act.
(c) Physician assistant services, as defined in section 1861(s)(2)(K)(i) of the Act.
(d) Certified nurse-midwife services, as defined in section 1861(gg) of the Act.
(e) Services of qualified psychologists, as defined in section 1861(ii) of the Act.
(f) Services of an anesthetist as defined in §410.69 of this chapter.
(g) Clinical social worker services as defined in section 1861(hh)(2) of the Act.
(h) Outpatient therapy services described in section 1833(a)(8) of the Act.
(i) Ambulance services, as described in section 1861(v)(1)(U) of the Act, or, if applicable, the fee schedule established under section 1834(l).
(j) Except as provided in §419.22(b)(11), prosthetic devices, prosthetics, prosthetic supplies, and orthotic devices.
(k) Except as provided in §419.2(b)(10), durable medical equipment supplied by the hospital for the patient to take home.
§ 419.30 Base expenditure target for calendar year 1999.

(a) HCFA estimates the aggregate amount that would be payable for hospital outpatient services in calendar year 1999 by summing—

(1) The total amounts that would be payable from the Trust Fund for covered hospital outpatient services without regard to the outpatient prospective payment system described in this part; and

(2) The total amounts of coinsurance that would be payable by beneficiaries to hospitals for covered hospital outpatient services without regard to the outpatient prospective payment system described in this part.

(b) The estimated aggregate amount under paragraph (a) of this section is determined as though the deductible required under section 1833(b) of the Act did not apply.

§ 419.31 Ambulatory payment classification (APC) system and payment weights.

(a) APC groups. (1) HCFA classifies outpatient services and procedures that are comparable clinically and in terms of resource use into APC groups. Except as specified in paragraph (a)(2) of this section, items and services within a group are not comparable with respect to the use of resources if the highest median cost for an item or service within the group is more than 2 times greater than the lowest median cost for an item or service within the group.

(2) HCFA may make exceptions to the requirements set forth in paragraph (a)(1) in unusual cases, such as low volume items and services, but may not make such an exception in the case of a drug or biological that has been designated as an orphan drug under section 526 of the Federal Food, Drug and Cosmetic Act.

(b) APC weighting factors. (1) Using hospital outpatient claims data from calendar year 1996 and data from the most recent available hospital cost reports, HCFA determines the median costs for the services and procedures within each APC group.

(2) HCFA assigns to each APC group an appropriate weighting factor to reflect the relative median costs for the services within the APC group compared to the median costs for the services in all APC groups.

(c) Standardizing amounts. (1) HCFA determines the portion of costs determined in paragraph (b)(1) of this section that is labor-related. This is known as the “labor-related portion” of hospital outpatient costs.

(2) HCFA standardizes the median costs determined in paragraph (b)(1) of this section by adjusting for variations in hospital labor costs across geographic areas.
§ 419.32 Calculation of prospective payment rates for hospital outpatient services.

(a) Conversion factor for 1999. HCFA calculates a conversion factor in such a manner that payment for hospital outpatient services furnished in 1999 would have equaled the base expenditure target calculated in §419.30, taking into account APC group weights and estimated service frequencies and reduced by the amounts that would be payable in 1999 as outlier payments under §419.43(d) and transitional pass-through payments under §419.43(e).

(b) Conversion factor for calendar year 2000 and subsequent years. (1) Subject to paragraph (b)(2) of this section, the conversion factor for a calendar year is equal to the conversion factor calculated for the previous year adjusted as follows:

(i) For calendar years 2000, 2001, and 2002, by the hospital inpatient market basket percentage increase applicable under section 1886(b)(3)(B)(iii) of the Act reduced by one percentage point.

(ii) For calendar years 2003 and subsequent years, by the hospital inpatient market basket percentage increase applicable under section 1886(b)(3)(B)(iii) of the Act.

(2) Beginning in calendar year 2000, HCFA may substitute for the hospital inpatient market basket percentage in paragraph (b) of this section a market basket percentage increase that is determined and applied to hospital outpatient services in the same manner that the hospital inpatient market basket percentage increase is determined and applied to inpatient hospital services.

(c) Payment rates. The payment rate for services and procedures for which payment is made under the hospital outpatient prospective payment system is the product of the conversion factor calculated under paragraph (a) or paragraph (b) of this section and the relative weight determined under §419.31(b).

(d) Budget neutrality. HCFA adjusts the conversion factor as needed to ensure that updates and adjustments under §419.50(a) are budget neutral.

Subpart D—Payments to Hospitals

§ 419.40 Payment concepts.

(a) In addition to the payment rate described in §419.32, for each APC group there is a predetermined beneficiary coinsurance amount as described in §419.41(a). The Medicare program payment amount for each APC group is calculated by applying the program payment percentage as described in §419.41(b).

(b) For purposes of this section—

(1) Coinsurance percentage is calculated as the difference between the program payment percentage and 100 percent. The coinsurance percentage in any year is thus defined for each APC group as the greater of the following: the ratio of the APC group unadjusted copayment amount to the annual APC group payment rate, or 20 percent.

(2) Program payment percentage is calculated as the lower of the following: the ratio of the APC group payment rate minus the APC group unadjusted coinsurance amount, to the APC group payment rate, or 80 percent.

(3) Unadjusted coinsurance amount is calculated as 20 percent of the wage-adjusted national median of charges for services within an APC group furnished during 1996, updated to 1999 using an actuarial projection of charge increases for hospital outpatient department services during the period 1996 to 1999.

(c) Limitation of coinsurance amount to inpatient hospital deductible amount. The coinsurance amount for a procedure performed in a year cannot exceed the amount of the inpatient hospital deductible established under section 1813(b) of the Act for that year.

§ 419.41 Calculation of national beneficiary coinsurance amounts and national Medicare program payment amounts.

(a) To calculate the unadjusted coinsurance amount for each APC group, HCFA—

(1) Standardizes 1996 hospital charges for the services within each APC group to offset variations in hospital labor costs across geographic areas;

(2) Identifies the median of the wage-neutralized 1996 charges for each APC group; and
§ 419.42 Hospital election to reduce coinsurance.

(a) A hospital may elect to reduce coinsurance for any or all APC groups on a calendar year basis. A hospital may not elect to reduce copayment for some, but not all, services within the same group.

(b) A hospital must notify its fiscal intermediary of its election to reduce coinsurance no later than—

(1) June 1, 2000, for coinsurance elections for the period July 1, 2000 through December 31, 2000; or

(2) December 1 preceding the beginning of each subsequent calendar year.

(c) The hospital’s election must be properly documented. It must specifically identify the APCs to which it applies and the coinsurance amount (within the limits identified below) that the hospital has selected for each group.

(d) The election of reduced coinsurance remains in effect unchanged during the year for which the election was made.

(e) In electing reduced coinsurance, a hospital may elect a level that is less than that year’s wage-adjusted coinsurance amount for the group but not less than 20 percent of the APC payment rate as determined in § 419.32.

(f) The hospital may advertise and otherwise disseminate information concerning the reduced level of coinsurance that it has elected. All advertisements and information furnished to Medicare beneficiaries must specify that the coinsurance reductions advertised apply only to the specified services of that hospital and that coinsurance reductions are available only for hospitals that choose to reduce coinsurance for hospital outpatient services and are not allowed in any other ambulatory settings or physician offices.

§ 419.43 Adjustments to national program payment and beneficiary coinsurance amounts.

(a) General rule. HCFA determines national prospective payment rates for hospital outpatient department services and determines a wage adjustment factor to adjust the portion of the APC payment and national beneficiary coinsurance amount attributable to labor-related costs for relative differences in labor and labor-related costs across geographic regions in a budget neutral manner.

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(b) Labor-related portion of payment and copayment rates for hospital outpatient services. HCFA determines the portion of hospital outpatient costs attributable to labor and labor-related costs (known as the "labor-related portion" of hospital outpatient costs) in accordance with §419.31(c)(1).

(c) Wage index factor. HCFA uses the hospital inpatient prospective payment system wage index established in accordance with part 412 of this chapter to make the adjustment referred to in paragraph (a) of this section.

(d) Outlier adjustment—(1) General rule. Subject to paragraph (d)(4) of this section, HCFA provides for an additional payment for each hospital outpatient service (or group of services) for which a hospital's charges, adjusted to cost, exceed the following:

(i) A fixed multiple of the sum of—
(A) The applicable Medicare hospital outpatient payment amount determined under §419.32(c), as adjusted under §419.43 (other than for adjustments under this paragraph (d) or paragraph (e) of this section); and
(B) Any transitional pass-through payment under paragraph (e) of this section.

(ii) At the option of HCFA, a fixed dollar amount.

(2) Amount of adjustment. The amount of the additional payment under paragraph (d)(1) of this section is determined by HCFA and approximates the marginal cost of care beyond the applicable cutoff point under paragraph (d)(1) of this section.

(3) Limit on aggregate outlier adjustments—(i) In general. The total of the additional payments made under this paragraph (d) for covered hospital outpatient department services furnished in a year (as estimated by HCFA before the beginning of the year) may not exceed the applicable percentage specified in paragraph (d)(3)(ii) of this section of the total program payments (sum of both the Medicare and beneficiary payments to the hospital) estimated to be made under this part for all hospital outpatient services furnished in that year. If this paragraph is first applied to less than a full year, the limit applies only to the portion of the year.

(ii) Applicable percentage. For purposes of paragraph (d)(3)(i) of this section, the term "applicable percentage" means a percentage specified by HCFA up to (but not to exceed)—
(A) For a year (or portion of a year) before 2004, 2.5 percent; and
(B) For 2004 and thereafter, 3.0 percent.

(4) Transitional authority. In applying paragraph (d)(1) of this section for hospital outpatient services furnished before January 1, 2002, HCFA may—

(i) Apply paragraph (d)(1) of this section to a bill for these services related to an outpatient encounter (rather than for a specific service or group of services) using hospital outpatient payment amounts and transitional pass-through payments covered under the bill; and

(ii) Use an appropriate cost-to-charge ratio for the hospital or CMHC (as determined by HCFA), rather than for specific departments within the hospital.

(e) Transitional pass-through for additional costs of innovative medical devices, drugs, and biologicals—(1) General rule. HCFA provides for an additional payment under this paragraph for any of the following that are provided as part of a hospital outpatient service (or group of services):

(i) Current orphan drugs. A drug or biological that is used for a rare disease or condition with respect to which the drug or biological has been designated as an orphan drug under section 526 of the Federal Food, Drug and Cosmetic Act if payment for the drug or biological as an outpatient hospital service under this part was being made on the first date that the system under this part is implemented.

(ii) Current cancer therapy drugs and biologicals and brachytherapy. A drug or biological that is used in cancer therapy, including, but not limited to, a chemotherapeutic agent, an antiemetic, a hematopoietic growth factor, a colony stimulating factor, a biological response modifier, a bisphosphonate, and a device of brachytherapy, if payment for the drug, biological, or device as an outpatient hospital service under this part was being made on the first date that
the system under this part is implemented.

(iii) Current radiopharmaceutical drugs and biological products. A radiopharmaceutical drug or biological product used in diagnostic, monitoring, and therapeutic nuclear medicine procedures if payment for the drug or biological as an outpatient hospital service under this part was being made on the first date that the system under this part is implemented.

(iv) New medical devices, drugs, and biologicals. A medical device, drug, or biological not described in paragraph (e)(1)(i), (e)(1)(ii), or (e)(1)(iii) of this section if—

(A) Payment for the device, drug, or biological as an outpatient hospital service under this part was not being made as of December 31, 1996; and

(B) The cost of the device, drug, or biological is not insignificant (as defined in paragraph (e)(1)(iv)(C) and (D) of this section) in relation to the hospital outpatient fee schedule amount (as calculated under §419.32(c)) payable for the service (or group of services) involved.

(C) In the case of a new device, drug, or biological for which a transitional pass-through payment is first made before January 1, 2003, the cost of the device, drug, or biological is considered not insignificant if its expected reasonable cost exceeds 10 percent of the applicable fee schedule amount for the associated service.

(D) In the case of a new device, drug, or biological for which a transitional pass-through payment is first made on or after January 1, 2003, the cost of the device, drug, or biological is considered not insignificant if it meets all of the following thresholds:

(1) Its expected reasonable cost exceeds 10 percent of the applicable fee schedule amount for the associated service.

(2) The expected reasonable cost of the new drug, biological, or device must exceed the current portion of the fee schedule amount determined to be associated with the drug, biological, or device by 25 percent.

(3) The difference between the expected reasonable cost of the item and the portion of the hospital outpatient fee schedule amount determined to be associated with the item exceeds 10 percent of the applicable hospital outpatient fee schedule amount.

(2) Limited period of payment. The payment under this paragraph (e) with respect to a medical device, drug, or biological applies during a period of at least 2 years, but not more than 3 years, that begins—

(i) On the first date this section is implemented in the case of a drug, biological, or device described in paragraphs (e)(2)(i), (e)(2)(ii), or (e)(2)(iii) of this section and in the case of a device, drug, or biological described in paragraph (e)(1)(iv) of this section and for which payment under this part is made as an outpatient hospital service before the first date; or

(ii) In the case of a device, drug, or biological described in paragraph (e)(1)(iv) of this section not described in paragraph (e)(2)(i) of this section, on the first date on which payment is made under this part for the device, drug, or biological as an outpatient hospital service.

(3) Amount of additional payment. Subject to paragraph (e)(4)(iii) of this section, the amount of the payment under this paragraph is—

(i) In the case of a drug or biological, the amount by which the amount determined under section 1842(o) of the Act for the drug or biological exceeds the portion of the otherwise applicable Medicare hospital outpatient fee schedule amount that HCFA determines is associated with the drug or biological; or

(ii) In the case of a medical device, the amount by which the hospital’s charges for the device, adjusted to cost, exceeds the portion of the otherwise applicable Medicare hospital outpatient fee schedule amount that HCFA determines is associated with the device.

(4) Criteria to define new or innovative medical devices eligible for pass-through payments. HCFA makes pass-through payment for new or innovative medical devices that meet all of the following criteria:

(i) They were not recognized for payment as a hospital outpatient service prior to 1997.

(ii) They have been approved/cleared for use by the FDA.
(iii) They are determined to be reasonable and necessary for the diagnosis or treatment of an illness or injury or to improve the functioning of a malformed body part, as required by section 1862(a)(1)(A) of the Act. Some investigational devices are refinements of existing technologies or replications of existing technologies and may be considered reasonable and necessary. If such devices have received an FDA investigational device exemption (IDE) and are classified by the FDA as Category B devices in accordance with sections §§405.203 to 405.215 of this chapter, excluding §405.209, they will be considered for coverage under the hospital outpatient prospective payment system.

(iv) They are an integral and subordinate part of the procedure performed, are used for one patient only, are single use, come in contact with human tissue, and are surgically implanted or inserted whether or not they remain with the patient when the patient is released from the hospital outpatient department.

(v) The associated cost is not insignificant, as determined under paragraph (e)(3)(v) of this section, in relation to the APC payment for the service in which the related medical device is packaged.

(vi) They are not equipment, instruments, apparatuses, implements, or such items for which depreciation and financing expenses are recovered as depreciable assets as defined in Chapter 1 of the Medicare Provider Reimbursement Manual (HCFA Pub. 15-1).

(vii) They are not materials and supplies such as sutures, customized surgical kits, or clips, other than radiological site markers, furnished incident to a service or procedure. Supplies include pharmacological imaging and stressing agents other than radiopharmaceutical (for which transitional pass-through payment is authorized under section 1833(t)(6)(A) of the Act).

(viii) They are not materials such as biologicals or synthetics that may be used to replace human skin.

(5) Limit on aggregate annual adjustment—(i) General rule. The total of the additional payments made under this paragraph for hospital outpatient services furnished in a year, as estimated by HCFA before the beginning of the year, may not exceed the applicable percentage specified in paragraph (e)(4)(iii) of this section of the total program payments estimated to be made under this section for all hospital outpatient services furnished in that year. If this paragraph is first applied to less than a full year, the limit applies only to the portion of the year.

(ii) Applicable percentage. For purposes of paragraph (e)(4)(i) of this section, the term “applicable percentage” means—

(A) For a year (or portion of a year) before 2004, 2.5 percent; and

(B) For 2004 and thereafter, a percentage specified by HCFA up to (but not to exceed) 2.0 percent.

(iii) Uniform prospective reduction if aggregate limit projected to be exceeded. If HCFA estimates before the beginning of a year that the amount of the additional payments under this paragraph (e) for the year (or portion thereof) as determined under paragraph (e)(4)(i) of this section without regard to this paragraph (e)(4)(iii) would exceed the limit established under this paragraph (e)(4)(iii), HCFA reduces pro rata the amount of each of the additional payments under this paragraph for that year (or portion thereof) in order to ensure that the aggregate additional payments under this paragraph (as so estimated) do not exceed the limit.

(f) Budget neutrality. Outlier adjustments under paragraph (d) of this section and transitional pass-through payments under paragraph (e) of this section are established in a budget-neutral manner.

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(2) One-half of the full program and the beneficiary payment amounts for all other covered procedures.

(b) Terminated procedures. When a surgical procedure is terminated prior to completion due to extenuating circumstances or circumstances that threaten the well-being of the patient, the Medicare program payment amount and the beneficiary copayment amount are based on—

(1) The full amounts if the procedure is discontinued after the induction of anesthesia or after the procedure is started; or

(2) One-half of the full program and the beneficiary coinsurance amounts if the procedure is discontinued after the patient is prepared for surgery and taken to the room where the procedure is to be performed but before anesthesia is induced.

Subpart E—Updates

§ 419.50 Annual review.

(a) General rule. Not less often than annually, HCFA reviews and updates groups, relative payment weights, and the wage and other adjustments to take into account changes in medical practice, changes in technology, the addition of new services, new cost data, and other relevant information and factors.

(b) Consultation requirement. HCFA will consult with an expert outside advisory panel composed of an appropriate selection of representatives of providers to review (and advise HCFA concerning) the clinical integrity of the groups and weights. The panel may use data collected or developed by entities and organizations (other than the Department of Health and Human Services) in conducting the review.

(c) Effective dates. HCFA conducts the first annual review under paragraph (a) of this section in 2001 for payments made in 2002.

Subpart F—Limitations on Review

§ 419.60 Limitations on administrative and judicial review.

There can be no administrative or judicial review under sections 1869 and 1878 of the Act or otherwise of the following:

(a) The development of the APC system, including—

(1) Establishment of the groups and relative payment weights;

(2) Wage adjustment factors;

(3) Other adjustments; and

(4) Methods for controlling unnecessary increases in volume.

(b) The calculation of base amounts described in section 1833(t)(3) of the Act.

(c) Periodic adjustments described in section 1833(t)(9) of the Act.

(d) The establishment of a separate conversion factor for hospitals described in section 1886(d)(1)(B)(v) of the Act.

(e) The determination of the fixed multiple, or a fixed dollar cutoff amount, the marginal cost of care, or applicable percentage under §419.43(d) or the determination of insignificance of cost, the duration of the additional payments (consistent with §419.43(e)), the portion of the Medicare hospital outpatient fee schedule amount associated with particular devices, drugs, or biologicals, and the application of any pro rata reduction under §419.43(e).

Subpart G—Transitional Corridors

§ 419.70 Transitional adjustment to limit decline in payment.

(a) Before 2002. Except as provided in paragraph (d) of this section, for covered hospital outpatient services furnished before January 1, 2002, for which the prospective payment system amount (as defined in paragraph (e) of this section) is—

(1) At least 90 percent, but less than 100 percent, of the pre-BBA amount (as defined in paragraph (f) of this section), the amount of payment under this part is increased by 80 percent of the amount of this difference;

(2) At least 80 percent, but less than 90 percent, of the pre-BBA amount, the amount of payment under this part is increased by the amount by which the product of 0.71 and the pre-BBA amount exceeds the product of 0.70 and the prospective payment system amount;

(3) At least 70 percent, but less than 80 percent, of the pre-BBA amount, the amount of payment under this part is increased by the amount by which the
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product of 0.63 and the pre-BBA amount, exceeds the product of 0.60 and the PPS amount; or
(4) Less than 70 percent of the pre-BBA amount, the amount of payment under this part shall be increased by 21 percent of the pre-BBA amount.

(b) For 2002. Except as provided in paragraph (d) of this section, for covered hospital outpatient services furnished during 2002, for which the prospective payment system amount is—
(1) At least 90 percent, but less than 100 percent, of the pre-BBA amount, the amount of payment under this part is increased by 70 percent of the amount of this difference;
(2) At least 80 percent, but less than 90 percent, of the pre-BBA amount, the amount of payment under this part is increased by the amount by which the product of 0.61 and the pre-BBA amount exceeds the product of 0.60 and the prospective payment system amount; or
(3) Less than 80 percent of the pre-BBA amount, the amount of payment under this part is increased by 13 percent of the pre-BBA amount.

(c) For 2003. Except as provided in paragraph (d) of this section, for covered hospital outpatient services furnished during 2003, for which the prospective payment system amount is—
(1) At least 90 percent, but less than 100 percent, of the pre-BBA amount, the amount of payment under this part is increased by 60 percent of the amount of this difference; or
(2) Less than 90 percent of the pre-BBA amount, the amount of payment under this part is increased by 6 percent of the pre-BBA amount.

(d) Hold harmless provisions—(1) Temporary treatment for small rural hospitals. For covered hospital outpatient services furnished in a calendar year before January 1, 2004 for which the prospective payment system amount is less than the pre-BBA amount, the amount of payment under this part is increased by the amount of that difference if the hospital—
(i) Is located in a rural area as defined in §412.63(b) of this chapter or is treated as being located in a rural area under section 1886(d)(8)(E) of the Act; and
(ii) Has 100 or fewer beds as defined in §412.105(b) of this chapter.

(2) Permanent treatment for cancer hospitals. In the case of a hospital described in §412.23(f) of this chapter for which the prospective payment system amount is less than the pre-BBA amount for covered hospital outpatient services, the amount of payment under this part is increased by the amount of this difference.

(e) Prospective payment system amount defined. In this paragraph, the term “prospective payment system amount” means, with respect to covered hospital outpatient services, the amount payable under this part for these services (determined without regard to this paragraph or any reduction in coinsurance elected under §419.42), including amounts payable as copayment under §419.41, coinsurance under section 1866(a)(2)(A)(ii) of the Act, and the deductible under section 1833(b) of the Act.

(f) Pre-BBA amount defined—(1) General rule. In this paragraph, the “pre-BBA amount” means, with respect to covered hospital outpatient services furnished by a hospital or a community mental health center (CMHC) in a year, an amount equal to the product of the reasonable cost of the provider for these services for the portions of the provider’s cost reporting period (or periods) occurring in the year and the base provider outpatient payment-to-cost ratio for the provider (as defined in paragraph (f)(2) of this section).

(2) Base payment-to-cost-ratio defined. For purposes of this paragraph, HCFA shall determine these ratios as if the amendments to sections 1833(i)(3)(B)(ii) and 1833(n)(1)(B)(i) of the Act made by section 4521 of the BBA, to require that the full amount beneficiaries paid as coinsurance under section 1862(a)(2)(A) of the Act be taken into account in determining Medicare Part B Trust Fund payment to the hospital, were in effect in 1996. The “base payment-to-cost ratio” for a hospital or CMHC means the ratio of—

(i) The provider’s payment under this part for covered outpatient services furnished during the cost reporting period ending in 1996, including any payment for these services through cost-
sharing described in paragraph (e) of this section; and
(ii) The reasonable cost of these services for this period, without applying the cost reductions under section 1861(v)(3)(S) of the Act.

(g) Interim payments. HCFA makes payments under this paragraph to hospitals and CMHCs on an interim basis, subject to retrospective adjustments based on settled cost reports.
(h) No effect on coinsurance. No payment made under this section affects the unadjusted coinsurance amount or the coinsurance amount described in §419.41.
(i) Application without regard to budget neutrality. The additional payments made under this paragraph—
(1) Are not considered an adjustment under §419.43(f); and
(2) Are not implemented in a budget neutral manner.

PART 420—PROGRAM INTEGRITY: MEDICARE

Subpart A—General Provisions

§ 420.1 Scope and purpose. This part sets forth requirements for Medicare providers, intermediaries, and carriers to disclose ownership and control information. It also deals with access to records pertaining to certain contracts entered into by Medicare providers. These rules are aimed at protecting the integrity of the Medicare program. The statutory basis for these requirements is explained in each of the other subparts.

[51 FR 34787, Sept. 30, 1986]

Subpart B—[Reserved]

Subpart C—Disclosure of Ownership and Control Information

§ 420.200 Purpose.
§ 420.201 Definitions.
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§ 420.203 Disclosure of hiring of intermediary’s former employees.
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§ 420.205 Disclosure by providers and part B suppliers of business transaction information.
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Subpart D—Access to Books, Documents, and Records of Subcontractors

§ 420.300 Basis, purpose, and scope.
§ 420.301 Definitions.
§ 420.302 Requirement for access clause in contracts.
§ 420.303 HHS criteria for requesting books, documents, and records.
§ 420.304 Procedures for obtaining access to books, documents, and records.

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Subpart E—Rewards for Information Relating to Medicare Fraud and Abuse, and Establishment of a Program to Collect Suggestions for Improving Medicare Program Efficiency and to Reward Suggesters for Monetary Savings

§ 420.400 Basis and scope.
§ 420.401 Rewards for information relating to Medicare fraud and abuse.
§ 420.410 Establishment of a program to collect suggestions for improving Medicare program efficiency and to reward suggesters for monetary savings.

Authority: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh).

Source: 44 FR 31142, May 30, 1979, unless otherwise noted.

Subpart A—General Provisions

§ 420.1 Scope and purpose.

This part sets forth requirements for Medicare providers, intermediaries, and carriers to disclose ownership and control information. It also deals with access to records pertaining to certain contracts entered into by Medicare providers. These rules are aimed at protecting the integrity of the Medicare program. The statutory basis for these requirements is explained in each of the other subparts.

[51 FR 34787, Sept. 30, 1986]

§ 420.2 Other related regulations.

(a) Appeals procedures. Part 496 of this chapter sets forth the appeals procedures available to providers whose provider agreements HCFA terminates for failure to comply with the disclosure of information requirements set forth in subpart C of this part.

(b) Exclusion, termination, or suspension. Part 1001 of this title sets forth the rules applicable to exclusion, termination, or suspension from the Medicare program because of fraud or abuse or conviction of program-related crimes.


Subpart B—[Reserved]
Subpart C—Disclosure of Ownership and Control Information

§ 420.200 Purpose.

This subpart implements sections 1124, 1124A, 1126, and 1861(v)(1)(i) of the Social Security Act. It sets forth requirements for providers, Part B suppliers, intermediaries, and carriers to disclose ownership and control information and the identities of managing employees. It also sets forth requirements for disclosure of information about a provider’s or Part B supplier’s owners, those with a controlling interest, or managing employees convicted of criminal offenses against Medicare, Medicaid, or the title V (Maternal and Child Health Services) and title XX (Social Services) programs.

§ 420.201 Definitions.

As used in this subpart unless the context indicates otherwise:

Agent means any person who has been delegated the authority to obligate or act on behalf of a provider.

Disclosing entity means:
(1) A provider of services, an independent clinical laboratory, a renal disease facility, a rural health clinic, a Federally qualified health center, or a health maintenance organization (as defined in section 1301(a) of the Public Health Service Act);
(2) A carrier or other agency or organization that is acting for one or more providers of services for purposes of part A and part B of Medicare; and
(3) A part B supplier, as defined in §400.202 of this chapter.

Other disclosing entity means any other Medicare disclosing entity and any entity that does not participate in Medicare, but is required to disclose certain ownership and control information because of participation in any of the programs established under title V, XIX, or XX of the Act. This includes:
(1) An entity (other than an individual practitioner or group of practitioners) that furnishes, or arranges for the furnishing of, health-related services for which payment may be claimed by the entity under any plan or program established under title V of the Social Security Act or under an approved State Medicaid plan;
(2) An entity (other than an individual practitioner or group of practitioners) that furnishes, or arranges for the furnishing of, health-related services for which payment may be claimed by the entity under an approved State plan and services program under title XX of the Act; or
(3) A Medicaid fiscal agent.

Group of practitioners means two or more health care practitioners who practice their profession at a common location (whether or not they share common facilities, common supporting staff, or common equipment).

Indirect ownership interest means any ownership interest in an entity that has an ownership interest in the disclosing entity. The term includes an ownership interest in any entity that has an indirect ownership interest in the disclosing entity.

Managing employee means a general manager, business manager, administrator, director, or other individual who exercises operational or managerial control over, or who directly or indirectly conducts, the day-to-day operation of the institution, organization, or agency.

Ownership interest means the possession of equity in the capital, the stock, or the profits of the disclosing entity.

Person with an ownership or control interest means a person or corporation that—
(1) Has an ownership interest totaling 5 percent or more in a disclosing entity;
(2) Has an indirect ownership interest equal to 5 percent or more in a disclosing entity;
(3) Has a combination of direct and indirect ownership interests equal to 5 percent or more in a disclosing entity;
(4) Owns an interest of 5 percent or more in any mortgage, deed of trust, note, or other obligation secured by the disclosing entity if that interest equals at least 5 percent of the value of the property or assets of the disclosing entity;
(5) Is an officer or director of a disclosing entity that is organized as a corporation; or
(6) Is a partner in a disclosing entity that is organized as a partnership.
§ 420.202 Determination of ownership or control percentages.

(a) Indirect ownership interest. The amount of indirect ownership interest is determined by multiplying the percentages of ownership in each entity. For example, if A owns 10 percent of the stock in a corporation that owns 80 percent of the disclosing entity, A's interest equates to an 8 percent indirect ownership interest in the disclosing entity and must be reported. Conversely, if B owns 80 percent of the stock of a corporation that owns 5 percent of the stock of the disclosing entity, B's interest equates to a 4 percent indirect ownership interest in the disclosing entity and need not be reported.

(b) Person with an ownership or control interest. In order to determine the percentage of ownership interest in any mortgage, deed of trust, note, or other obligation, the percentage of interest owned in obligation is multiplied by the percentage of the disclosing entity's assets used to secure the obligation. For example, if A owns 10 percent of a note secured by 60 percent of the provider's assets, A's interest in the provider's assets equates to 6 percent and must be reported. Conversely, if B owns 40 percent of a note secured by 10 percent of the provider's assets, B's interest in the provider's assets equates to 4 percent and need not be reported.

§ 420.203 Disclosure of hiring of intermediary's former employees.

A provider must notify the Secretary promptly if it, or its home office (in the case of a chain organization), employs or obtains the services of an individual who, at any time during the year preceding such employment, was employed in a managerial, accounting, auditing, or similar capacity by an agency or organization which currently serves, or at any time during the preceding year, served as a Medicare fiscal intermediary or carrier for the provider. Similar capacity means the performance of essentially the same work functions as those of a manager, accountant, or auditor even though the individual is not so designated by title.

§ 420.204 Principals convicted of a program-related crime.

(a) Information required. Prior to HCFA's acceptance of a provider agreement or issuance or reissuance of a supplier billing number, or at any time upon written request by HCFA, the provider or part B supplier must furnish HCFA with the identity of any person who:

(1) Has an ownership or control interest in the provider or part B supplier;
(2) Is an agent or managing employee of the provider or part B supplier; or
(3) Is a person identified in paragraph (a)(1) or (a)(2) of this section and has been convicted of, or was an owner of, had a controlling interest in, or was a managing employee of a corporation that has been convicted of a criminal offense, subjected to any civil monetary penalty, or excluded from the programs for any activities related to involvement in the Medicare, Medicaid, title V or title XX social services program, since the inception of those programs.

(b) Refusal to enter into or renew agreement or to issue or reissue billing numbers. HCFA may refuse to enter into or renew an agreement with a provider of services, or to issue or reissue a billing.
number to a part B supplier, if any person who has an ownership or control interest in the provider or supplier, or who is an agent or managing employee, has been convicted of a criminal offense or subjected to any civil penalty or sanction related to the involvement of that person in Medicare, Medicaid, title V or title XX social services programs. In making this decision, HCFA considers the facts and circumstances of the specific case, including the nature and severity of the crime, penalty or sanction and the extent to which it adversely affected beneficiaries and the programs involved. HCFA also considers whether it has been given reasonable assurance that the person will not commit any further criminal or civil offense against the programs.

(c) Notification of Inspector General. HCFA promptly notifies the Inspector General of the Department of the receipt of any application or request for participation, certification, re-certification, or for a billing number that identifies any person described in paragraph (a)(3) of this section and the action taken on that application or request.

§ 420.205 Disclosure by providers and part B suppliers of business transaction information.

A provider or part B supplier must submit to HCFA, within 35 days after the date of a written request, full and complete information on—

(a) The ownership of a subcontractor with which the provider or part B supplier has had, during the previous 12 months, business transactions in an aggregate amount in excess of $25,000;

(b) Any significant business transactions between the provider or part B supplier and any wholly owned supplier or between the provider or part B supplier and any subcontractor, during the 5 year period ending on the date of the request;

(c) The names of managing employees of the subcontractors;

(d) The identity of any other entities to which payment may be made by Medicare, which a person with an ownership or control interest or a managing employee in the subcontractor has or has had an ownership or control interest in the 3-year period preceding disclosure; and

(e) Any penalties, assessments, or exclusions under sections 1128, 1128A and 1128B of the Act incurred by the subcontractor, its owners, managing employees or those with a controlling interest in the subcontract.

§ 420.206 Disclosure of persons having ownership, financial, or control interest.

(a) Information that must be disclosed. A disclosing entity must submit the following information in the manner specified in paragraph (b) of this section:

(1) The name and address of each person with an ownership or control interest in the entity or in any subcontractor in which the entity has direct or indirect ownership interest totaling 5 percent or more. In the case of a part B supplier that is a joint venture, ownership of 5 percent or more of any company participating in the joint venture should be reported. Any physician who has been issued a Unique Physician Identification Number by the Medicare program must provide this number.

(2) Whether any of the persons named, in compliance with paragraph (a)(1) of this section, is related to another as spouse, parent, child, or sibling.

(3) The name of any other disclosing entity in which any person with an ownership or control interest, or who is a managing employee in the reporting disclosing entity, has, or has had in the previous three-year period, an ownership or control interest or position as managing employee, and the nature of the relationship with the other disclosing entity. If any of these other disclosing entities has been convicted of a criminal offense or received a civil monetary or other administrative sanction related to participation in Medicare, Medicaid, title V (Maternal and Child Health) or title XX (Social Services) programs, such as penalties assessments and exclusions under sections 1128, 1128A or 1128B of the Act, the disclosing entity must also provide that information.

(b) Time and manner of disclosure. (1) Any disclosing entity that is subject to
§ 420.300 Basis, purpose, and scope.

This subpart implements section 1861(v)(1)(I) of the Act, which requires, for Medicare payment under certain provider contracts, access by the Secretary, upon written request, and the Comptroller General, and their duly authorized representatives, to certain contracts for services and to books, documents, and records necessary to verify the costs of the services. The contracts affected are those between providers and their subcontractors, and between the subcontractors and organizations related to the subcontractor by control or common ownership. It also specifies the criteria by which HHS will determine whether to request access to books, documents, and records.

§ 420.301 Definitions.

For purposes of this subpart—

Books, documents, and records means all writings, recordings, transcriptions and tapes of any description necessary to verify the nature and extent of the costs of the services provided by the subcontractor.

Common ownership means that an individual or individuals possess significant ownership or equity in the subcontractor and the entity providing the services under the contract.

Contract for services means a contract through which a provider obtains the performance of an act or acts, as distinguished from supplies or equipment. It includes any contract for both goods and services to the extent the value or cost of the service component is $10,000 or more within a 12-month period.

Control means that an individual or an organization has the power, directly or indirectly, significantly to influence or direct the actions of policies of an organization.

Provider means a hospital, skilled nursing facility, home health agency, hospice or comprehensive outpatient rehabilitation facility, or a related organization (as defined in §413.17 of this chapter) of any of these providers.

Related to the subcontractor means that the subcontractor is, to a significant extent, associated or affiliated with, owns, or is owned by, or has control of or is controlled by, the organization furnishing the services, facilities, or supplies.

Subcontractor means any entity, including an individual or individuals,
that contracts with a provider to supply a service, either to the provider or directly to a beneficiary, for which Medicare reimburses the provider the cost of the service. This includes organizations related to the subcontractor that have a contract with the subcontractor for which the cost or value is $10,000 or more in a 12-month period.

§ 420.302 Requirement for access clause in contracts.

(a) Applicability. This subpart applies to contracts—

(1) Between a provider and a subcontractor and, where subject to section 1861(v)(l)(I)(ii) of the Act, between a subcontractor and an organization related to the subcontractor;

(2) Entered into or renewed after December 5, 1980; and

(3) For services the cost or value of which is $10,000 or more over a 12-month period, including contracts for both goods and services in which the service component is worth $10,000 or more over a 12-month period.

(b) Requirement. Any contract meeting the conditions of paragraph (a) of this section must include a clause that allows the Comptroller General of the United States, HHS, and their duly authorized representatives access to the subcontractor’s contract, books, documents, and records until the expiration of four years after the services are furnished under the contract or subcontract. The access must be provided for in accordance with the provisions of this subpart. The clause must also allow similar access by HHS, the Comptroller General, and their duly authorized representatives to contracts subject to section 1861(v)(l)(I)(ii) of the Act between a subcontractor and organizations related to the subcontractor and to books, documents, and records.

(c) Prohibition against Medicare reimbursement. If a contract subject to the requirements of this subpart does not contain the clause required by paragraph (b) of this section, HCFA will not reimburse the provider for the cost of the services furnished under the contract and will recoup any payments previously made for services under the contract. However, in order to avoid nonreimbursement or recoupment, providers will have until July 30, 1983, to amend those contracts entered into or renewed after December 5, 1980, and before January 31, 1983, that do not conform to the requirements of paragraph (b) of this section.

§ 420.303 HHS criteria for requesting books, documents, and records.

HHS will generally request books, documents, and records from a subcontractor only if one of the following situations exists and the question cannot satisfactorily and efficiently be resolved without access to the books, documents, and records:

(a) HHS has reason to believe that the costs claimed for services of the subcontractor are excessive or inappropriate.

(b) There is insufficient information to judge the appropriateness of the costs.

(c) There is a written accusation with suitable evidence against the provider or subcontractor of kickbacks, bribes, rebates, or other illegal activities.

(d) There is evidence of a possible nondisclosure of the existence of a related organization.

§ 420.304 Procedures for obtaining access to books, documents, and records.

(a) Contents of the request. Requests for access will be in writing and contain the following elements:

(1) Reasonable identification of the books, documents, and records to which access is being requested.

(2) Identification of the contract or subcontract in which costs are being questioned as excessive or inappropriate.

(3) The reason that the appropriateness of the costs or value of the services of the subcontractor in question cannot be adequately or efficiently determined without access to the subcontractor’s books and records.

(4) The authority in the statute and regulations for the access requested.

(5) To the extent possible, the identification of those individuals who will be visiting the subcontractor to obtain

access to the books, documents, and records.
(6) The time and date of the scheduled visit.
(7) The name of the duly authorized representative of HHS to contact if there are any questions.

(b) Subcontractor response to a request for access to books, documents, and records.
(1) The subcontractor will have 30 days from the date of a written request for access to books, documents, and records to make them available in accordance with the request.
(2) If the subcontractor believes the request is inadequate because it does not fully meet one or more of the required elements in paragraph (a) of this section, the subcontractor must advise the requesting organization of the additional information needed.
(i) The subcontractor must notify the requesting organization within 20 days of the date of the request that it was improperly completed.
(ii) The subcontractor must make the books, documents, and records available within 20 days after the date of the requesting organization’s response.
(3) If the subcontractor believes, for good cause, that the requested books, documents, and records cannot be made available as requested within the 30-day period under paragraph (b)(1) of this section, the subcontractor may request an extension of time within which to comply with the request from the requesting organization. The requesting organization may, at its discretion, grant the request for an extension, in whole or in part, for good cause shown.
(4) The subcontractor must make the books, documents, and records available during its regular business hours for inspection, audit, and reproduction.
(5) If HHS asks the subcontractor to reproduce books, documents, and records, HHS will pay the reasonable cost of reproduction. However, if the subcontractor reproduces books, documents, and records as a means of making them available, the subcontractor must bear the cost of the reproduction and no Medicare reimbursement will be made for that purpose.
(6) HHS reserves the right to examine the originals of any requested contracts, books, documents, and records, if they exist.

(c) Refusal by subcontractor to furnish access to records. If HCFA determines that the books, documents, and records are necessary for the reimbursement determination and the subcontractor refuses to make them available, HHS may initiate legal action against the subcontractor.

Subpart E—Rewards for Information Relating to Medicare Fraud and Abuse, and Establishment of a Program to Collect Suggestions for Improving Medicare Program Efficiency and to Reward Suggesters for Monetary Savings

SOURCE: 63 FR 31128, June 8, 1998, unless otherwise noted.

§ 420.400 Basis and scope.
This subpart implements sections 203(b) and (c) of Public Law 104-191, which require the establishment of programs to encourage individuals to report suspected cases of fraud and abuse and submit suggestions on methods to improve the efficiency of the Medicare program. Sections 203(b) and (c) of Public Law 104-191 also provide the authority for HCFA to reward individuals for reporting fraud and abuse and for submitting suggestions that could improve the efficiency of the Medicare program. This subpart sets forth procedures for rewarding individuals.
[64 FR 66401, Nov. 26, 1999]

§ 420.405 Rewards for information relating to Medicare fraud and abuse.
(a) General rule. HCFA pays a monetary reward for information that leads to the recovery of at least $100 of Medicare funds from individuals and entities that are engaging in, or have engaged in, acts or omissions that constitute grounds for the imposition of a sanction under section 1128, section 1128A, or section 1128B of the Act or that have otherwise engaged in sanctionable fraud and abuse against the Medicare program. The determination of whether an individual meets the criteria for an award, and the amount of the award, is at the discretion of
Health Care Financing Administration, HHS § 420.405

HCFA. HCFA pays rewards only if a reward is not otherwise provided for by law. When HCFA applies the criteria specified in paragraphs (b), (c), and (e) of this section to determine the eligibility and the amount of the reward, it notifies the recipient as specified in paragraph (d) of this section.

(b) Information eligible for reward. (1) In order for an individual to be eligible to receive a reward, the information he or she supplied must relate to the activities of a specific individual or entity and must specify the time period of the alleged activities.

(2) HCFA does not give a reward for information relating to an individual or entity that, at the time the information is provided, is already the subject of a review or investigation by HCFA or its contractors, or the OIG, the Department of Justice, the Federal Bureau of Investigation, or any other Federal, State, or local law enforcement agency.

(c) Persons eligible to receive a reward—

(1) General rule. Any person (other than one excluded under paragraph (c)(2) of this section) is eligible to receive a reward under this section if the person submits the information in the manner set forth in paragraph (f) of this section.

(2) Excluded individuals. (i) An individual who was, or is an immediate family member of, an officer or employee of HHS or its contractors, the SSA, the OIG, a State Medicaid Agency, or the Department of Justice, the Federal Bureau of Investigation, or any other Federal, State, or local law enforcement agency at the time he or she came into possession of, or divulged, information leading to a recovery of Medicare funds is not eligible to receive a reward under this section.

(ii) Any other Federal or State employee or contractor or an HHS grantee is not eligible for a reward under this section if the information submitted came to his or her knowledge in the course of his or her official duties.

(iii) An individual who illegally obtained the information he or she submitted is excluded from receiving a reward under this section.

(iv) An individual who participated in the sanctionable offense with respect to which payment would be made is excluded from receiving a reward under this section.

(d) Notification of eligibility—(1) General rule. After all Medicare funds have been recovered and HCFA has determined a participant eligible to receive a reward under the provisions of this section, it notifies the informant of his or her eligibility, by mail, at the most recent address supplied by the individual. It is the individual's responsibility to ensure that the reward program has been notified of any change in his or her address or other relevant personal information (for example, change of name, phone number).

(2) Special circumstances. (i) If the individual has relocated to an unknown address, the individual or his or her legal representative may claim the reward by contacting HCFA within 1 year from the date on which HCFA first attempted to notify the individual about a reward. HCFA does not consider the individual or his or her legal representative eligible for a reward more than 1 year after the date on which it first attempted to give notice. HCFA does not pay interest on rewards that are not immediately claimed.

(ii) If the individual has become incapacitated or has died, an executor, administrator, or other legal representative may claim the reward on behalf of the individual or the individual's estate. The claimant must submit certified copies of the letters testamentary, letters of administration, or other similar evidence to show his or her authority to claim the reward. The claim must be filed within 1 year from the date on which HCFA first gave or attempted to give notice of the reward.

(e) Amount and payment of reward. (1) In determining whether it will pay a reward and, if so, the amount of the reward, HCFA takes into account all relevant factors, including the significance of the information furnished in relation to the ultimate resolution of the case and the recovery of Medicare funds.

(2) The amount of a reward represents what HCFA considers to be adequate compensation in the particular case, not to exceed 10 percent of the overpayments recovered in the case or $1,000, whichever is less.
§ 420.410 Establishment of a program to collect suggestions for improving Medicare program efficiency and to reward suggesters for monetary savings.

(a) Definitions. As used in this section, the following definitions apply:

Payment means a monetary award given to a suggester in recognition of, and as a reward for, a suggestion adopted by HCFA that improves the efficiency of, and results in monetary savings to, the Medicare program.

Savings means the monetary value of the net benefits the Medicare program derives from implementing the suggestion.

Suggester means an individual, a group of individuals, or a legal entity such as a corporation, partnership, or professional association, not otherwise excluded under §420.410(d), who submits a suggestion under this section.

Suggestion means an original idea submitted in writing.

Suggestion program means the specific procedures and requirements established by HCFA for receiving suggestions from the suggester on methods to improve the efficiency of the Medicare program, evaluating the suggestions and, if appropriate, paying a reward to the suggester for adopted suggestions that result in improved efficiency and produce monetary savings to the Medicare program.

(b) General rule. HCFA may make payment for adopted suggestions that increase the efficiency of the Medicare program and result in monetary savings. HCFA only makes payment for suggestions in instances in which a reward is not otherwise provided by law. The determination to adopt a suggestion, to reward the suggester, and the method of calculating a reward are at the sole discretion of HCFA.

(c) Eligibility. Except as specified in paragraph (d) of this section, any individual, group of individuals or legal entity, such as a corporation, partnership or professional association, is eligible to submit a suggestion and be considered for a reward under this suggestion program if the suggestion is submitted to HCFA in the manner set forth in paragraph (e) of this section.

(d) Exclusions. Medicare contractors, their officers and employees, individuals who work for Federal agencies under a contract, employees of Federally-sponsored research and demonstration projects, Federal officers and employees, and immediate family members of these individuals, are excluded from receiving payment under the suggestion program. If, after the suggester receives a reward payment, HCFA determines that the suggester was ineligible to receive the reward, HCFA is
Health Care Financing Administration, HHS § 420.410

not liable for the reward payment and the suggester must refund all monies received.

(e) Requirements for submitting suggestions—(1) To be considered, the suggestion must be in writing, mailed to HCFA, and must include the following information:
   (i) A description of an existing problem or need;
   (ii) A suggested method for solving the problem or filling the need; and
   (iii) If known, an estimate of the savings potential that could result from implementing the suggestion.

(2) Suggestions must be mailed to: Health Care Financing Administration Suggestion Program, 7500 Security Blvd., Baltimore, Maryland 21244-1850.

(3) Any suggesters interested in receiving a reward must provide HCFA with the following information: An individual suggester must provide his or her name, a group of suggesters must provide the names of all the group members, and a legal entity must provide its name and the name of its representative. All suggesters must provide an address, telephone number, and any other identifying information that HCFA needs to contact the suggester for additional information and, where applicable, to mail the reward.

(f) Evaluation process—(1) Relevant factors. HCFA evaluates all suggestions on the basis of the following factors:
   (i) Originality of suggestion.
   (ii) An estimate of potential monetary savings to the Medicare program.
   (iii) The extent to which Medicare program efficiency would be improved if HCFA adopts the suggestion.
   (iv) Accuracy of the information reflected in the suggestion.
   (v) Feasibility of implementation.
   (vi) Nature and complexity of the suggestion.
   (vii) Any other factors that appear to be relevant.

(2) Evaluation time limit. HCFA concludes the evaluation process in a reasonable amount of time, not to exceed 2 years from the receipt date, taking into consideration the complexity of the suggestion, the number of possible implementation strategies, and HCFA’s current workload.

(g) Basis for reward payment—(1) General rule. If HCFA determines that it is appropriate to make a reward payment for a suggestion adopted in whole or in part, that results in improved efficiency and monetary savings to the Medicare program, the payment is based on—
   (i) The actual first-year net savings to the Medicare program, or
   (ii) The average annual net savings to the Medicare program expected to be realized over a period of not more than 3 years if—
      (A) An improvement is expected to yield monetary savings for more than 1 year and implementation involves substantial costs; or
      (B) Monetary savings are negligible in the first year but are expected to substantially increase in subsequent years.

(2) Reward payment amount. HCFA determines the amount of a reward payment using the following formula:
   (i) Net savings from $1,000 to $10,000—10 percent of the savings, with a minimum award amount of $100;
   (ii) Net savings of $10,001 to $100,000—$1,000 for the first $10,000 of savings, plus 3 percent of the net savings over $10,000;
   (iii) Net savings of more than $100,000—$3,700 for the first $100,000 of savings, plus 0.5 percent of savings over $100,000, with a maximum award amount of $25,000.

(h) Adoption of suggestion and issuance of reward payment—(1) Adoption. Upon completing its evaluation, HCFA decides whether to adopt a suggestion. If HCFA receives the same or an overlapping suggestion from two or more unrelated parties, HCFA will consider a reward only for the suggestion HCFA received first, if the suggestion or overlapping part of the suggestion are identical, and HCFA has adopted that part. If the suggestions are not identical, HCFA will consider rewarding the suggestion received first, if it is feasible and HCFA is able to adopt and implement the suggestion. If the first suggestion cannot be implemented, HCFA may consider rewarding the suggestion received next, even if it is similar, provided HCFA can adopt and implement the suggestion.

(2) Issuance of reward payment. After the reward payment amount is determined, as described in paragraph (g) of
this section, HCFA mails payment to the suggester (or to the legal representatives referenced in paragraph (k) of this section) only after the suggestion has been in operation for 1 year.

(i) Group suggestions. When HCFA deems that a reward payment is appropriate for a suggestion submitted by a group of individuals, HCFA pays an equal share of the reward to each of the individuals identified in the group. If an organization such as a corporation, partnership, or professional association submits a suggestion, HCFA makes a single reward payment to that organization.

(j) Change in name or address. It is the suggester’s responsibility to notify HCFA of any change of address or other relevant information. If the suggester fails to update HCFA on any change in this information, and the reward payment mailed to the suggester is returned to HCFA, the suggester must claim the reward payment by contacting HCFA within 1 year from the date HCFA first mailed the reward payment to the suggester. HCFA does not pay interest on rewards that, for any reason, are delayed or are not immediately claimed.

(k) Incapacitated or deceased suggester. If the suggester is incapacitated or has died, an executor, administrator, or other legal representative may claim the reward on behalf of the suggester or the suggester’s estate. The claimant must submit certified copies of the letters testamentary, letters of administration, or other similar evidence to HCFA showing his or her authority to claim the reward. The claim must be filed within 1 year from the date on which HCFA first attempted to pay the reward to the individual who submitted the suggestion.

(l) Maintenance of records—(1) HCFA retains records related to the administration of the suggestion program in accordance with 36 CFR part 1228 (the regulations for the National Archives and Records Administration).

(2) HCFA does not disclose information submitted under the suggestion program, except as required by law.

[64 FR 66401, Nov. 26, 1999]
Subpart A—Scope, Definitions, and General Provisions

§ 421.1 Basis and scope.
(a) This part is based on the indicated provisions of the following sections of the Act:
1124—Requirements for disclosure of certain information.
1816 and 1842—Use of organizations and agencies in making Medicare payments to providers and suppliers of services.

(b) Section 421.118 is also based on 42 U.S.C. 1395b-1(a)(I)(F), which authorizes demonstration projects involving intermediary agreements and carrier contracts.

(c) The provisions of this part apply to agreements with Part A (Hospital Insurance) intermediaries and contracts with Part B (Supplementary Medical Insurance) carriers. They also state that HCFA may perform certain functions directly or by contract. They specify criteria and standards to be used in selecting intermediaries and evaluating their performance, in assigning or reassigning a provider or providers to particular intermediaries, and in designating regional or national intermediaries for certain classes of providers. The provisions set forth the instances where there is the opportunity for a hearing for intermediaries and carriers affected by certain adverse actions. In some circumstances, the adversely affected intermediaries may request a judicial review of hearings decisions on—
   (1) Assignment or reassignment of a provider or providers; or
   (2) Designation of an intermediary or intermediaries to serve a class of providers.

§ 421.3 Definitions.

Intermediary means an entity that has a contract with HCFA to determine and make Medicare payments for Part A or Part B benefits payable on a cost basis (or under the Prospective Payment System for hospitals) and to perform other related functions. For purposes of designating regional or alternative regional intermediaries for home health agencies and of designating intermediaries for hospices under §421.117 as well as for applying the performance criteria in §421.120 and the performance standards in §421.122 and any adverse action resulting from such application, the term intermediary also means a Blue Cross Plan which has entered into a subcontract approved by HCFA with the Blue Cross and Blue Shield Association to perform intermediary functions.

§ 421.5 General provisions.
(a) Competitive bidding not required for carriers. HCFA may enter into contracts with carriers, or with intermediaries to act as carriers in certain circumstances, without regard to section 3709 of the U.S. Revised Statutes or any other provision of law that requires competitive bidding.

(b) Indemnification of intermediaries and carriers. Intermediaries and carriers act on behalf of HCFA in carrying out certain administrative responsibilities that the law imposes. Accordingly, their agreements and contracts contain clauses providing for indemnification with respect to actions taken on behalf of HCFA and HCFA is the real party of interest in any litigation involving the administration of the program.

(c) Use of intermediaries to perform carrier functions. HCFA may contract with an intermediary to perform carrier functions with respect to services for which Part B payment is made to a provider.

(d) Nonrenewal of agreement or contract. Notwithstanding any of the provisions of this part, HCFA has the authority not to renew an agreement or contract when its term expires.

(e) Intermediary availability in an area. For more effective and efficient administration of the program, HCFA retains the right to expand or diminish the geographical area in which an intermediary is available to serve providers.

(f) Provision for automatic renewal. Agreements and contracts under this part may contain automatic renewal clauses for continuation from term to term unless either party gives notice, within timeframes specified in the
agreement or contract, of its intention not to renew.

§ 421.100 Intermediary functions.
An agreement between HCFA and an intermediary specifies the functions to be performed by the intermediary, which must include, but are not necessarily limited to, the following:
(a) Coverage. (1) The intermediary ensures that it makes payments only for services that are:
(i) Furnished to Medicare beneficiaries;
(ii) Covered under Medicare; and
(iii) In accordance with PRO determinations when they are services for which the PRO has assumed review responsibility under its contract with HCFA.
(2) The intermediary takes appropriate action to reject or adjust the claim if—
(i) The intermediary or the PRO determines that the services furnished or proposed to be furnished were not reasonable, not medically necessary, or not furnished in the most appropriate setting; or
(ii) The intermediary determines that the claim does not properly reflect the kind and amount of services furnished.
(b) Fiscal management. The intermediary must receive, disburse, and account for funds in making Medicare payments.
(c) Provider audits. The intermediary must audit the records of providers of services as necessary to assure proper payments.
(d) Utilization patterns. The intermediary must assist providers to—
(1) Develop procedures relating to utilization practices;
(2) Make studies of the effectiveness of those procedures and recommend methods to improve them;
(3) Evaluate the results of utilization review activity; and
(4) Assist in the application of safeguards against unnecessary utilization of services.
(e) Resolution of cost report disputes. The intermediary must establish and maintain procedures approved by HCFA to consider and resolve any disputes that may result from provider dissatisfaction with an intermediary’s determinations concerning provider cost reports.
(f) Reconsideration of determinations. The intermediary must establish and maintain procedures approved by HCFA for the reconsideration of its determinations to deny payments to an individual or to the provider that furnished services to the individual. The PRO performs reconsideration of cases in which it made a determination subject to reconsideration.
(g) Information and reports. The intermediary must furnish to HCFA any information and reports that HCFA requests in order to carry out its responsibilities in the administration of the Medicare program.
(h) Other terms and conditions. The intermediary must comply with all applicable laws and regulations and with any other terms and conditions included in its agreement.
(i) Dual intermediary responsibilities. With respect to the responsibility for service to provider-based HHAs and provider-based hospices, where the HHA or hospice and its parent provider will be served by different intermediaries under § 421.117 of this part, the designated regional intermediary will process bills, make coverage determinations and make payments to the HHAs and hospices. The intermediary serving the parent provider will perform all fiscal functions, including audits and settlement of the Medicare cost reports and the HHA and hospice supplement workbooks.

§ 421.103 Options available to providers and HCFA.
(a) Except for hospices (which are covered under § 421.117), a provider may elect to receive payment for covered services furnished to Medicare beneficiaries—
(1) Directly from HCFA (subject to the provisions of paragraph (b) of this section); or
(2) Through an intermediary, when both HCFA and the intermediary consent.

(b) Whenever HCFA determines it appropriate, it may contract with any organization (including an intermediary with which HCFA has previously entered into an agreement under §421.105 and §421.110 or designated as a regional or alternative regional intermediary under §421.117) for the purposes of making payments to any provider that does not elect to receive payment from an intermediary.

§421.104 Nominations for intermediary.

(a) Nomination by groups or associations of providers. (1) An association of providers, except for hospices, may nominate an organization or agency to serve as intermediary for its members.

(2) The nomination is not binding on any member of the association if it notifies HCFA of its nonconcurrence with the nomination.

(3) The nomination must be made in writing, to HCFA, and must—

(i) Identify the proposed intermediary by giving the complete name and address;

(ii) Include, or furnish as an attachment, the name, address, and bed capacity (or patient care capacity in the case of home health agencies) of each member of the association;

(iii) List the members that have concurred in the nomination of the proposed intermediary; and

(iv) Be signed by an authorized representative of the association.

(b) Action by nonmembers or nonconcurring members. Providers that nonconcur in their association’s nomination, or are not members of an association, may—

(1) Form a group of 2 or more providers for the specific purpose of nominating an intermediary, in accordance with provisions of paragraph (a) of this section;

(2) Elect to receive payments from a fiscal intermediary with which HCFA already has an agreement, if HCFA and the intermediary agree to it (see §421.106); or

(3) Elect to receive payment from HCFA as provided in §421.103.

(c) HCFA is not required to enter into an agreement with a proposed intermediary solely because it has been nominated.


§421.105 Notification of action on nomination.

(a) HCFA will send, to each member of a nominating association or group, written notice of a decision to enter into or not enter into an agreement with the nominated organization or agency.

(b) Any member of a group or association having more than one nominated intermediary approved by HCFA to act on its behalf must withdraw its nomination from all but one or exercise the option provided in §421.103(a), subject to §421.103(b), to receive payment directly from HCFA.


§421.106 Change to another intermediary or to direct payment.

(a) Any provider may request a change of intermediary, or except for a hospice, that it be paid directly by HCFA, by—

(1) Giving HCFA written notice of its desire at least 120 days before the end of its current fiscal year; and

(2) Concurrently giving written notice to its intermediary.

(b) If HCFA finds the change is consistent with effective and efficient administration of the program and approves the request under paragraph (a) of this section, it will notify the provider, the outgoing intermediary, and the newly-elected intermediary (if any) that the change will be effective on the first day following the close of the fiscal year in which the request was filed.


§421.110 Requirements for approval of an agreement.

Before entering into or renewing an intermediary agreement, HCFA will—
§ 421.112 Considerations relating to the effective and efficient administration of the program.

(a) In order to accomplish the most effective and efficient administration of the Medicare program, determinations may be made by the Secretary with respect to the termination of an intermediary agreement, or by HCFA with respect to the—
1. Renewal of an intermediary agreement (§ 421.110);
2. Assignment or reassignment of providers to an intermediary (§ 421.114); or
3. Designation of a regional or national intermediary to serve a class of providers (§ 421.116).

(b) When taking the actions listed in paragraph (a), the Secretary or HCFA will consider the performance of the individual intermediary in its Medicare operations using the factors contained in the performance criteria (§ 421.120) and performance standards (§ 421.122).

(c) In addition, when taking the actions listed in paragraph (a) of this section, the Secretary or HCFA may consider factors relating to—
1. Consistency in the administration of program policy;
2. Development of intermediary expertise in difficult areas of program administration;
3. Individual capacity of available intermediaries to serve providers as it is affected by such considerations as—
   (i) Program emphasis on the number or type of providers to be served; or
   (ii) Changes in data processing technology;
4. Overdependence of the program on the capacity of an intermediary to an extent that services could be interrupted;
5. Economy in the delivery of intermediary services;
6. Timeliness in the delivery of intermediary services;
7. Duplication in the availability of intermediaries;
8. Conflict of interest between an intermediary and provider; and
9. Any additional pertinent factors.

[45 FR 42179, June 23, 1980, as amended at 59 FR 662, Jan. 6, 1994]
§ 421.114 Assignment and reassignment of providers by HCFA.

HCFA may assign or reassign any provider to any intermediary if it determines that the assignment or reassignment will result in a more effective and efficient administration of the Medicare program. Before making this determination HCFA will consider—
(a) The preferences of the provider;
(b) The availability of an intermediary as specified in §421.5(e); and
(c) Intermediary performance measured against the criteria and standards specified in §§421.120 and 421.122.


§ 421.116 Designation of national or regional intermediaries.

(a) After considering intermediary performance measured against the criteria and standards specified in §§421.120 and 421.122, HCFA may designate a particular intermediary to serve a class of providers nationwide or in any geographic area it defines. HCFA may make this designation if it determines that the designation will result in a greater degree of effectiveness and efficiency in the administration of the Medicare program than could be achieved by an assignment of providers to an intermediary preferred by the providers.

(b) No designation may be made until the affected providers and intermediaries are given an explanation and the intermediaries are advised of their right to a hearing and judicial review as specified in §421.128. This provision does not apply to experimental contracts awarded under §421.118.

(c) To designate an intermediary, HCFA may establish classes of providers on the basis of—
(1) The type of provider, for example, hospital, skilled nursing facility, home health agency; or
(2) Common characteristics.


§ 421.117 Designation of regional and alternative designated regional intermediaries for home health agencies and hospices.

(a) This section is based on section 1816(e)(4) of the Social Security Act, which requires the Secretary to designate regional intermediaries for home health agencies (HHAs) other than hospital-based HHAs but permits him or her to designate regional intermediaries for hospital-based HHAs only if the designation meets promulgated criteria concerning administrative efficiency and effectiveness; on section 1816(e)(5) of the Social Security Act, which requires the Secretary to designate intermediaries for hospices; and on section 1874 of the Act, which permits HCFA to contract with any organization for the purpose of making payments to any provider that elects to receive payment directly from HCFA.

(b) HCFA applies the following criteria to determine whether the assignment of hospital-based HHAs to designated regional intermediaries will result in the more effective and efficient administration of the Medicare program:
(1) Uniform interpretation of Medicare rules;
(2) Expertise in bill processing;
(3) Control of administrative costs;
(4) Ease of communication of program policy and issues to affected providers;
(5) Ease of data collection;
(6) Ease of HCFA’s monitoring of intermediary performance; and
(7) Other criteria as the Secretary believes to be pertinent.

(c) Except as provided in paragraphs (e), (f), and (g) of this section, an HHA must receive payment through a regional intermediary designated by HCFA.

(d) Except as provided in paragraphs (f) through (h) of this section, a hospice must receive payment through a regional intermediary designated by HCFA.

(e) An HHA chain not desiring to receive payment from designated regional intermediaries may request service by one lead intermediary with the assistance of a local designated regional intermediary. Alternatively, the chain may request to be serviced by a single intermediary. A lead, local, or a single intermediary must be an organization that is a designated regional intermediary. Any request made under this paragraph is evaluated by HCFA in
accordance with the criteria contained at §421.106 of this subpart.

(f) An HHA or hospice not wishing to receive payment from a regional intermediary designated under paragraph (c) or (d) of this section may submit a request to the HCFA Regional Office to receive payment through an alternative regional intermediary designated by HCFA.

(g) Except as provided in paragraph (h) of this section, any request that an HHA or hospice may make to change from a designated regional intermediary to an alternative designated regional intermediary, in accordance with paragraph (f) of this section, is evaluated by HCFA in accordance with the criteria set forth at §421.106(b) of this subpart and must be filed within the timeframe established at §421.106(a) of this subpart.

(h) Exception: An HHA or a hospice that, as of June 20, 1988 is receiving payment from a designated regional intermediary may, without regard to the limitations contained in §421.106 of this subpart, continue to receive payment from that intermediary. It may do so even if that intermediary is not the designated regional intermediary or the alternative designated regional intermediary for the particular State in which the HHA or hospice is located.

[53 FR 17944, May 19, 1988]

§ 421.118 Awarding of experimental contracts.

Notwithstanding the provisions of §§421.103 and 421.104, HCFA may award a fixed price or performance incentive contract under the experimental authority contained in 42 U.S.C. 1395b-1 for performance of any of the functions specified in §421.100. Action taken by HCFA under this paragraph is not subject to—

(a) The administrative and judicial review which would otherwise be available under §421.128; or

(b) Performance criteria and performance standards review as provided for in §§421.120 and 421.122.

§ 421.120 Performance criteria.

(a) Application of performance criteria. As part of the intermediary evaluations authorized by section 1816(f) of the Act, HCFA periodically assesses the performance of intermediaries in their Medicare operations using performance criteria. The criteria measure and evaluate intermediary performance of functional responsibilities such as—

(1) Correct coverage and payment determinations;

(2) Responsiveness to beneficiary concerns; and

(3) Proper management of administrative funds.

(b) Basis for criteria. HCFA will base the performance criteria on—

(1) Nationwide intermediary experience;

(2) Changes in intermediary operations due to fiscal constraints; and

(3) HCFA’s objectives in achieving better performance.

(c) Publication of criteria. The development and revision of criteria for evaluating intermediary performance is a continuing process. Therefore, before the beginning of each evaluation period, HCFA will publish the performance criteria as a notice in the FEDERAL REGISTER.

[48 FR 7178, Feb. 18, 1983]

§ 421.122 Performance standards.

(a) Development of standards. In addition to the performance criteria (§421.120), HCFA develops detailed performance standards for use in evaluating intermediary performance which may be based on historical performance, application of acceptable statistical measures of variation to nationwide intermediary experience during a base period, or changing program emphases or requirements. These standards are also developed considering intermediary experience and evaluate the specific requirements of each functional responsibility or criterion.

(b) Factors beyond intermediary’s control. To identify measurable factors that significantly affect an intermediary’s performance, but that are not within the intermediary’s control, HCFA will—
§ 421.128 Intermediary's opportunity for hearing and right to judicial review.

(a) Basis for appeal. An intermediary adversely affected by any of the following actions shall be granted an opportunity for a hearing:

(1) Assignment or reassignment of providers to another intermediary.

[59 FR 682, Jan. 6, 1994]
(2) Designation of a national or regional intermediary to serve a class of providers.

(3) Termination of the agreement.

(b) Request for hearing. The intermediary shall file the request with HCFA within 20 days from the date on the notice of intended action.

(c) Hearing procedures. The hearing officer shall be a representative of the Secretary and not otherwise a party to the initial administrative decision. The intermediary may be represented by counsel and may present evidence and examine witnesses. A complete recording of the proceedings at the hearing will be made and transcribed.

(d) Judicial review. An adverse hearing decision concerning action under paragraph (a)(1) or (a)(2) of this section is subject to judicial review in accordance with 5 U.S.C. chapter 7.

(e) As specified in §421.118, contracts awarded under the experimental authority of HCFA are not subject to the provisions of this section.

(f) Exception. An intermediary adversely affected by the designation of a regional intermediary or an alternative regional intermediary for HHAs, or an intermediary for hospices, under §421.117 of this subpart is not entitled to a hearing or judicial review concerning adverse effects caused by the designation of an intermediary.


Subpart C—Carriers

§421.200  Carrier functions.

A contract between HCFA and a carrier, other than a regional DMEPOS carrier, specifies the functions to be performed by the carrier which must include, but are not necessarily limited to, the following:

(a) Coverage. (1) The carrier ensures that payment is made only for services that are:

(i) Furnished to Medicare beneficiaries;

(ii) Covered under Medicare; and

(iii) In accordance with PRO determinations when they are services for which the PRO has assumed review responsibility under its contract with HCFA.

(2) The carrier takes appropriate action to reject or adjust the claim if—

(i) The carrier or the PRO determines that the services furnished or proposed to be furnished were not reasonable, not medically necessary, or not furnished in the most appropriate setting;

(ii) The carrier determines that the claim does not properly reflect the kind and amount of services furnished.

(b) Payment on a cost basis. If payment is on a cost basis, the carrier must assure that payments are based on reasonable costs, as determined under part 413 of this chapter.

(c) Payment on a charge basis. If payment is on a charge basis, under part 405, subpart E of this chapter, the carrier must ensure that—

(1) Charges are reasonable and not higher than the charge for a comparable service furnished under comparable circumstances to the carrier's policy holders and subscribers; and

(2) The payment is based on one of the following—

(i) An itemized bill.

(ii) An assignment under the terms of which the reasonable charge is the full charge for the service, as specified in §424.55 of this chapter.

(iii) If the beneficiary has died, the procedures set forth in §§424.62 and 424.64 of this chapter.

(d) Fiscal management. The carrier must receive, disburse, and account for funds in making payments under Medicare.

(e) Provider audits. The carrier must audit the records of providers to whom it makes Medicare Part B payments to assure that payments are made properly.

(f) Utilization patterns. (1) The carrier must have methods and procedures for identifying utilization patterns that deviate from professionally established norms and bring the deviant patterns to the attention of appropriate professional groups.

(2) The carrier must assist providers and other persons who furnish Medicare Part B services to—

(i) Develop procedures relating to utilization practices;

(ii) Make studies of the effectiveness of those procedures and devise methods to improve them;
§ 421.203 Carrier's failure to perform efficiently and effectively.

(a) Failure by a carrier to meet, or demonstrate the capacity to meet, the criteria and standards specified in

(iii) Apply safeguards against unnecessary utilization of services; and

(iv) Develop procedures for utilization review, and establish groups to perform such reviews of providers to whom it makes Medicare Part B payments.

(g) Information and reports. The carrier must furnish to HCFA any information and reports that HCFA requests in order to carry out HCFA’s responsibilities in the administration of the Medicare program. The carrier must be responsive to requests for information from the public.

(h) Maintenance and availability of records. The carrier must maintain and make available to HCFA the records necessary for verification of payments and for other related purposes.

(i) Hearings to Part B beneficiaries. (1) The carrier must provide an opportunity for a fair hearing if it denies the beneficiary’s request for payment, does not act upon the request with reasonable promptness, or pays less than the amount claimed.

(2) The hearing procedures must be in accordance with part 405, subpart H, of this chapter (Review and Hearing Under the Supplementary Medical Insurance Program).

(j) Other terms and conditions. The carrier must comply with any other terms and conditions included in its contract.

§ 421.202 Requirements and conditions.

Before entering into or renewing a carrier contract, HCFA determines that the carrier—

(a) Has the capacity to perform its contractual responsibilities effectively and efficiently;

(b) Has the financial responsibility and legal authority necessary to carry out its responsibilities; and

(c) Will be able to meet any other requirements HCFA considers pertinent, and, if designated a regional DMEPOS carrier, any special requirements for regional carriers under §421.210 of this subpart.

§ 421.201 Performance criteria and standards.

(a) Application of performance criteria and standards. As part of the carrier evaluations mandated by section 1842(b)(2) of the Act, HCFA periodically assesses the performance of carriers in their Medicare operations using performance criteria and standards.

(1) The criteria measure and evaluate carrier performance of functional responsibilities such as—

(i) Accurate and timely payment determinations;

(ii) Responsiveness to beneficiary, physician, and supplier concerns; and

(iii) Proper management of administrative funds.

(2) The standards evaluate the specific requirements of each functional responsibility or criterion.

(b) Basis for criteria and standards. HCFA bases the performance criteria and standards on—

(1) Nationwide carrier experience;

(2) Changes in carrier operations due to fiscal constraints; and

(3) HCFA’s objectives in achieving better performance.

(c) Publication of criteria and standards. Before the beginning of each evaluation period, which usually coincides with the Federal fiscal year period of October 1—September 30, HCFA publishes the performance criteria and standards as a notice in the Federal Register. HCFA may not necessarily publish the criteria and standards every year. HCFA interprets the statutory phrase “before the beginning of each evaluation period” as allowing publication of the criteria and standards after the Federal fiscal year begins, as long as the evaluation period of the carriers for the new criteria and standards begins after the publication of the notice. [59 FR 682, Jan. 6, 1994]
§ 421.205 Termination by the Secretary.

(a) Cause for termination. The Secretary may terminate a contract with a carrier at any time if he or she determines that the carrier has failed substantially to carry out any material terms of the contract or has performed its function in a manner inconsistent with the effective and efficient administration of the Medicare Part B program.

(b) Notice and opportunity for hearing. Upon notification of the Secretary's intent to terminate the contract, the carrier may request a hearing within 20 days after the date on the notice of intent to terminate.

(c) Hearing procedures. The hearing procedures will be those specified in § 421.128(c).

§ 421.210 Designations of regional carriers to process claims for durable medical equipment, prosthetics, orthotics and supplies.

(a) Basis. This section is based on sections 1834(a) and 1834(h) of the Act which authorize the Secretary to designate one or more carriers by specific regions to process claims for durable medical equipment, prosthetic devices, orthotics and other supplies (DMEPOS). This authority has been delegated to HCFA.

(b) Types of claims. Claims for the following, except for items incident to a physician's professional service as defined in §410.26, incident to a physician's service in a rural health clinic as defined in §405.2413, or bundled into payment to a provider, ambulatory surgical center, or other facility, are processed by the designated carrier for its designated region and not by other carriers—

(1) Durable medical equipment (and related supplies) as defined in section 1861(n) of the Act;

(2) Prosthetic devices (and related supplies) as described in section 1861(s)(8) of the Act, (including intraocular lenses and parenteral and enteral nutrients, supplies, and equipment, when furnished under the prosthetic device benefit);

(3) Orthotics and prosthetics (and related supplies) as described in section 1861(s)(9);

(4) Home dialysis supplies and equipment as described in section 1861(s)(2)(F);

(5) Surgical dressings and other devices as described in section 1861(s)(5);

(6) Immunosuppressive drugs as described in section 1861(s)(2)(J); and

(7) Other items or services which are designated by HCFA.

(c) Region designation. The boundaries of the four regions for processing claims described in paragraph (b) of this section coincide with the boundaries of 1 or more sectors or areas designated for the Common Working File. These four regions contain the following States and territories: Region A: Maine, New Hampshire, Vermont, Massachusetts, Connecticut, Rhode Island, New York, New Jersey, Pennsylvania, and Delaware. Region B: Maryland, the District of Columbia, Virginia, West Virginia, Ohio, Michigan, Indiana, Illinois, Wisconsin and Minnesota. Region C: North Carolina, South Carolina, Kentucky, Tennessee, Georgia, Florida, Alabama, Mississippi, Louisiana, Texas, Arkansas, Oklahoma, New Mexico, Colorado, Puerto Rico and the Virgin Islands. Region D: Alaska, Hawaii, American Samoa, Guam, the Northern Mariana Islands, California, Nevada, Arizona, Washington, Oregon, Montana, Idaho, Utah, Wyoming, North Dakota, South Dakota, Nebraska, Kansas, Iowa and Missouri.

(d) Criteria for designating regional carriers. HCFA designates regional carriers to achieve a greater degree of effectiveness and efficiency in the administration of the Medicare program as measured by—
§ 421.214 Advance payments to suppliers furnishing items or services under Part B. 

(a) Scope and applicability. This section provides for the following: 

(1) Sets forth requirements and procedures for the issuance and recovery of advance payments to suppliers of Part B services and the rights and responsibilities of suppliers under the payment and recovery process. 

(2) Does not limit HCFA's right to recover unadjusted advance payment balances. 

(3) Does not affect suppliers' appeal rights under part 405, subpart H of this chapter relating to substantive determinations on suppliers' claims. 

(4) Does not apply to claims for Part B services furnished by suppliers that have in effect provider agreements under section 1866 of the Act and part 499 of this chapter, and are paid by intermediaries. 

(b) Definition. As used in this section, advance payment means a conditional partial payment made by the carrier in...
(c) When advance payments may be made. An advance payment may be made if all of the following conditions are met:

1. The carrier is unable to process the claim timely.

2. HCFA determines that the prompt payment interest provision specified in section 1842(c) of the Act is insufficient to make a claimant whole.

3. HCFA approves, in writing to the carrier, the making of an advance payment by the carrier.

(d) When advance payments are not made. Advance payments are not made to any supplier that meets any of the following conditions:

1. Is delinquent in repaying a Medicare overpayment.

2. Has been advised of being under active medical review or program integrity investigation.

3. Has not submitted any claims.

4. Has not accepted claims’ assignments within the most recent 180-day period preceding the system malfunction.

(e) Requirements for suppliers. (1) Except as provided for in paragraph (g)(1) of this section, a supplier must request, in writing to the carrier, an advance payment for Part B services it furnished.

(2) A supplier must accept an advance payment as a conditional payment subject to adjustment, recoupment, or both, based on an eventual determination of the actual amount due on the claim and subject to the provisions of this section.

(f) Requirements for carriers. (1) A carrier must notify a supplier as soon as it is determined that payment will not be made in a timely manner, and an advance payment option is to be offered to the supplier.

(i) A carrier must calculate an advance payment for a particular claim at no more than 80 percent of the anticipated payment for that claim based upon the historical assigned claims payment data for claims paid the supplier.

(ii) “Historical data” are defined as a representative 90-day assigned claims payment trend within the most recent 180-day experience before the system malfunction.

(iii) Based on this amount and the number of claims pending for the supplier, the carrier must determine and issue advance payments.

(iv) If historical data are not available or if backlogged claims cannot be identified, the carrier must determine and issue advance payments based on some other methodology approved by HCFA.

(v) Advance payments can be made no more frequently than once every 2 weeks to a supplier.

(2) Generally, a supplier will not receive advance payments for more assigned claims than were paid, on a daily average, for the 90-day period before the system malfunction.

(3) A carrier must recover an advance payment by applying it against the amount due on the claim on which the advance was made. If the advance payment exceeds the Medicare payment amount, the carrier must apply the unadjusted balance of the advance payment against future Medicare payments due the supplier.

(4) In accordance with HCFA instructions, a carrier must maintain a financial system of data in accordance with the Statement of Federal Financial Accounting Standards for tracking each advance payment and its recoupment.

(g) Requirements for HCFA. (1) In accordance with the provisions of this section, HCFA may determine that circumstances warrant the issuance of advance payments to all affected suppliers furnishing Part B services. HCFA may waive the requirement in paragraph (e)(1) of this section as part of that determination.

(2) If adjusting Medicare payments fails to recover an advance payment, HCFA may authorize the use of any other recoupment method available (for example, lump sum repayment or an extended repayment schedule) including, upon written notice from the carrier to the supplier, converting any unpaid balances of advance payments to overpayments. Overpayments are recovered in accordance with part 401, subpart F of this chapter concerning claims collection and compromise and part 405, subpart C of this chapter concerning recovery of overpayments.
(h) Prompt payment interest. An advance payment is a “payment” under section 1842(c)(2)(C) of the Act for purposes of meeting the time limit for the payment of clean claims, to the extent of the advance payment.

(i) Notice, review, and appeal rights. (1) The decision to advance payments and the determination of the amount of any advance payment are committed to HCFA’s discretion and are not subject to review or appeal.

(2) The carrier must notify the supplier receiving an advance payment about the amounts advanced and recouped and how any Medicare payment amounts have been adjusted.

(3) The supplier may request an administrative review from the carrier if it believes the carrier’s reconciliation of the amounts advanced and recouped is incorrectly computed. If a review is requested, the carrier must provide a written explanation of the adjustments.

(4) The review and explanation described in paragraph (i)(3) of this section is separate from a supplier’s right to appeal the amount and computation of benefits paid on the claim, as provided at part 405, subpart H of this chapter. The carrier’s reconciliation of amounts advanced and recouped is not an initial determination as defined at §405.803 of this chapter, and any written explanation of a reconciliation is not subject to further administrative review.

[61 FR 49275, Sept. 19, 1996]
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§ 422.2 Definitions.

As used in this part—

ACR stands for adjusted community rate.

Additional benefits are health care services not covered by Medicare, and reductions in premiums or cost-sharing for Medicare covered services, funded from adjusted excess amounts as calculated in the ACR.

Adjusted community rate (ACR) is the equivalent of the maximum amount allowed under § 422.310.
Arrangement means a written agreement between an M+C organization and a provider or provider network, under which—

(1) The provider or provider network agrees to furnish for a specific M+C plan(s) specified services to the organization's M+C enrollees;

(2) The organization retains responsibilities for the services; and

(3) Medicare payment to the organization discharges the enrollee's obligation to pay for the services.

Balance billing generally refers to an amount billed by a provider that represents the difference between the amount the provider charges an individual for a service and the sum of the amount the individual's health insurer (for example, the original Medicare program) will pay for the service plus any cost-sharing by the individual.

Basic benefits means all Medicare-covered benefits (except hospice services) and additional benefits.

Benefits are health care services that are intended to maintain or improve the health status of enrollees, for which the M+C organization incurs a cost or liability under an M+C plan (not solely an administrative processing cost). Benefits are submitted and approved through the ACR process.

Coinsurance is a fixed percentage of the total amount paid for a health care service that can be charged to an M+C plan enrollee on a per-service basis.

Copayment is a fixed amount that can be charged to an M+C plan enrollee on a per-service basis.

Cost-sharing includes deductibles, coinsurance, and copayments.

Licensed by the State as a risk-bearing entity means the entity is licensed or otherwise authorized by the State to assume risk for offering health insurance or health benefits coverage, such that the entity is authorized to accept prepaid capitation for providing, arranging, or paying for comprehensive health services under an M+C contract.

M+C stands for Medicare+Choice.

M+C eligible individual means an individual who meets the requirements of §422.50.

M+C organization means a public or private entity organized and licensed by a State as a risk-bearing entity (with the exception of provider-sponsored organizations receiving waivers) that is certified by HCFA as meeting the M+C contract requirements.

M+C plan means health benefits coverage offered under a policy or contract by an M+C organization that includes a specific set of health benefits offered at a uniform premium and uniform level of cost-sharing to all Medicare beneficiaries residing in the service area of the M+C plan (or in individual segments of a service area, under §422.304(b)(2)).

M+C plan enrollee is an M+C eligible individual who has elected an M+C plan offered by an M+C organization.

Mandatory supplemental benefits are health services not covered by Medicare that an M+C enrollee must purchase as part of an M+C plan that are paid for in full, directly by (or on behalf of) Medicare enrollees, in the form of premiums or cost-sharing.

MSA stands for medical savings account.

M+SA trustee means a person or business with which an enrollee establishes an M+SA MSA. A trustee may be a bank, an insurance company, or any other entity that—

(1) Is approved by the Internal Revenue Service to be a trustee or custodian of an individual retirement account (IRA); and

(2) Meets the requirements of §422.262(b).

National coverage determination (NCD) means a national policy determination regarding the coverage status of a particular service that HCFA makes under section 1862(a)(1) of the Act, and publishes as a Federal Register notice or HCFA ruling. (The term does not include coverage changes mandated by statute.)

Optional supplemental benefits are health services not covered by Medicare that are purchased at the option of the M+C enrollee and paid for in full, directly by (or on behalf of) the Medicare enrollee, in the form of premiums or cost-sharing. These services may be grouped or offered individually.

Original Medicare means health insurance available under Medicare Part A and Part B through the traditional fee-for-service payment system.

Point of service (POS) is a benefit option that an M+C coordinated care plan
can offer to its Medicare enrollees as an additional, mandatory supplemental, or optional supplemental benefit. Under the POS benefit option, the M+C plan allows members the option of receiving specified services outside of the M+C plan’s provider network. In return for this flexibility, members typically have higher cost-sharing requirements for services received and, where offered as a mandatory or optional supplemental benefit, may also be charged a premium for the POS benefit option.

Provider means—
(1) Any individual who is engaged in the delivery of health care services in a State and is licensed or certified by the State to engage in that activity in the State; and
(2) Any entity that is engaged in the delivery of health care services in a State and is licensed or certified to deliver those services if such licensing or certification is required by State law or regulation.

Provider network means the providers with which an M+C organization contracts or makes arrangements to furnish covered health care services to Medicare enrollees under an M+C coordinated care or network MSA plan.

Religious and fraternal benefit (RFB) society means an organization that—
(1) Is described in section 501(c)(8) of the Internal Revenue Code of 1986 and is exempt from taxation under section 501(a) of that Act; and
(2) Is affiliated with, carries out the tenets of, and shares a religious bond with, a church or convention or association of churches or an affiliated group of churches.

RFB plan means an M+C plan that is offered by an RFB society.

Service area means a geographic area approved by HCFA within which an M+C-eligible individual may enroll in a particular M+C plan offered by an M+C organization. Each M+C plan must be available to all M+C-eligible individuals within the plan’s service area. In deciding whether to approve an M+C plan’s proposed service area, HCFA considers the following criteria:

(1) Whether the area meets the ‘‘county integrity rule’’ that a service area generally consists of a full county or counties. However, HCFA may approve a service area that includes a portion of a county if it determines that the ‘‘partial county’’ area is necessary, nondiscriminatory, and in the best interests of the beneficiaries.

(2) The extent to which the proposed service area mirrors service areas of existing commercial health care plans or M+C plans offered by the organization.

(3) For M+C coordinated care plans and network M+C MSA plans, whether the contracting provider network meets the access and availability standards set forth in §422.112. Although not all contracting providers must be located within the plan’s service area, HCFA must determine that all services covered under the plan are accessible from the service area.

(4) For non-network M+C MSA plans, HCFA may approve single county non-network M+C MSA plans even if the M+C organization’s commercial plans have multiple county service areas.


§ 422.4 Types of M+C plans.

(a) General rule. An M+C plan may be a coordinated care plan, a combination of an M+C MSA plan and a contribution into an M+C MSA established in accordance with §422.262, or an M+C private fee-for-service plan.

(1) A coordinated care plan. A coordinated care plan is a plan that includes a network of providers that are under contract or arrangement with the organization to deliver the benefit package approved by HCFA.

(i) The network is approved by HCFA to ensure that all applicable requirements are met, including access and availability, service area, and quality.

(ii) Coordinated care plans may include mechanisms to control utilization, such as referrals from a gatekeeper for an enrollee to receive services within the plan, and financial arrangements that offer incentives to providers to furnish high quality and cost-effective care.

(iii) Coordinated care plans include plans offered by health maintenance organizations (HMOs), provider-sponsored organizations (PSOs), preferred
§ 422.6 Application requirements.

(a) Scope. This section sets forth application requirements for entities that seek a contract as an M+C organization offering an M+C plan.

(b) Completion of an application. (1) In order to obtain a determination on whether it meets the requirements to become an M+C organization and is qualified to provide a particular type of M+C plan, an entity, or an individual authorized to act for the entity (the applicant) must complete a certified application, in the form and manner required by HCFA, including the following:

(i) Documentation of appropriate State licensure or State certification that the entity is able to offer health insurance or health benefits coverage that meets State-specified standards applicable to M+C plans, and is authorized by the State to accept prepaid capitation for providing, arranging, or paying for the comprehensive health care services to be offered under the M+C contract; or

(ii) Federal waiver as described in subpart H of this part.

(i) Pays providers of services at a rate determined by the plan on a fee-for-service basis without placing the provider at financial risk;

(ii) Does not vary the rates for a provider based on the utilization of that provider’s services; and

(iii) Does not restrict enrollees’ choices among providers that are lawfully authorized to provide services and agree to accept the plan’s terms and conditions of payment.

(b) Multiple plans. Under its contract, an M+C organization may offer multiple plans, regardless of type, provided that the M+C organization is licensed or approved under State law to provide those types of plans (or, in the case of a PSO plan, has received from HCFA a waiver of the State licensing requirement). If an M+C organization has received a waiver for the licensing requirement to offer a PSO plan, that waiver does not apply to the licensing requirement for any other type of M+C plan.

(2) The authorized individual must describe thoroughly how the entity and M+C plan meet, or will meet, the requirements described in this part.

(c) Responsibility for making determinations. HCFA is responsible for determining whether an entity qualifies as an M+C organization and whether proposed M+C plans meet the requirements of this part.

(d) Resubmittal of application. An application that has been denied by HCFA may not be resubmitted for 4 months after the date of the notice from HCFA denying the application.

(e) Disclosure of application information under the Freedom of Information Act. An applicant submitting material that he or she believes is protected from disclosure under 5 U.S.C. 552, the Freedom of Information Act, or because of exceptions provided in 45 CFR part 5 (the Department's regulations providing exceptions to disclosure), should label the material "privileged" and include an explanation of the applicability of an exception described in 45 CFR part 5.

§422.8 Evaluation and determination procedures.

(a) Basis for evaluation and determination. (1) HCFA evaluates an application for an M+C contract on the basis of information contained in the application itself and any additional information that HCFA obtains through on-site visits, public hearings, and any other appropriate procedures.

(2) If the application is incomplete, HCFA notifies the contract applicant and allows 60 days from the date of the notice for the contract applicant to furnish the missing information.

(3) After evaluating all relevant information, HCFA determines whether the contract applicant's application meets the applicable requirements of §422.6.

(b) Use of information from a prior contracting period. If an M+C organization, HMO, competitive medical plan, or health care prepayment plan has failed to comply with the terms of a previous year's contract with HCFA under title XVIII of the Act, or has failed to complete a corrective action plan during the term of the contract, HCFA may deny an application from a contract applicant based on the contract applicant's failure to comply with that prior contract with HCFA even if the contract applicant meets all of the current requirements.

(c) Notice of determination. HCFA notifies each applicant that applies for an M+C contract under this part of its determination and the basis for the determination. The determination may be approval, intent to deny, or denial.

(d) Approval of application. If HCFA approves the application, it gives written notice to the contract applicant, indicating that it meets the requirements for an M+C contract.

(e) Intent to deny. (1) If HCFA finds that the contract applicant does not appear to meet the requirements for an M+C organization and appears to be able to meet those requirements within 60 days, HCFA gives the contract applicant notice of intent to deny the application for an M+C contract and a summary of the basis for this preliminary finding.

(2) Within 60 days from the date of the notice, the contract applicant may respond in writing to the issues or other matters that were the basis for HCFA's preliminary finding and may revise its application to remedy any defects HCFA identified.

(f) Denial of application. If HCFA denies the application, it gives written notice to the contract applicant indicating—

(1) That the contract applicant does not meet the contract requirements under part C of title XVIII of the Act;

(2) The reasons why the contract applicant does not meet the contract requirements; and

(3) The contract applicant's right to request reconsideration in accordance with the procedures specified in subpart N of this part.

(g) Oversight of continuing compliance.

(1) HCFA oversees an M+C organization's continued compliance with the requirements for an M+C organization.

(2) If an M+C organization no longer meets those requirements, HCFA terminates the contract in accordance with §422.510.

[65 FR 40315, June 29, 2000]
§ 422.10 Cost-sharing in enrollment-related costs (M+C user fee).

(a) Basis and scope. This section implements that portion of section 1857 of the Act that pertains to cost-sharing in enrollment-related costs. It sets forth the procedures that HCFA follows to determine the aggregate annual ‘user fee’ to be contributed by M+C organizations and to assess the required user fees for M+C plans offered by M+C organizations.

(b) Purpose of assessment. Section 1857(e)(2) of the Act authorizes HCFA to charge and collect from each M+C plan offered by an M+C organization its pro rate share of fees for administering section 1851 of the Act, relating to dissemination of enrollment information; and section 4360 of the Omnibus Budget Reconciliation Act of 1990, relating to the health insurance counseling and assistance program.

(c) Applicability. The fee assessment also applies to those demonstrations for which enrollment is effected or coordinated under section 1851 of the Act.

(d) Collection of fees.

(1) Timing of collection. HCFA collects the fees over 9 consecutive months beginning with January of each fiscal year.

(2) Amount to be collected. The aggregate amount of fees for a fiscal year is the lesser of—

(i) The estimated costs to be incurred by HCFA in that fiscal year to carry out the activities described in paragraph (b) of this section; or

(ii) For fiscal year 2000, $100 million and for fiscal year 2001 and each succeeding year, the M+C portion (as defined in paragraph (e) of this section) of $100 million.

(e) M+C portion. In this section, the term ‘M+C portion’ means, for a fiscal year, the ratio, as estimated by the Secretary of the average number of individuals enrolled in M+C plans during the fiscal year to the average number of individuals entitled to benefits under part A, and enrolled under part B, during the fiscal year.

(f) Assessment methodology. (1) The amount of the M+C portion of the user fee each M+C organization must pay is assessed as a percentage of the total Medicare payments to each organization. HCFA determines this percentage rate using the following formula:

\[ \frac{A \times B}{C} \]

where—

A is the total estimated January payments to all organizations subject to the assessment;

B is the 9-month (January through September) assessment period; and

C is the total fiscal year M+C user fee assessment amount determined in accordance with paragraph (d)(2) of this section.

(2) HCFA determines each organization’s pro rata share of the annual fee on the basis of the organization’s calculated monthly payment amount during the 9 consecutive months beginning with January. HCFA calculates each organization’s monthly pro rata share by multiplying the established percentage rate by the total monthly calculated Medicare payment amount to the organization as recorded in HCFA’s payment system on the first day of the month.

(3) HCFA deducts the organization’s fee from the amount of Federal funds otherwise payable to the organization for that month under the M+C program.

(4) If assessments reach the amount authorized for the year before the end of September, HCFA discontinues assessment.

(5) If there are delays in determining the amount of the annual aggregate fees specified in paragraph (d)(2) of this section, or the fee percentage rate specified in paragraph (f)(2), HCFA may adjust the assessment time period and the fee percentage amount.

[65 FR 40315, June 29, 2000]

Subpart B—Eligibility, Election, and Enrollment

SOURCE: 63 FR 35071, June 26, 1998, unless otherwise noted.

§ 422.50 Eligibility to elect an M+C plan.

(a) An individual is eligible to elect an M+C plan if he or she—

(1) Is entitled to Medicare under Part A and enrolled in Part B (except that an individual entitled only to Part B and who was enrolled in an HMO or CMP with a risk contract under part 417 of this chapter on December 31, 1998 may continue to be enrolled in the M+C organization as an M+C plan enrollee);
§ 422.54 Continuation of enrollment.

(a) Definition. Continuation area means an additional area (outside the service area) within which the M+C organization furnishes or arranges for furnishing services to its continuation-of-enrollment enrollees. Enrollees must reside in a continuation area on a permanent basis. A continuation area does not expand the service area of any plan.

(b) Basic rule. An M+C organization may offer a continuation of enrollment option to enrollees when they no longer reside in the service area of a plan and permanently move into the geographic area designated by the M+C organization as a continuation of enrollment area. The intent to no longer reside in an area and permanently live in another area is verified through documentation that establishes residency, such as, driver’s license, voter registration.

(c) General requirements. (1) An M+C organization that wishes to offer a continuation of enrollment option must meet the following requirements:

(i) Obtain HCFA’s approval of the continuation area, the marketing materials that describe the option, and the M+C organization’s assurances of access to services.

(ii) Describe the option(s) in the member materials it offers and make the option available to all enrollees residing in the service area.

(2) An enrollee who moves out of the service area into the geographic area designated as the continuation area has the choice of continuing enrollment or disenrolling from the plan. The enrollee must make the choice of continuing enrollment in a manner specified by HCFA. If no choice is made, the enrollee must be disenrolled from the plan.

(d) Specific requirements—

(1) Continuation of enrollment benefits. The M+C organization must, at a minimum, provide or arrange for the Medicare-covered benefits as described in §422.101(a).

(2) Reasonable access. The M+C organization must ensure reasonable access in the continuation area—

(i) Through contracts with providers, or through direct payment of claims that satisfy the requirements in §422.100(b)(2), to other providers who meet the requirement in subpart E of this part; and

(2) Has not been medically determined to have end-stage renal disease, except that an individual who develops end-stage renal disease while enrolled in an M+C plan or in a health plan offered by the M+C organization is eligible to elect an M+C plan offered by that organization;

(3) Meets either of the following residency requirements:

(i) Resides in the service area of the M+C plan.

(ii) Resides outside of the service area of the M+C plan and is enrolled in a health plan offered by the M+C organization during the month immediately preceding the month in which the individual is entitled to both Medicare Part A and Part B, provided that an M+C organization chooses to offer this option and that HCFA determines that all applicable M+C access requirements of §422.112 are met for that individual through the M+C plan's established provider network. The M+C organization must furnish the same benefits to these enrollees as to enrollees who reside in the service area;

(4) Has been a member of an Employer Group Health Plan (EGHP) that includes the elected M+C plan, even if the individual lives outside of the M+C plan service area, provided that an M+C organization chooses to offer this option and that HCFA determines that all applicable M+C access requirements at §422.12 are met for that individual through the M+C plan’s established provider network. The M+C organization must furnish the same benefits to all enrollees, regardless of whether they reside in the service area;

(5) Completes and signs an election form and gives information required for enrollment; and

(6) Agrees to abide by the rules of the M+C organization after they are disclosed to him or her in connection with the election process.

An M+C eligible individual may not be enrolled in more than one M+C plan at any given time.

(ii) By ensuring that the access requirements of §422.112 are met.

(3) Reasonable cost-sharing. For services furnished in the continuation area, an enrollee’s cost-sharing liability is limited to the cost-sharing amounts required in the M+C plan’s service area (in which the enrollee no longer resides).

(4) Protection of enrollee rights. An M+C organization that offers a continuation of enrollment option must convey all enrollee rights conferred under this rule, with the understanding that—

(i) The ultimate responsibility for all appeals and grievance requirements remain with the organization that is receiving payment from HCFA; and

(ii) Organizations that require enrollees to give advance notice of intent to use the continuation of enrollment option, must stipulate the notification process in the marketing materials.

(e) Capitation payments. HCFA’s capitation payments to all M+C organizations, for all Medicare enrollees, are based on rates established on the basis of the enrollee’s permanent residence, regardless of where he or she receives services.

§ 422.57 Limited enrollment under M+C RFB plans.

An RFB society that offers an M+C RFB plan may offer that plan only to members of the church, or convention or group of churches with which the society is affiliated.

§ 422.60 Election process.

(a) Acceptance of enrollees: General rule. (1) Except for the limitations on enrollment in an M+C MSA plan provided by §422.62(d)(1) and except as specified in paragraph (a)(2) of this section, each M+C organization must accept without restriction (except for an M+C RFB plan as provided by §422.57) individuals who are eligible to elect an M+C plan that the M+C organization offers and who elect an M+C plan during initial coverage election periods under §422.62(a)(1), annual election periods under §422.62(a)(2), and under the circumstances described in §422.62(b)(1) through (b)(4).

(2) M+C organizations must accept elections during the open enrollment periods specified in §422.62(a)(1), (a)(2), and (a)(3) if their M+C plans are open to new enrollees.

(b) Capacity to accept new enrollees. (1) M+C organizations may submit information on enrollment capacity of plans under 38 U.S.C. chapter 17, may not enroll in an M+C MSA plan.

(c) Individuals eligible for Medicare cost-sharing under Medicaid State plans. An individual who is entitled to coverage of Medicare cost-sharing under a State plan under title XIX of the Act is not eligible to enroll in an M+C MSA plan.

(d) Other limitations. An individual who receives health benefits that cover all or part of the annual deductible under the M+C MSA plan may not enroll in an M+C MSA plan. Examples of this type of coverage include, but are not limited to, primary health care coverage other than Medicare, current coverage under the Medicare hospice benefit, supplemental insurance policies not specifically permitted under §422.104, and retirement health benefits.

[63 FR 35071, June 26, 1998; 63 FR 52612, Oct. 1, 1998]
they offer by July 1 of each year as provided by §422.303(a)(1).

(2) If HCFA determines that an M+C plan offered by an M+C organization has a capacity limit, and the number of M+C eligible individuals who elect to enroll in that plan exceeds the limit, the M+C organization offering the plan may limit enrollment in the plan under this part, but only if it provides priority in acceptance as follows:

(i) First, for individuals who elected the plan prior to the HCFA determination that capacity has been exceeded, elections will be processed in chronological order by date of receipt of their election forms.

(ii) Then for other individuals in a manner that does not discriminate on the basis of any factor related to health as described in §422.110.

(3) HCFA considers enrollment limit requests for an M+C plan service area, other than those submitted with the adjusted community rate proposal, or for a portion of the plan service area, only if the health and safety of beneficiaries is at risk, such as if the provider network is not available to serve the enrollees in all or a portion of the service area.

(c) Election forms. (1) The election form must comply with HCFA instructions regarding content and format and have been approved by HCFA as described in §422.80. The form must be completed and signed by the M+C eligible individual (or the individual who will soon become entitled to Medicare benefits) and include authorization for disclosure and exchange of necessary information between the U.S. Department of Health and Human Services and its designees and the M+C organization. Persons who assist beneficiaries in completing forms must sign the form and indicate their relationship to the beneficiary.

(2) The M+C organization must file and retain election forms for the period specified in HCFA instructions.

(d) When an election is considered to have been made. An election in an M+C plan is considered to have been made on the date the election form is received by the M+C organization.

(e) Handling of election forms. The M+C organization must have an effective system for receiving, controlling, and processing election forms. The system must meet the following conditions and requirements:

(1) Each election form is dated as of the day it is received.

(2) Election forms are processed in chronological order, by date of receipt.

(3) The M+C organization gives the beneficiary prompt written notice of acceptance or denial in a format specified by HCFA.

(4) In a format specified by HCFA, a notice of acceptance—

(i) Informs the beneficiary of the date on which enrollment will be effective under §422.68; and

(ii) If the M+C plan is enrolled to capacity, explains the procedures that will be followed when vacancies occur.

(5) A notice of denial explains the reasons for denial in a format specified by HCFA.

(f) Exception for employer group health plans. (1) In cases in which an M+C organization has both a Medicare contract and a contract with an employer group health plan, and in which the M+C organization arranges for the employer to process election forms for Medicare-entitled group members, who wish to enroll under the Medicare contract, the effective date of the election may be retroactive. Consistent with §422.250(b), payment adjustments based on a retroactive effective date may be made for up to a 90-day period.

(2) In order to obtain the effective date described in paragraph (f)(1) of this section, the beneficiary must certify that, at the time of enrollment in the M+C organization, he or she received the disclosure statement specified in §422.111.
§ 422.62 Election of coverage under an M+C plan.

(a) General: Coverage election periods—
(1) Initial coverage election period. The initial coverage election period is the period during which a new M+C eligible individual may make an initial election. This period begins 3 months prior to the month the individual is first entitled to both Part A and Part B and ends the last day of the month preceding the month of entitlement.

(2) Annual election period. (i) Beginning in 1999, the month of November is the annual election period for the following calendar year. Organizations offering M+C plans in January 1999 must open enrollment to Medicare beneficiaries in November 1998.

(ii) During the annual election period, an individual eligible to enroll in an M+C plan may change his or her election from an M+C plan to original Medicare or to a different M+C plan, or from original Medicare to an M+C plan.

(3) Open enrollment and disenrollment opportunities through 2001. From 1998 through 2001, the number of elections or changes that an M+C eligible individual may make is not limited (except as provided for in paragraph (d) of this section for M+C MSA plans). Subject to the M+C plan being open to enrollees as provided under § 422.60(a)(2), an individual eligible to elect an M+C plan may change his or her election from an M+C plan to original Medicare or to a different M+C plan, or from original Medicare to an M+C plan.

(4) Open enrollment and disenrollment during 2002. (i) Except as provided in paragraphs (a)(4)(ii), (a)(4)(iii), and (a)(6) of this section, an individual who is eligible to elect an M+C plan in 2002 may elect an M+C plan or change his or her election from an M+C plan to original Medicare or to a different M+C plan, or from original Medicare to an M+C plan, but only once during the first 6 months of the year.

(ii) Newly eligible M+C individual. An individual who becomes an M+C eligible individual during 2002 may elect an M+C plan or original Medicare and then change his or her election once during the period that begins the month the individual is entitled to both Part A and Part B and ends on the last day of the 6th month of such entitlement, or on December 31, whichever is earlier. The individual can change the election from an M+C plan to original Medicare or to a different M+C plan, or from original Medicare to an M+C plan during this period.

(iii) The limitation to one election or change in paragraphs (a)(4)(i) and (a)(4)(ii) of this section does not apply to elections or changes made during the annual election period specified in (a)(2) of this section or during a special enrollment period specified in paragraph (b) of this section.

(5) Open enrollment and disenrollment beginning in 2003. (i) For 2003 and subsequent years, except as provided in paragraphs (a)(5)(ii), (a)(5)(iii), and (a)(6) of this section, an individual who is eligible to elect an M+C plan may elect an M+C plan, change his or her election from an M+C plan to original Medicare or to a different M+C plan, or from original Medicare to an M+C plan, but only once during the first 3 months of the year.

(ii) Newly eligible M+C individual. An individual who becomes an M+C eligible individual during 2003 or later may elect an M+C plan or original Medicare and then change his or her election once during the period that begins the month the individual is entitled to both Part A and Part B and ends on the last day of the 3rd month of such entitlement, or on December 31, whichever is earlier. The individual can change the election from an M+C plan to original Medicare or to a different M+C plan, or from original Medicare to an M+C plan during this period.

(iii) The limitation to one election or change in paragraphs (a)(5)(i) and (a)(5)(ii) of this section does not apply to elections or changes made during the annual election period specified in paragraph (a)(2) of this section or during a special election period specified in paragraph (b) of this section.
(6) Open enrollment period for institutionalized individuals. After 2001, an individual who is eligible to elect an M+C plan and who is institutionalized, as defined by HCFA, is not limited (except as provided for in paragraph (d) of this section for M+C MSA plans) in the number of elections or changes he or she may make. Subject to the M+C plan being open to enrollees as provided under §422.60(a)(2), an M+C eligible institutionalized individual may at any time elect an M+C plan or change his or her election from an M+C plan to original Medicare, to a different M+C plan, or from original Medicare to an M+C plan.

(b) Special election periods. Effective as of January 1, 1999 for M+C MSA plans, and as of January 1, 2002, for all other types of M+C plans, an individual may at any time (that is, not limited to the annual election period) discontinue the election of an M+C plan offered by an M+C organization and change his or her election, in the form and manner specified by HCFA, from an M+C plan to original Medicare or to a different M+C plan under any of the following circumstances:

(1) HCFA or the organization has terminated the organization’s contract for the plan, discontinued the plan in the area in which the individual resides, or the organization has notified the individual of the impending termination of the plan, or the impending discontinuation of the plan in the area in which the individual resides.

(2) The individual is not eligible to remain enrolled in the plan because of a change in his or her place of residence to a location out of the service area or continuation area or other change in circumstances as determined by HCFA but not including terminations resulting from a failure to make timely payment of an M+C monthly or supplemental beneficiary premium, or from disruptive behavior.

(3) The individual demonstrates to HCFA, in accordance with guidelines issued by HCFA, that—

(A) Failure to provide the beneficiary on a timely basis medically necessary services for which benefits are available under the plan.

(B) Failure to provide medical services in accordance with applicable quality standards; or

(ii) The organization (or its agent, representative, or plan provider) materially misrepresented the plan’s provisions in marketing the plan to the individual.

(4) The individual meets such other exceptional conditions as HCFA may provide.

(c) Special election period for individual age 65. Effective January 1, 2002, an M+C eligible individual who elects an M+C plan during the initial enrollment period, as defined under section 1837(d) of the Act, that surrounds his or her 65th birthday (this period begins 3 months before and ends 3 months after the month of the individual’s 65th birthday) may discontinue the election of that plan and elect coverage under original Medicare at any time during the 12-month period that begins on the effective date of enrollment in the M+C plan.

(d) Special rules for M+C MSA plans—

(1) Enrollment. An individual may enroll in an M+C MSA plan only during an initial or annual election period described in paragraphs (a)(1) and (a)(2) of this section or during November 1998.

(2) Disenrollment. (i) Except as provided in paragraph (d)(2)(ii) of this section, an individual may disenroll from an M+C MSA plan only during—

(A) November 1998;

(B) An annual election period; or

(C) The special election period described in paragraph (b) of this section.

(ii) Exception. An individual who elects an M+C MSA plan during an annual election period and has never before elected an M+C MSA plan may revoke that election, no later than December 15 of that same year, by submitting to the organization that offers the M+C MSA plan a signed and dated request in the form and manner prescribed by HCFA or by filing the appropriate disenrollment form through
§ 422.64 Information about the M+C program.

Each M+C organization must provide, on an annual basis, and in a format and using standard terminology that may be specified by HCFA, the information necessary to enable HCFA to provide to current and potential beneficiaries the information they need to make informed decisions with respect to the available choices for Medicare coverage.

§ 422.66 Coordination of enrollment and disenrollment through M+C organizations.

(a) Enrollment. An individual who wishes to elect an M+C plan offered by an M+C organization may make or change his or her election during the election periods specified in § 422.62 by filing the appropriate election form with the organization or through other mechanisms as determined by HCFA.

(b) Disenrollment—(1) Basic rule. An individual who wishes to disenroll from an M+C plan may change his or her election during the election periods specified in § 422.62 in either of the following manners:

(i) Elect a different M+C plan by filing the appropriate election form with the M+C organization or through other mechanisms as determined by HCFA.

(ii) Submit a signed and dated request for disenrollment to the M+C organization in the form and manner prescribed by HCFA or file the appropriate disenrollment form through other mechanisms as determined by HCFA.

(2) When a disenrollment request is considered to have been made, a disenrollment request is considered to have been made on the date the disenrollment request is received by the M+C organization.

(3) Responsibilities of the M+C organization. The M+C organization must—

(i) Submit a disenrollment notice to HCFA within timeframes specified by HCFA;

(ii) Provide the enrollee with a copy of the request for disenrollment; and

(iii) In the case of a plan where lock-in applies, also provide the enrollee with a statement explaining that he or she—

(A) Remains enrolled until the effective date of disenrollment; and

(B) Until that date, neither the M+C organization nor HCFA pays for services not provided or arranged for by the M+C plan in which the enrollee is enrolled; and

(iv) File and retain disenrollment requests for the period specified in HCFA instructions.

(4) Effect of failure to submit disenrollment notice to HCFA promptly. If the M+C organization fails to submit the correct and complete notice required in paragraph (b)(3)(i) of this section, the M+C organization must reimburse HCFA for any capitation payments received after the month in which payment would have ceased if the requirement had been met timely.

(5) Retroactive disenrollment. HCFA may grant retroactive disenrollment in the following cases:

(i) There never was a legally valid enrollment.

(ii) A valid request for disenrollment was properly made but not processed or acted upon.

(c) Election by default: Initial coverage election period. An individual who fails to make an election during the initial coverage election period is deemed to have elected original Medicare.

(d) Conversion of enrollment (seamless continuation of coverage)—(1) Basic rule. An M+C plan offered by an M+C organization must accept any individual (regardless of whether the individual has end-stage renal disease) who is enrolled in a health plan offered by the M+C organization during the month immediately preceding the month in which he or she is entitled to both Part A and Part B, and who meets the eligibility requirements at § 422.50.

(2) Reserved vacancies. Subject to HCFA’s approval, an M+C organization may set aside a reasonable number of vacancies in order to accommodate enrollment of conversions. Any set aside vacancies that are not filled within a reasonable time must be made available to other M+C eligible individuals.
(3) Effective date of conversion. If an individual chooses to remain enrolled with the M+C organization as an M+C enrollee, the individual’s conversion to an M+C enrollee is effective the month in which he or she is entitled to both Part A and Part B in accordance with the requirements in paragraph (d)(5) of this section.

(4) Prohibition against disenrollment. The M+C organization may disenroll an individual who is converting under the provisions of paragraph (a) of this section only under the conditions specified in §422.74.

(5) Election form. The individual who is converting must complete and sign an election form as described in §422.60(c)(1).

(6) Submittal of information to HCFA. The M+C organization must transmit the information necessary for HCFA to add the individual to its records as specified in §422.60(e)(6).

(e) Maintenance of enrollment. An individual who has made an election under this section is considered to have continued to have made that election until either of the following, which ever occurs first:

(1) The individual changes the election under this section.

(2) The elected M+C plan is discontinued or no longer serves the area in which the individual resides, the organization does not offer, or the individual does not elect, the option of continuing enrollment, as provided under either §422.54 or §422.74(b)(3)(ii).

(f) Exception for employer group health plans. (1) In cases when an M+C organization has both a Medicare contract and a contract with an employer group health plan, and in which the M+C organization arranges for the employer to process election forms for Medicare-entitled group members who wish to disenroll from the Medicare contract, the effective date of the election may be retroactive. Consistent with §422.250(b), payment adjustments based on a retroactive effective date may be made for up to a 90-day period.

(2) Upon receipt of the election form from the employer, the M+C organization must submit a disenrollment notice to HCFA within timeframes specified by HCFA.

§422.68 Effective dates of coverage and change of coverage.

(a) Initial coverage election period. An election made during an initial coverage election period as described in §422.62(a)(1) is effective as of the first day of the month of entitlement to both Part A and Part B.

(b) Annual election periods. For an election or change of election made during an annual election period as described in §422.62(a)(2), coverage is effective as of the first day of the following calendar year.

(c) Open enrollment periods. For an election, or change in election, made during an open enrollment period as described in §422.62(a)(3) through (a)(6), coverage is effective as of the first day of the first calendar month following the month in which the election is made, except that, if the election or change in election is made after the 10th day of any calendar month, then the election shall not take effect until the first day of the second calendar month following the date on which the election is made.

(d) Special election periods. For an election or change of election made during a special election period as described in §422.62(b), the effective date of coverage shall be determined by HCFA, to the extent practicable, in a manner consistent with protecting the continuity of health benefits coverage.

(e) Special election period for individual age 65. For an election of coverage under original Medicare made during a special election period for an individual age 65 as described in §422.62(c), coverage is effective as of the first day of the first calendar month following the month in which the election is made.

§422.74 Disenrollment by the M+C organization.

(a) General rule. Except as provided in paragraphs (b) through (d) of this section, an M+C organization may not—
§ 422.74

(1) Disenroll an individual from any M+C plan it offers; or
(2) Orally or in writing, or by any action or inaction, request or encourage an individual to disenroll.

(b) Basis for disenrollment—(1) Optional disenrollment. An M+C organization may disenroll an individual from an M+C plan it offers in any of the following circumstances:
(i) Any monthly basic and supplementary beneficiary premiums are not paid on a timely basis, subject to the grace period for late payment established under paragraph (d)(1) of this section.
(ii) The individual has engaged in disruptive behaviors specified at paragraph (d)(2) of this section.
(iii) The individual provides fraudulent information on his or her election form or permits abuse of his or her enrollment card as specified in paragraph (d)(3) of this section.

(2) Required disenrollment. An M+C organization must disenroll an individual from an M+C plan it offers in any of the following circumstances:
(i) The individual no longer resides in the M+C plan’s service area as specified under paragraph (d)(4) of this section, is no longer eligible under § 422.50(a)(3)(ii), and optional continued enrollment has not been offered or elected under § 422.54.
(ii) The individual loses entitlement to Part A or Part B benefits as described in paragraph (d)(5) of this section.
(iii) Death of the individual as described in paragraph (d)(6) of this section.

(3) Plan termination or reduction of area where plan is available.

(c) Notice requirement. If the disenrollment is for any of the reasons specified in paragraphs (b)(1), (b)(2)(i), or (b)(3) of this section (that is, other than death or loss of entitlement to Part A or Part B), the M+C organization must give the individual a written notice of the disenrollment with an explanation of why the M+C organization is planning to disenroll the individual. Notices for reasons specified in paragraphs (b)(1) through (b)(2)(i) must—
(1) Be mailed to the individual before submission of the disenrollment notice to HCFA; and
(2) Include an explanation of the individual’s right to a hearing under the M+C organization’s grievance procedures.

(d) Process for disenrollment—(1) Monthly basic and supplementary premiums are not paid timely. An M+C organization may disenroll an individual from the M+C plan for failure to pay any basic and supplementary premiums under the following circumstances:
(i) The M+C organization makes a reasonable effort to collect unpaid premium amounts by sending a written notice of nonpayment to the enrollee within 20 days after the date the delinquent charges were due—
(A) Alerting the individual that the premiums are delinquent;
(B) Providing the individual with an explanation of the disenrollment procedures and any lock-in requirements of the M+C plan; and
(C) Advising that failure to pay the premiums within the 90-day grace period will result in termination of M+C coverage;
(ii) The M+C organization only disenrolls a Medicare enrollee when the organization has not received payment within 90 days after the date it has sent the notice of nonpayment to the enrollee.

(iii) The M+C organization gives the individual a written notice of disenrollment that meets the requirement set forth in paragraph (c) of this section.

(iv) If the enrollee fails to pay the premium for optional supplemental benefits (that is, a package of benefits that an enrollee is not required to accept), but pays the basic premium and any mandatory supplemental premium, the M+C organization has the option to discontinue the optional supplemental benefits and retain the individual as an M+C enrollee.

(2) Disenrollment for disruptive behavior—(i) Basis for disenrollment. An M+C organization may disenroll an individual from the M+C plan if the individual's behavior is disruptive, unruly, abusive, or uncooperative to the extent that his or her continued enrollment in the plan seriously impairs the M+C plan's ability to furnish services to either the particular individual or other individuals enrolled in the plan.

(ii) Effort to resolve the problem. The M+C organization must make a serious effort to resolve the problems presented by the individual, including the use (or attempted use) of the M+C organization's grievance procedures. The beneficiary has a right to submit any information or explanation that he or she may wish to submit to the M+C organization.

(iii) Consideration of extenuating circumstances. The M+C organization must establish that the individual's behavior is not related to the use of medical services or to diminished mental capacity.

(iv) Documentation. The M+C organization must document the enrollee's behavior, its own efforts to resolve any problems, and any extenuating circumstances, as described in paragraphs (d)(2)(i) through (d)(2)(iii) of this section.

(v) HCFA review of the M+C organization's proposed disenrollment. (A) HCFA decides after reviewing the documentation submitted by the M+C organization and any information submitted by the beneficiary (which the M+C organization must forward to HCFA) whether the M+C organization has met the disenrollment requirements.

(B) HCFA makes the decision within 20 working days after receipt of the documentation and notifies the M+C organization within 5 working days after making its decision.

(i) Effective date of disenrollment. If HCFA permits an M+C organization to disenroll an individual for disruptive behavior, the termination is effective the first day of the calendar month after the month in which the M+C organization gives the individual written notice of the disenrollment that meets the requirements set forth in paragraph (c) of this section.

(3) Individual commits fraud or permits abuse of enrollment card.—(i) Basis for disenrollment. An M+C organization may disenroll the individual from an M+C plan if the individual—

(A) Knowingly provides, on the election form, fraudulent information that materially affects the individual's eligibility to enroll in the M+C plan; or

(B) Intentionally permits others to use his or her enrollment card to obtain services under the M+C plan.

(ii) Notice of disenrollment. The M+C organization must give the individual a written notice of the disenrollment that meets the requirements set forth in paragraph (c) of this section.

(iii) Report to HCFA. The M+C organization must report to HCFA any disenrollment based on fraud or abuse by the individual.

(4) Individual no longer resides in the M+C plan's service area. (i) Basis for disenrollment. Unless continuation of enrollment is elected under §422.54, the M+C organization must disenroll an individual if the M+C organization establishes, on the basis of a written statement from the individual or other evidence acceptable to HCFA, that the individual has permanently moved out of the plan's service area. If the individual has not moved from the M+C plan's service area, but has left the plan's service area for more than 6 months, the M+C organization must disenroll the individual.

(ii) Special rule. The M+C organization must disenroll an individual who
is enrolled in the M+C plan, under the eligibility requirements at §422.50(a)(3)(ii) or (a)(4), if the organization establishes, on the basis of a written statement from the individual or other evidence acceptable to HCFA, that the individual has permanently moved from the residence in which she or he resided at the time of enrollment in the M+C plan, to an area outside the M+C plan service area (unless continuation of enrollment is elected under §422.54). If the individual has not permanently moved from the residence in which she or he resided at the time of enrollment in the M+C plan, but has left the residence for over 6 months, the M+C organization must disenroll the individual.

(iii) Notice of disenrollment. The M+C organization must give the individual a written notice of the disenrollment that meets the requirements set forth in paragraph (c) of this section.

(5) Loss of entitlement to Part A or Part B benefits. If an individual is no longer entitled to Part A or Part B benefits, HCFA notifies the M+C organization that the disenrollment is effective the first day of the calendar month following the last month of entitlement to Part A or Part B benefits.

(6) Death of the individual. If the individual dies, disenrollment is effective the first day of the calendar month following the month of death.

(7) Plan termination or area reduction.

(i) When an M+C organization has its contract for an M+C plan terminated, terminates an M+C plan, or discontinues offering the plan in any portion of the area where the plan had previously been available, the M+C organization must give each affected M+C plan enrollee a written notice of the effective date of the plan termination or area reduction and a description of alternatives for obtaining benefits under the M+C program.

(ii) The notice must be sent before the effective date of the plan termination or area reduction, and in the timeframes specified in §422.506(a)(2).

(e) Consequences of disenrollment—(1) Disenrollment for non-payment of premiums, disruptive behavior, fraud or abuse, loss of Part A or Part B. An individual who is disenrolled under paragraph (b)(1)(i), (b)(1)(ii), (b)(1)(iii), or paragraph (b)(2)(ii) of this section is deemed to have elected original Medicare.

(2) Disenrollment based on plan termination, area reduction, or individual moves out of area. (i) An individual who is disenrolled under paragraph (b)(2)(i) or (b)(3) of this section has a special election period in which to make a new election as provided in §422.62(b)(1) and (b)(2).

(ii) An individual who fails to make an election during the special election period is deemed to have elected original Medicare.


§422.80 Approval of marketing materials and election forms.

(a) HCFA review of marketing materials. An M+C organization may not distribute any marketing materials (as defined in paragraph (b)), or election forms, or make such materials or forms available to individuals eligible to elect an M+C plan, unless—

(1) At least 45 days before the date of distribution the M+C organization has submitted the material or form to HCFA for review under the guidelines in paragraph (c); and

(2) HCFA has not disapproved the distribution of the material or form.

(b) Definition of marketing materials. Marketing materials include any informational materials targeted to Medicare beneficiaries which:

(1) Promote the M+C organization, or any M+C plan offered by the M+C organization;

(2) Inform Medicare beneficiaries that they may enroll, or remain enrolled in, an M+C plan offered by the M+C organization;

(3) Explain the benefits of enrollment in an M+C plan, or rules that apply to enrollees;

(4) Explain how Medicare services are covered under an M+C plan, including conditions that apply to such coverage;

(5) Examples of marketing materials include, but are not limited to:

(i) General audience materials such as general circulation brochures, newspapers, magazines, television, radio, billboards, yellow pages, or the internet.
(ii) Marketing representative materials such as scripts or outlines for telemarketing or other presentations.
(iii) Presentation materials such as slides and charts.
(iv) Promotional materials such as brochures or leaflets, including materials for circulation by third parties (e.g., physicians or other providers).
(v) Membership communication materials such as membership rules, subscriber agreements (evidence of coverage), member handbooks and wallet card instructions to enrollees.
(vi) Letters to members about contractual changes; changes in providers, premiums, benefits, plan procedures etc.
(vii) Membership or claims processing activities (e.g., materials on rules involving non-payment of premiums, confirmation of enrollment or disenrollment, or annual notification information).
(c) Guidelines for HCFA review. In reviewing marketing material or election forms under paragraph (a) of this section, HCFA determines that the marketing materials:
(1) Provide, in a format (and, where appropriate, print size), and using standard terminology that may be specified by HCFA, the following information to Medicare beneficiaries interested in enrolling:
   (i) Adequate written description of rules (including any limitations on the providers from whom services can be obtained), procedures, basic benefits and services, and fees and other charges.
   (ii) Adequate written description of any supplemental benefits and services.
   (iii) Adequate written explanation of the grievance and appeals process, including differences between the two, and when it is appropriate to use each.
   (iv) Any other information necessary to enable beneficiaries to make an informed decision about enrollment.
(2) Notify the general public of its enrollment period (whether time-limited or continuous) in an appropriate manner, through appropriate media, throughout its service and continuation area.
(3) Include in the written materials notice that the M+C organization is authorized by law to refuse to renew its contract with HCFA, that HCFA also may refuse to renew the contract, and that termination or non-renewal may result in termination of the beneficiary’s enrollment in the plan.
(4) Are not materially inaccurate or misleading or otherwise make material misrepresentations.
(5) For markets with a significant non-English speaking population, provide materials in the language of these individuals.
(d) Deemed approval (one-stop shopping). If HCFA has not disapproved the distribution of marketing materials or forms submitted by an M+C organization with respect to an M+C plan in an area, HCFA is deemed not to have disapproved the distribution in all other areas covered by the M+C plan and organization except with regard to any portion of the material or form that is specific to the particular area.
(e) Standards for M+C organization marketing. (1) In conducting marketing activities, M+C organizations may not:
   (i) Provide for cash or other monetary rebates as an inducement for enrollment or otherwise. This does not prohibit explanation of any legitimate benefits the beneficiary might obtain as an enrollee of the M+C plan, such as eligibility to enroll in a supplemental benefit plan that covers deductibles and coinsurance, or preventive services.
   (ii) Engage in any discriminatory activity such as, for example, attempts to recruit Medicare beneficiaries from higher income areas without making comparable efforts to enroll Medicare beneficiaries from lower income areas.
   (iii) Solicit door-to-door for Medicare beneficiaries.
   (iv) Engage in activities that could mislead or confuse Medicare beneficiaries, or misrepresent the M+C organization. The M+C organization may not claim that it is recommended or endorsed by HCFA or Medicare or that HCFA or Medicare recommends that the beneficiary enroll in the M+C plan.
   (v) Engage in activities that could mislead or confuse Medicare beneficiaries, or misrepresent the M+C organization. The M+C organization may not claim that it is recommended or endorsed by HCFA or Medicare or that HCFA or Medicare recommends that the beneficiary enroll in the M+C plan.
   (vi) Engage in activities that could mislead or confuse Medicare beneficiaries, or misrepresent the M+C organization. The M+C organization may not claim that it is recommended or endorsed by HCFA or Medicare or that HCFA or Medicare recommends that the beneficiary enroll in the M+C plan.
   (vii) Include in the written materials notice that the M+C organization is not authorized by law to refuse to renew its contract with HCFA, that HCFA also may refuse to renew the contract, and that termination or non-renewal may result in termination of the beneficiary’s enrollment in the plan.
§ 422.100 General requirements.

(a) Basic rule. Subject to the conditions and limitations set forth in this subpart, an M+C organization offering an M+C plan must provide enrollees in that plan with coverage of the basic benefits described in paragraph (c) of this section (and, to the extent applicable, the benefits described in § 422.102) by furnishing the benefits directly or through arrangements, or by paying for the benefits. HCFA reviews these benefits subject to the requirements of § 422.100(g) and the requirements in subpart G of this part.

(b) Services of noncontracting providers and suppliers. (1) An M+C organization must make timely and reasonable payment to or on behalf of the plan enrollee for the following services obtained from a provider or supplier that does not contract with the M+C organization to provide services covered by the M+C plan:

(i) Ambulance services dispatched through 911 or its local equivalent as provided in § 422.113.

(ii) Emergency and urgently needed services as provided in § 422.113.

(iii) Maintenance and post-stabilization care services as provided in § 422.113.

(iv) Renal dialysis services provided while the enrollee was temporarily outside the plan's service area.

(v) Services for which coverage has been denied by the M+C organization and found (upon appeal under subpart M of this part) to be services the enrollee was entitled to have furnished, or paid for, by the M+C organization.

(2) An M+C plan (other than an M+C MSA plan) offered by an M+C organization satisfies paragraph (a) of this section with respect to benefits for services furnished by a noncontracting provider if that M+C plan provides payment in an amount the provider would have received under original Medicare (including balance billing permitted under Medicare Part A and Part B).

(c) Types of benefits. An M+C plan includes at a minimum basic benefits, and also may include mandatory and optional supplemental benefits.

(1) Basic benefits are all Medicare-covered services, except hospice services, and additional benefits as defined
§ 422.101 Requirements relating to basic benefits.

Except as specified in §422.264 (for entitlement that begins or ends during a hospital stay) and §422.266 (with respect to hospice care), each M+C organization must meet the following requirements:

(a) Provide coverage of, by furnishing, arranging for, or making payment for, all services that are covered by Part A and Part B of Medicare (if the enrollee is entitled to benefits under both parts) or by Medicare Part B (if entitled only under Part B) and that are available to beneficiaries residing in the plan’s service area. Services may be provided outside of the service area of the plan if the services are accessible and available to enrollees.

(b) Comply with—
(1) HCFA’s national coverage determinations;
(2) General coverage guidelines included in original Medicare manuals and instructions unless superseded by operational policy letters or regulations in this part; and
(3) Written coverage decisions of local carriers and intermediaries with jurisdiction for claims in the geographic area in which services are covered under the M+C plan.

[65 FR 40319, June 29, 2000]
§ 422.102 Supplemental benefits.

(a) Mandatory supplemental benefits.

(1) Subject to HCFA’s approval, an M+C organization may require Medicare enrollees of an M+C plan other than an MSA plan to accept and pay for services in addition to Medicare-covered services described in §422.101 and additional benefits described in §422.312.

(2) If the M+C organization imposes mandatory supplemental benefits, it must impose them on all Medicare beneficiaries enrolled in the M+C plan.

(3) HCFA approves mandatory supplemental benefits if the benefits are designed in accordance with HCFA’s guidelines and requirements as stated in this part and instructions and operational policy letters.

(b) Optional supplemental benefits. Except as provided in §422.104 in the case of MSA plans, each M+C organization may offer (for election by the enrollee and without regard to health status) services that are not included in the basic benefits as described in §422.100(c) and any mandatory supplemental benefits described in paragraph (a) of this section. Optional supplemental benefits are purchased at the discretion of the enrollee and must be offered to all Medicare beneficiaries enrolled in the M+C plan.

(c) Payment for supplemental services. All supplemental benefits are paid in full, directly by (or on behalf of) the enrollee of the M+C plan.

(d) Marketing of supplemental benefits. M+C organizations may offer enrollees a group of services as one optional supplemental benefit, offer services individually, or offer a combination of groups and individual services.

[65 FR 40320, June 29, 2000]

§ 422.103 Benefits under an M+C MSA plan.

(a) General rule. An M+C organization offering an M+C MSA plan must make available to an enrollee, or provide reimbursement for, at least the services described under §422.101 after the enrollee incurs countable expenses equal to the amount of the plan’s annual deductible.

(b) Countable expenses. An M+C organization offering an M+C MSA plan must count toward the annual deductible at least all amounts that would be paid for the particular service under original Medicare, including amounts that would be paid by the enrollee as deductibles or coinsurance.

(c) Services after the deductible. For services received by the enrollee after the annual deductible is satisfied, an M+C organization offering an M+C MSA plan must pay, at a minimum, the lesser of the following amounts:

(1) 100 percent of the expense of the services.

(2) 100 percent of the amounts that would have been paid for the services under original Medicare, including amounts that would be paid by the enrollee as deductibles and coinsurance.

(d) Annual deductible. The annual deductible for an M+C MSA plan—

(1) For contract year 1999, may not exceed $6,000; and

(2) For subsequent contract years may not exceed the deductible for the preceding contract year, increased by the national per capita growth percentage determined under §422.252(b).

§ 422.104 Special rules on supplemental benefits for M+C MSA plans.

(a) An M+C organization offering an M+C MSA plan may not provide supplemental benefits that cover expenses that count towards the deductible specified in §422.103(d).

(b) In applying the limitation of paragraph (a) of this section, the following kinds of policies are not considered as covering the deductible:

(1) A policy that provides coverage (whether through insurance or otherwise) for accidents, disability, dental care, vision care, or long-term care.

(2) A policy of insurance in which substantially all of the coverage relates to liabilities incurred under workers’ compensation laws, tort liabilities, liabilities relating to use or ownership of property, and any other similar liabilities that HCFA may specify by regulation.

(3) A policy of insurance that provides coverage for a specified disease or illness or pays a fixed amount per day (or other period) of hospitalization.
§ 422.105 Special rules for point of service option.

(a) General rule. A POS benefit is an option that an M+C organization may offer in an M+C coordinated care plan or network M+C MSA plan to provide enrollees with additional choice in obtaining specified health care services. The organization may offer a POS option—

(1) Under a coordinated care plan only as an additional benefit as described in § 422.312;

(2) Under a coordinated care plan only as a mandatory supplemental benefit as described in § 422.102(a); or

(3) Under a coordinated care plan or network MSA plan as an optional supplemental benefit as described in § 422.102(b).

(b) Approval required. An M+C organization may not implement a POS benefit until it has been approved by HCFA.

(c) Ensuring availability and continuity of care. An M+C network plan that includes a POS benefit must continue to provide all benefits and ensure access as required under this subpart.

(d) Enrollee information and disclosure. The disclosure requirements specified in § 422.111 apply in addition to the following requirements:

(1) Written rules. M+C organizations must maintain written rules on how to obtain health benefits through the POS benefit.

(2) Evidence of coverage document. The M+C organization must provide to beneficiaries enrolling in a plan with a POS benefit an “evidence of coverage” document, or otherwise provide written documentation, that specifies all costs and possible financial risks to the enrollee, including—

(i) Any premiums and cost-sharing for which the enrollee is responsible;

(ii) Annual limits on benefits and on out-of-pocket expenditures;

(iii) Potential financial responsibility for services for which the plan denies payment because they were not covered under the POS benefit, or exceeded the dollar limit for the benefit; and

(iv) The annual maximum out-of-pocket expense an enrollee could incur.

(e) Prompt payment. Health benefits payable under the POS benefit are subject to the prompt payment requirements in § 422.520.

(f) POS-related data. An M+C organization that offers a POS benefit through an M+C plan must report enrollee utilization data at the plan level by both plan contracting providers (in-network) and by non-contracting providers (out-of-network) including enrollee use of the POS benefit, in the form and manner prescribed by HCFA.


§ 422.106 Coordination of benefits with employer group health plans and Medicaid.

(a) General rule. If an M+C organization contracts with an employer group health plan (EGHP) that covers enrollees in an M+C plan, or contracts with a State Medicaid agency to provide Medicaid benefits to individuals who are eligible for both Medicare and Medicaid, and who are enrolled in an M+C plan, the enrollees must be provided the same benefits as all other enrollees in the M+C plan, with the EGHP or Medicaid benefits supplementing the M+C plan benefits. Jurisdiction regulating benefits under these circumstances is as follows:

(1) All requirements of this part that apply to the M+C program apply to the M+C plan coverage provided to enrollees eligible for benefits under an EGHP or Medicaid contract.

(2) Employer benefits that complement an M+C plan, and the marketing materials associated with the benefits, are not subject to review or approval by HCFA. M+C plan benefits provided to members of the EGHP, and the associated marketing materials, are subject to HCFA review and approval.

(3) Medicaid benefits are not reviewed under this part, but are subject to appropriate HCFA review under the Medicaid program. M+C plan benefits provided to individuals entitled to Medicaid benefits provided by the M+C organization under a contract with the State Medicaid agency are subject to M+C rules and requirements.

(b) Examples. Employer/Medicaid benefits, permissible EGHP or Medicaid plan benefits include the following:
§ 422.108 Medicare secondary payer (MSP) procedures.

(a) Basic rule. HCFA does not pay for services to the extent that Medicare is the primary payer under section 1862(b) of the Act and part 411 of this chapter.

(b) Responsibilities of the M+C organization. The M+C organization must, for each M+C plan—

(1) Identify payers that are primary to Medicare under section 1862(b) of the Act and part 411 of this chapter;

(2) Identify the amounts payable by those payers; and

(3) Coordinate its benefits to Medicare enrollees with the benefits of the primary payers.

(c) Collecting from other entities. The M+C organization may bill, or authorize a provider to bill, other individuals or entities for covered Medicare services for which Medicare is not the primary payer, as specified in paragraphs (d) and (e) of this section.

(d) Collecting from other insurers or the enrollee. If a Medicare enrollee receives services from an M+C organization covered by an M+C organization, the enrollee may receive services from an M+C organization covered services that are also covered under State or Federal workers’ compensation, any no-fault insurance, or any liability insurance policy or plan, including a self-insured plan, the M+C organization may bill, or authorize a provider to bill any of the following—

(1) The insurance carrier, the employer, or any other entity that is liable for payment for the services under section 1862(b) of the Act and part 411 of this chapter.

(2) The Medicare enrollee, to the extent that he or she has been paid by the carrier, employer, or entity for covered medical expenses.

(e) Collecting from group health plans (GHPs) and large group health plans (LGHPs). An M+C organization may bill a GHP or LGHP for services it furnishes to a Medicare enrollee who is also covered under the GHP or LGHP and may bill the Medicare enrollee to the extent that he or she has been paid by the GHP or LGHP.

(f) MSP rules and State laws. Consistent with § 422.402 concerning the Federal preemption of State law, the rules established under this section supersede any State laws, regulations, contract requirements, or other standards that would otherwise apply to M+C plans only to the extent that those State laws are inconsistent with the standards established under this part. A State cannot take away an M+C organization’s right under Federal law and the MSP regulations to bill, or to authorize providers and suppliers to bill, for services for which Medicare is not the primary payer. Section 1952(a)(4) of the Social Security Act does not prohibit a State from limiting the amount of the recovery; thus, State law could modify, but not negate, an M+C organization’s rights in this regard.

[65 FR 40320, June 29, 2000]

§ 422.109 Effect of national coverage determinations (NCDs).

(a) If HCFA determines and announces that an NCD meets the criteria for “significant cost” described in paragraph (c) of this section, an M+C organization is not required to assume risk for the costs of that service until the contract year for which the annual M+C capitation rate is determined on a basis that includes the cost of the NCD service.

(b) The M+C organization must furnish, arrange or pay for an NCD “significant cost” service before the adjustment of the annual M+C capitation rate. The following rules apply to these services:

(1) Medicare payment for the service is:

(i) In addition to the capitation payment to the M+C organization; and

(ii) Made directly by the fiscal intermediary and carrier to the M+C organization in accordance with original
Medicare payment rules, methods, and requirements.

(2) NCD costs for which HCFA intermediaries and carriers will not make payment and are the responsibility of the M+C organization are—

(i) Services necessary to diagnose a condition covered by the NCD;

(ii) Most services furnished as follow-up care to the NCD service;

(iii) Any service that is already a Medicare-covered service and included in the annual M+C capitation rate; and

(iv) Any service, including the costs of the NCD service itself, to the extent the M+C organization is already obligated to cover it as an additional benefit under §422.102 or supplemental benefit under §422.102.

(3) NCD costs for which HCFA intermediaries and carriers make payment are—

(i) Costs relating directly to the provision of services related to the NCD that were noncovered services prior to the issuance of the NCD; and

(ii) A service that is not included in the M+C per capita payment rate.

(4) If the M+C organization does not provide or arrange for the service consistent with HCFA's NCD, enrollees may obtain the services through qualified providers not under contract to the M+C organization, and the organization will pay for the services consistent with §422.109(c).

(5) Beneficiaries are liable for any applicable coinsurance amounts, but are not responsible for the Part A deductible.

(c) The term “significant cost” as it relates to a particular NCD means either of the following:

(1) The average cost of furnishing a single service exceeds a cost threshold that—

(i) For calendar years 1998 and 1999, is $100,000;

(ii) For calendar year 2000 and subsequent calendar years, is the preceding year’s dollar threshold adjusted to reflect the national per capita growth percentage described in §422.254(b).

(2) The estimated cost of all Medicare services furnished nationwide as a result of a particular NCD represents at least 0.1 percent of the national standardized annual capitation rate (see §422.254(f)), multiplied by the total number of Medicare beneficiaries nationwide for the applicable calendar year.

§ 422.111 Disclosure requirements.

(a) Detailed description. An M+C organization must disclose the information specified in paragraph (b) of this section—

§ 422.110 Discrimination against beneficiaries prohibited.

(a) General prohibition. Except as provided in paragraph (b) of this section, an M+C organization may not deny, limit, or condition the coverage or furnishing of benefits to individuals eligible to enroll in an M+C plan offered by the organization on the basis of any factor that is related to health status, including, but not limited to the following:

(1) Medical condition, including mental as well as physical illness.

(2) Claims experience.

(3) Receipt of health care.

(4) Medical history.

(5) Genetic information.

(6) Evidence of insurability, including conditions arising out of acts of domestic violence.

(7) Disability.

(b) Exception. An M+C organization may not enroll an individual who has been medically determined to have end-stage renal disease. However, an enrollee who develops end-stage renal disease while enrolled in a particular M+C organization may not be disenrolled for that reason. An individual who is an enrollee of a particular M+C organization, and resides in the M+C plan service area at the time he or she first becomes M+C eligible, is considered to be “enrolled” in the M+C organization for purposes of the preceding sentence.

(c) Additional requirements. An M+C organization is required to observe the provisions of the Civil Rights Act, Age Discrimination Act, Rehabilitation Act of 1973, and Americans with Disabilities Act (see §422.502(h)).
(f) Content of plan description. The description must include the following information:

(1) Service area. The M+C plan's service area and any enrollment continuation area.

(2) Benefits. The benefits offered under the plan, including applicable conditions and limitations, premiums and cost-sharing (such as copayments, deductibles, and coinsurance) and any other conditions associated with receipt or use of benefits; and for purposes of comparison—

(i) The benefits offered under original Medicare, including the content specified in paragraph (f)(1) of this section;

(ii) For an M+C MSA plan, the benefits under other types of M+C plans; and

(iii) The availability of the Medicare hospice option and any approved hospices in the service area, including those the M+C organization owns, controls, or has a financial interest in.

(3) Access. The number, mix, and distribution (addresses) of providers from whom enrollees may obtain services; any out-of-network coverage; any point-of-service option, including the supplemental premium for that option; and how the M+C organization meets the requirements of §§ 422.112 and 422.114 for access to services offered under the plan.

(4) Out-of-area coverage provided under the plan, including coverage provided to individuals eligible to enroll in the plan under §422.50(a)(3)(i).

(5) Emergency coverage. Coverage of emergency services, including—

(i) Explanation of what constitutes an emergency, referencing the definitions of emergency services and emergency medical condition at §422.113;

(ii) The appropriate use of emergency services, stating that prior authorization cannot be required;

(iii) The process and procedures for obtaining emergency services, including use of the 911 telephone system or its local equivalent; and

(iv) The locations where emergency care can be obtained and other locations at which contracting physicians and hospitals provide emergency services and post-stabilization care included in the M+C plan.

(6) Supplemental benefits. Any mandatory or optional supplemental benefits and the premium for those benefits.

(7) Prior authorization and review rules. Prior authorization rules and other review requirements that must be met in order to ensure payment for the services. The M+C organization must instruct enrollees that, in cases where noncontracting providers submit a bill directly to the enrollee, the enrollee should not pay the bill, but submit it to the M+C organization for processing and determination of enrollee liability, if any.

(8) Grievance and appeals procedures. All grievance and appeals rights and procedures.

(9) Quality assurance program. A description of the quality assurance program required under §422.152.

(10) Disenrollment rights and responsibilities.

(c) Disclosure upon request. Upon request of an individual eligible to elect an M+C plan, an M+C organization must provide to the individual the following information:

(1) The information required paragraph (f) of this section.

(2) The procedures the organization uses to control utilization of services and expenditures.

(3) The number of disputes, and the disposition in the aggregate, in a manner and form described by the Secretary. Such disputes shall be categorized as

(i) Grievances according to §422.564; and

(ii) Appeals according to §422.578 et. seq.

(4) A summary description of the method of compensation for physicians.

(5) Financial condition of the M+C organization, including the most recently audited information regarding, at least, a description of the financial condition of the M+C organization offering the plan.
(d) Changes in rules. If an M+C organization intends to change its rules for an M+C plan, it must:
   (1) Submit the changes for HCFA review under the procedures of §422.80.
   (2) For changes that take effect on January 1, notify all enrollees by the previous October 15.
   (3) For all other changes, notify all enrollees at least 30 days before the intended effective date of the changes.
(e) Changes to provider network. The M+C organization must make a good faith effort to provide written notice of a termination of a contracted provider at least 30 calendar days before the termination effective date to all enrollees who are patients seen on a regular basis by the provider whose contracted is terminating, irrespective of whether the termination was for cause or without cause. When a contract termination involves a primary care professional, all enrollees who are patients of that primary care professional must be notified.
(f) Disclosable information—
   (1) Benefits under original Medicare. (i) Covered services.
   (ii) Beneficiary cost-sharing, such as deductibles, coinsurance, and copayment amounts.
   (iii) Any beneficiary liability for balance billing.
   (2) Enrollment procedures. Information and instructions on how to exercise election options under this subpart.
   (3) Rights. A general description of procedural rights (including grievance and appeals procedures) under original Medicare and the M+C program and the right to be protected against discrimination based on factors related to health status in accordance with §422.110.
   (4) Medigap and Medicare Select. A general description of the benefits, enrollment rights, and requirements applicable to Medicare supplemental policies under section 1882 of the Act, and provisions relating to Medicare Select policies under section 1882(t) of the Act.
   (5) Potential for contract termination. The fact that an M+C organization may terminate or refuse to renew its contract, or reduce the service area included in its contract, and the effect that any of those actions may have on individuals enrolled in that organization's M+C plan.
   (6) Comparative information. A list of M+C plans that are or will be available to residents of the service area in the following calendar year, and, for each available plan, information on the aspects described in paragraphs (c)(7) through (c)(11) of this section, presented in a manner that facilitates comparison among the plans.
   (7) Benefits. (i) Covered services beyond those provided under original Medicare.
   (ii) Any beneficiary cost-sharing.
   (iii) Any maximum limitations on out-of-pocket expenses.
   (iv) In the case of an M+C MSA plan, the amount of the annual MSA deposit and the differences in cost-sharing, enrollee premiums, and balance billing, as compared to M+C plans.
   (v) In the case of an M+C private fee-for-service plan, differences in cost-sharing, enrollee premiums, and balance billing, as compared to M+C plans.
   (vi) The extent to which an enrollee may obtain benefits through out-of-network health care providers.
   (vii) The types of providers that participate in the plan's network and the extent to which an enrollee may select among those providers.
   (viii) The coverage of emergency and urgently needed services.
   (8) Premiums. (i) The M+C monthly basic beneficiary premiums.
   (ii) The M+C monthly supplemental beneficiary premium.
   (9) The plan's service area.
   (10) Quality and performance indicators for benefits under a plan to the extent they are available as follows (and how they compare with indicators under original Medicare):
   (i) Disenrollment rates for Medicare enrollees for the 2 previous years, excluding disenrollment due to death or moving outside the plan's service area, calculated according to HCFA guidelines.
   (ii) Medicare enrollee satisfaction.
   (iii) Health outcomes.
   (iv) Plan-level appeal data.
   (v) The recent record of plan compliance with the requirements of this part, as determined by the Secretary.
   (vi) Other performance indicators.
§ 422.112 Access to services.

(a) Rules for coordinated care plans and network M+C MSA plans. An M+C organization that offers an M+C coordinated care plan or network M+C MSA plan may specify the networks of providers from which enrollees may obtain services if the M+C organization ensures that all covered services, including additional or supplemental services contracted for by (or on behalf of) the Medicare enrollee, are available and accessible under the plan. To accomplish this, the M+C organization must meet the following requirements:

(1) Provider network. Maintain and monitor a network of appropriate providers that is supported by written agreements and is sufficient to provide adequate access to covered services to meet the needs of the population served. These providers are typically utilized in the network as primary care providers (PCPs), specialists, hospitals, skilled nursing facilities, home health agencies, ambulatory clinics, and other providers.

(2) PCP panel. Establish a panel of PCPs from which the enrollee may select a PCP. If an M+C organization requires its enrollees to obtain a referral in most situations before receiving services from a specialist, the M+C organization must either assign a PCP for purposes of making the needed referral or make other arrangements to ensure access to medically necessary specialty care.

(3) Specialty care. Provide or arrange for necessary specialty care, and in particular give women enrollees the option of direct access to a women’s health specialist within the network for women’s routine and preventive health care services provided as basic benefits (as defined in §422.2). The M+C organization arranges for specialty care outside of the plan provider network when network providers are unavailable or inadequate to meet an enrollee’s medical needs.

(4) Serious medical conditions. Ensure that for each plan, the M+C organization has in effect HCFA-approved procedures that enable the M+C organization, through appropriate health care professionals, to—

(i) Identify individuals with complex or serious medical conditions;

(ii) Assess those conditions, and use medical procedures to diagnose and monitor them on an ongoing basis; and

(iii) Establish and implement a treatment plan that—

(A) Is appropriate to those conditions;

(B) Includes an adequate number of direct access visits to specialists consistent with the treatment plan;

(C) Is time-specific and updated periodically; and

(D) Ensures adequate coordination of care among providers.

(5) Service area expansion. If seeking a service area expansion for an M+C plan, demonstrate that the number and type of providers available to plan enrollees are sufficient to meet projected needs of the population to be served.

(6) Credentialed providers. Demonstrate to HCFA that its providers in an M+C plan are credentialed through the process set forth at §422.204(a).

(7) Written standards. Establish written standards for the following:

(i) Timeliness of access to care and member services that meet or exceed standards established by HCFA. Timely access to care and member services within a plan’s provider network must be continuously monitored to ensure compliance with these standards, and the M+C organization must take corrective action as necessary.

(ii) Policies and procedures (coverage rules, practice guidelines, payment policies, and utilization management) that allow for individual medical necessity determinations.

(iii) Provider consideration of beneficiary input into the provider’s proposed treatment plan.

(8) Hours of operation. Ensure that—

(i) The hours of operation of its M+C plan providers are convenient to the population served under the plan and do not discriminate against Medicare enrollees; and
(ii) Plan services are available 24 hours a day, 7 days a week, when medically necessary.

(9) Cultural considerations. Ensure that services are provided in a culturally competent manner to all enrollees, including those with limited English proficiency or reading skills, and diverse cultural and ethnic backgrounds.

(10) Ambulance services, emergency and urgently needed services, and post-stabilization care services coverage. Provide coverage for ambulance services, emergency and urgently needed services, and post-stabilization care services in accordance with §422.113.

(b) Rules for all M+C organizations to ensure continuity of care. The M+C organization must ensure continuity of care and integration of services through arrangements that include, but are not limited to the following—

(1) Policies that specify under what circumstances services are coordinated and the methods for coordination;

(2) Offering to provide each enrollee with an ongoing source of primary care and providing a primary care source to each enrollee who accepts the offer;

(3) Programs for coordination of plan services with community and social services generally available through contracting or noncontracting providers in the area served by the M+C plan, including nursing home and community-based services; and

(4) Procedures to ensure that the M+C organization and its provider network have the information required for effective and continuous patient care and quality review, including procedures to ensure that—

(i) The M+C organization makes a "best-effort" attempt to conduct an initial assessment of each enrollee’s health care needs, including following up on unsuccessful attempts to contact an enrollee, within 90 days of the effective date of enrollment;

(ii) Each provider, supplier, and practitioner furnishing services to enrollees maintains an enrollee health record in accordance with standards established by the M+C organization, taking into account professional standards; and

(iii) There is appropriate and confidential exchange of information among provider network components.

(5) Procedures to ensure that enrollees are informed of specific health care needs that require follow-up and receive, as appropriate, training in self-care and other measures they may take to promote their own health; and

(6) Systems to address barriers to enrollee compliance with prescribed treatments or regimens.

[64 FR 7980, Feb. 17, 1999, as amended at 65 FR 40321, June 29, 2000]
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area (or, under unusual and extraordinary circumstances, provided when the enrollee is in the service or continuation area but the organization’s provider network is temporarily unavailable or inaccessible) when the services are medically necessary and immediately required—

(A) As a result of an unforeseen illness, injury, or condition; and

(B) It was not reasonable given the circumstances to obtain the services through the organization offering the M+C plan.

(2) M+C organization financial responsibility. The M+C organization is financially responsible for emergency and urgently needed services—

(i) Regardless of whether the services are obtained within or outside the M+C organization;

(ii) Regardless of whether there is prior authorization for the services.

(A) Instructions to seek prior authorization for emergency or urgently needed services may not be included in any materials furnished to enrollees (including wallet card instructions), and enrollees must be informed of their right to call 911.

(B) Instruction to seek prior authorization before the enrollee has been stabilized may not be included in any materials furnished to providers (including contracts with providers);

(iii) In accordance with the prudent layperson definition of emergency medical condition regardless of final diagnosis;

(iv) For which a plan provider or other M+C organization representative instructs an enrollee to seek emergency services within or outside the plan; and

(v) With a limit on charges to enrollees for emergency services of $50 or what it would charge the enrollee if he or she obtained the services through the M+C organization, whichever is less.

(3) Stabilized condition. The physician treating the enrollee must decide when the enrollee may be considered stabilized for transfer or discharge, and that decision is binding on the M+C organization.

(c) Maintenance care and post-stabilization care services (hereafter referred to as “post-stabilization care services”).

(1) Definition. Post-stabilization care services means covered services, related to an emergency medical condition, that are provided after an enrollee is stabilized in order to maintain the stabilized condition, or, under the circumstances described in paragraph (c)(2)(iii) of this section, to improve or resolve the enrollee’s condition.

(2) M+C organization financial responsibility. The M+C organization—

(i) Is financially responsible (consistent with §422.214) for post-stabilization care services obtained within or outside the M+C organization that are pre-approved by a plan provider or other M+C organization representative;

(ii) Is financially responsible for post-stabilization care services obtained within or outside the M+C organization that are not pre-approved by a plan provider or other M+C organization representative, but administered to maintain the enrollee’s stabilized condition within 1 hour of a request to the M+C organization for pre-approval of further post-stabilization care services;

(iii) Is financially responsible for post-stabilization care services obtained within or outside the M+C organization that are not pre-approved by a plan provider or other M+C organization representative, but administered to maintain, improve, or resolve the enrollee’s stabilized condition if—

(A) The M+C organization does not respond to a request for pre-approval within 1 hour;

(B) The M+C organization cannot be contacted; or

(C) The M+C organization representative and the treating physician cannot reach an agreement concerning the enrollee’s stabilized condition if a plan physician is not available for consultation. In this situation, the M+C organization must give the treating physician the opportunity to consult with a plan physician and the treating physician may continue with care of the patient until a plan physician is reached or one of the criteria in §422.113(c)(3) is met; and

(iv) Must limit charges to enrollees for post-stabilization care services to an amount no greater than what the organization would charge the enrollee
if he or she had obtained the services through the M+C organization.
(3) End of M+C organization’s financial responsibility. The M+C organization’s financial responsibility for post-stabilization care services it has not pre-approved ends when—
(i) A plan physician with privileges at the treating hospital assumes responsibility for the enrollee’s care;
(ii) A plan physician assumes responsibility for the enrollee’s care through transfer;
(iii) An M+C organization representative and the treating physician reach an agreement concerning the enrollee’s care; or
(iv) The enrollee is discharged.

[65 FR 40322, June 29, 2000]

§ 422.114 Access to services under an M+C private fee-for-service plan.
(a) Sufficient access. (1) An M+C organization that offers an M+C private fee-for-service plan must demonstrate to HCFA that it has sufficient number and range of providers willing to furnish services under the plan.
(2) HCFA finds that an M+C organization meets the requirement in paragraph (a)(1) of this section if, with respect to a particular category of health care providers, the M+C organization has—
(i) Payment rates that are not less than the rates that apply under original Medicare for the provider in question;
(ii) Contracts or agreements with a sufficient number and range of providers to furnish the services covered under the M+C private fee-for-service plan; or
(iii) A combination of paragraphs (a)(2)(i) and (a)(2)(ii) of this section.
(b) Freedom of choice. M+C fee-for-service plans must permit enrollees to obtain services from any entity that is authorized to provide services under Medicare Part A and Part B and agrees to provide services under the terms of the plan.

§ 422.118 Confidentiality and accuracy of enrollee records.
For any medical records or other health and enrollment information it maintains with respect to enrollees, an M+C organization must establish procedures to do the following:
(a) Abide by all Federal and State laws regarding confidentiality and disclosure of medical records, or other health and enrollment information. The M+C organization must safeguard the privacy of any information that identifies a particular enrollee and have procedures that specify—
(1) For what purposes the information will be used within the organization; and
(2) To whom and for what purposes it will disclose the information outside the organization.
(b) Ensure that medical information is released only in accordance with applicable Federal or State law, or pursuant to court orders or subpoenas.
(c) Maintain the records and information in an accurate and timely manner.
(d) Ensure timely access by enrollees to the records and information that pertain to them.

[65 FR 40323, June 29, 2000]

§ 422.128 Information on advance directives.
(a) Each M+C organization must maintain written policies and procedures that meet the requirements for advance directives, as set forth in subpart I of part 489 of this chapter. For purposes of this part, advance directive has the meaning given the term in §489.100 of this chapter.
(b) An M+C organization must maintain written policies and procedures concerning advance directives with respect to all adult individuals receiving medical care by or through the M+C organization.
(1) An M+C organization must provide written information to those individuals with respect to the following:
(i) Their rights under the law of the State in which the organization furnishes services (whether statutory or recognized by the courts of the State) to make decisions concerning their medical care, including the right to accept or refuse medical or surgical treatment and the right to formulate advance directives. Providers may contract with other entities to furnish this
information but remain legally responsible for ensuring that the requirements of this section are met. The information must reflect changes in State law as soon as possible, but no later than 90 days after the effective date of the State law.

(ii) The M+C organization’s written policies respecting the implementation of those rights, including a clear and precise statement of limitation if the M+C organization cannot implement an advance directive as a matter of conscience. At a minimum, this statement must do the following:

(A) Clarify any differences between institution-wide conscientious objections and those that may be raised by individual physicians.
(B) Identify the state legal authority permitting such objection.
(C) Describe the range of medical conditions or procedures affected by the conscience objection.
(D) Provide the information specified in paragraph (a)(1) of this section to each enrollee at the time of initial enrollment. If an enrollee is incapacitated at the time of initial enrollment and is unable to receive information (due to the incapacitating condition or a mental disorder) or articulate whether or not he or she has executed an advance directive, the M+C organization may give advance directive information to the enrollee’s family or surrogate in the same manner that it issues other materials about policies and procedures to the family of the incapacitated enrollee or to a surrogate or other concerned persons in accordance with State law. The M+C organization is not relieved of its obligation to provide this information to the enrollee once he or she is no longer incapacitated or unable to receive such information. Follow-up procedures must be in place to ensure that the information is given to the individual directly at the appropriate time.
(E) Document in a prominent part of the individual’s current medical record whether or not the individual has executed an advance directive.
(F) Not condition the provision of care or otherwise discriminate against an individual based on whether or not the individual has executed an advance directive.

(G) Ensure compliance with requirements of State law (whether statutory or recognized by the courts of the State) regarding advance directives.
(H) Provide for education of staff concerning its policies and procedures on advance directives.
(I) Provide for community education regarding advance directives that may include material required in paragraph (a)(1)(i) of this section, either directly or in concert with other providers or entities. Separate community education materials may be developed and used, at the discretion of the M+C organization. The same written materials are not required for all settings, but the material should define what constitutes an advance directive, emphasizing that an advance directive is designed to enhance an incapacitated individual’s control over medical treatment, and describe applicable State law concerning advance directives. An M+C organization must be able to document its community education efforts.

(2) The M+C organization—
(i) Is not required to provide care that conflicts with an advance directive; and
(ii) Is not required to implement an advance directive if, as a matter of conscience, the M+C organization cannot implement an advance directive and State law allows any health care provider or any agent of the provider to conscientiously object.

(3) The M+C organization must inform individuals that complaints concerning noncompliance with the advance directive requirements may be filed with the State survey and certification agency.

§ 422.132 Protection against liability and loss of benefits.

Enrollees of M+C organizations are entitled to the protections specified in §422.502(g).

Subpart D—Quality Assurance

SOURCE: 63 FR 35082, June 26, 1998, unless otherwise noted.
§ 422.152 Quality assessment and performance improvement program.

(a) General rule. Each M+C organization that offers one or more M+C plans must have, for each of those plans, an ongoing quality assessment and performance improvement program that meets the applicable requirements of this section for the services it furnishes to its M+C enrollees.

(b) Requirements for network M+C MSA plans and M+C coordinated care plans other than PPO plans. An organization offering a network M+C MSA plan or M+C coordinated care plan other than a PPO plan must do the following:

(1) Meet the requirements in paragraph (c)(1) of this section concerning performance measurement and reporting. With respect to an M+C coordinated care plan, an organization must also meet the requirements of paragraph (c)(2) of this section concerning the achievement of minimum performance levels. The requirements of paragraph (c)(2) of this section do not apply with respect to an M+C MSA plan.

(2) Conduct performance improvement projects as described in paragraph (d) of this section. These projects must achieve, through ongoing measurement and intervention, demonstrable and sustained improvement in significant aspects of clinical care and nonclinical care areas that can be expected to have a favorable effect on health outcomes and enrollee satisfaction.

(3) In processing requests for initial or continued authorization of services, follow written policies and procedures that reflect current standards of medical practice.

(4) Have in effect mechanisms to detect both underutilization and overutilization of services.

(5) Make available to HCFA information on quality and outcomes measures that will enable beneficiaries to compare health coverage options and select among them, as provided in §422.54(c)(1).

(c) Performance measurement and reporting. The organization offering the plan must do the following:

(1) Measure performance under the plan, using standard measures required by HCFA, and report its performance to HCFA. The standard measures may be specified in uniform data collection and reporting instruments required by HCFA, and will relate to—

(i) Clinical areas including effectiveness of care, enrollee perception of care, and use of services; and

(ii) Nonclinical areas including access to and availability of services, appeals and grievances, and organizational characteristics.

(2) Achieve any minimum performance levels that HCFA establishes locally, regionally, or nationally with respect to the standard measures.

(i) In establishing minimum performance levels, HCFA considers historical plan and original Medicare performance data and trends.

(ii) HCFA establishes the minimum performance levels prospectively upon contract initiation and renewal.

(iii) The organization must meet the minimum performance levels by the end of the contract year.

(iv) In accordance with §422.506, HCFA may decline to renew the organization’s contract in the year that HCFA determines that it did not meet the minimum performance levels.

(d) Performance improvement projects.

(1) Performance improvement projects are organization initiatives that focus on specified clinical and nonclinical areas and that involve the following:

(i) Measurement of performance.

(ii) System interventions, including the establishment or alteration of practice guidelines.

(iii) Improving performance.

(iv) Systematic follow-up on the effect of the interventions.

(2) Each project must address the entire population to which the measurement specified in paragraph (d)(1)(i) of this section is relevant.

(3) HCFA establishes M+C organization and M+C plan-specific obligations for the number and distribution of projects among the required clinical and nonclinical areas, in accordance with paragraphs (d)(4) and (d)(5) of this section, to ensure that the projects are representative of the entire spectrum of clinical and nonclinical care areas associated with a plan.

(4) The required clinical areas include:

(i) Prevention and care of acute and chronic conditions.
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(ii) High-volume services.
(iii) High-risk services.
(iv) Continuity and coordination of care.
(5) The required nonclinical areas include:
   (i) Appeals, grievances, and other complaints.
   (ii) Access to, and availability of, services.
(6) In addition to requiring that the organization initiate its own performance improvement projects, HCFA may require that the organization—
   (i) Conduct particular performance improvement projects that are specific to the organization; and
   (ii) Participate in national or statewide performance improvement projects.
(7) For each project, the organization must assess performance under the plan using quality indicators that are—
   (i) Objective, clearly and unambiguously defined, and based on current clinical knowledge or health services research; and
   (ii) Capable of measuring outcomes such as changes in health status, functional status and enrollee satisfaction, or valid proxies of those outcomes.
(8) Performance assessment on the selected indicators must be based on systematic ongoing collection and analysis of valid and reliable data.
(9) Interventions must achieve improvement that is significant and sustained over time.
(10) The organization must report the status and results of each project to HCFA as requested.
(e) Requirements for M+C PPO plans, non-network MSA plans, and M+C private fee-for-service plans. An organization offering an M+C plan, non-network MSA plan, or private fee-for-service plan must do the following:
   (1) Measure performance under the plan using standard measures required by HCFA and report its performance to HCFA. The standard measures may be specified in uniform data collection and reporting instruments required by HCFA and will relate to—
      (i) Clinical areas including effectiveness of care, enrollee perception of care, and use of services; and
      (ii) Nonclinical areas including access to and availability of services, appeals and grievances, and organizational characteristics.
   (2) Evaluate the continuity and coordination of care furnished to enrollees.
   (3) If the organization uses written protocols for utilization review, the organization must—
      (i) Base those protocols on current standards of medical practice; and
      (ii) Have mechanisms to evaluate utilization of services and to inform enrollees and providers of services of the results of the evaluation.
(f) Requirements for all types of plans—
   (1) Health information. For all types of plans that it offers, an organization must—
      (i) Maintain a health information system that collects, analyzes, and integrates the data necessary to implement its quality assessment and performance improvement program;
      (ii) Ensure that the information it receives from providers of services is reliable and complete; and
      (iii) Make all collected information available to HCFA.
   (2) Program review. For each plan, there must be in effect a process for formal evaluation, at least annually, of the impact and effectiveness of its quality assessment and performance improvement program.
   (3) Remedial action. For each plan, the organization must correct all problems that come to its attention through internal surveillance, complaints, or other mechanisms.
[63 FR 35082, June 26, 1998, as amended at 65 FR 40323, June 29, 2000]

§ 422.154  External review.

(a) Basic rule. Except as provided in paragraph (c) of this section, each M+C organization must, for each M+C plan it operates, have an agreement with an independent quality review and improvement organization (review organization) approved by HCFA to perform functions of the type described in part 466 of this chapter.
(b) Terms of the agreement. The agreement must be consistent with HCFA guidelines and include the following provisions:
   (1) Require that the organization—
(i) Allocate adequate space for use of the review organization whenever it is conducting review activities; and

(ii) Provide all pertinent data, including patient care data, at the time the review organization needs the data to carry out the reviews and make its determinations.

(2) Except in the case of complaints about quality, exclude review activities that HCFA determines would duplicate review activities conducted as part of an approved accreditation process or as part of HCFA monitoring.

(c) Exceptions. The requirement of paragraph (a) of this section does not apply for an M+C private fee-for-service plan or a non-network M+C MSA plan if the organization does not carry out utilization review with respect to the plan.

[63 FR 35082, June 26, 1998, as amended at 65 FR 40323, June 29, 2000]

§ 422.156 Compliance deemed on the basis of accreditation.

(a) General rule. An M+C organization is deemed to meet all of the requirements of any of the areas described in paragraph (b) of this section if—

(1) The M+C organization is fully accredited (and periodically reaccredited) for the standards related to the applicable area under paragraph (b) of this section by a private, national accreditation organization approved by HCFA; and

(2) The accreditation organization used the standards approved by HCFA for the purposes of assessing the M+C organization’s compliance with Medicare requirements.

(b) Deemable requirements. The requirements relating to the following areas are deemable:

(1) Quality assurance.
(2) Antidiscrimination.
(3) Access to services.
(4) Confidentiality and accuracy of enrollee records.
(5) Information on advance directives.
(6) Provider participation rules.

(c) Effective date of deemed status. The date on which the organization is deemed to meet the applicable requirements is the later of the following:

(1) The date the M+C organization is accredited by the accreditation organization.

(d) Obligations of deemed M+C organizations. An M+C organization deemed to meet Medicare requirements must—

(1) Submit to surveys by HCFA to validate its accreditation organization’s accreditation process; and

(2) Authorize its accreditation organization to release to HCFA a copy of its most recent accreditation survey, together with any survey-related information that HCFA may require (including corrective action plans and summaries of unmet HCFA requirements).

(e) Removal of deemed status. HCFA removes part or all of an M+C organization’s deemed status for any of the following reasons:

(1) HCFA determines, on the basis of its own survey or the results of the accreditation survey, that the M+C organization does not meet the Medicare requirements for which deemed status was granted.

(2) HCFA withdraws its approval of the accreditation organization that accredited the M+C organization.

(3) The M+C organization fails to meet the requirements of paragraph (d) of this section.

(f) Enforcement authority. HCFA retains the authority to initiate enforcement action against any M+C organization that it determines, on the basis of its own survey or the results of an accreditation survey, no longer meets the Medicare requirements for which deemed status was granted.

[63 FR 35082, June 26, 1998, as amended at 65 FR 40323, June 29, 2000]

§ 422.157 Accreditation organizations.

(a) Conditions for approval. HCFA may approve an accreditation organization with respect to a given standard under this part if it meets the following conditions:

(1) In accrediting M+C organizations, it applies and enforces standards that are at least as stringent as Medicare requirements with respect to the standard or standards in question.

(2) It complies with the application and reapplication procedures set forth in §422.158.

(3) It ensures that:
(i) Any individual associated with it, who is also associated with an entity it accredits, does not influence the accreditation decision concerning that entity.

(ii) The majority of the membership of its governing body is not comprised of managed care organizations or their representatives.

(iii) Its governing body has a broad and balanced representation of interests and acts without bias.

(b) Notice and comment—(1) Proposed notice. HCFA publishes a notice in the FEDERAL REGISTER whenever it is considering granting an accreditation organization’s application for approval. The notice—

(i) Announces HCFA’s receipt of the accreditation organization’s application for approval;

(ii) Describes the criteria HCFA will use in evaluating the application; and

(iii) Provides at least a 30-day comment period.

(2) Final notice. (i) After reviewing public comments, HCFA publishes a final FEDERAL REGISTER notice indicating whether it has granted the accreditation organization’s request for approval.

(ii) If HCFA grants the request, the final notice specifies the effective date and the term of the approval, which may not exceed 6 years.

(c) Ongoing responsibilities of an approved accreditation organization. An accreditation organization approved by HCFA must undertake the following activities on an ongoing basis:

(1) Provide to HCFA in written form and on a monthly basis all of the following:

(i) Copies of all accreditation surveys, together with any survey-related information that HCFA may require (including corrective action plans and summaries of unmet HCFA requirements).

(ii) Notice of all accreditation decisions.

(iii) Notice of all complaints related to deemed M+C organizations.

(iv) Information about any M+C organization against which the accrediting organization has taken remedial or adverse action, including revocation, withdrawal or revision of the M+C organization’s accreditation. (The accreditation organization must provide this information within 30 days of taking the remedial or adverse action.)

(v) Notice of any proposed changes in its accreditation standards or requirements or survey process. If the organization implements the changes before or without HCFA approval, HCFA may withdraw its approval of the accreditation organization.

(2) Within 30 days of a change in HCFA requirements, submit to HCFA—

(i) An acknowledgment of HCFA’s notification of the change;

(ii) A revised cross-walk reflecting the new requirements; and

(iii) An explanation of how the accreditation organization plans to alter its standards to conform to HCFA’s new requirements, within the time-frames specified in the notification of change it receives from HCFA.

(3) Permit its surveyors to serve as witnesses if HCFA takes an adverse action based on accreditation findings.

(4) Within 3 days of identifying, in an accredited M+C organization, a deficiency that poses immediate jeopardy to the organization’s enrollees or to the general public, give HCFA written notice of the deficiency.

(5) Within 10 days of HCFA’s notice of withdrawal of approval, give written notice of the withdrawal to all accredited M+C organizations.

(6) Provide, on an annual basis, summary data specified by HCFA that relate to the past year’s accreditation activities and trends.

(d) Continuing Federal oversight of approved accreditation organizations. This paragraph establishes specific criteria and procedures for continuing oversight and for withdrawing approval of an accreditation organization.

(1) Equivalency review. HCFA compares the accreditation organization’s standards and its application and enforcement of those standards to the comparable HCFA requirements and processes when—

(i) HCFA imposes new requirements or changes its survey process;

(ii) An accreditation organization proposes to adopt new standards or changes its survey process; or

(iii) The term of an accreditation organization’s approval expires.
(2) Validation review. HCFA or its agent may conduct a survey of an accredited organization, examine the results of the accreditation organization's own survey, or attend the accreditation organization's survey, in order to validate the organization's accreditation process. At the conclusion of the review, HCFA identifies any accreditation programs for which validation survey results—
(i) Indicate a 20 percent rate of disparity between certification by the accreditation organization and certification by HCFA or its agent on standards that do not constitute immediate jeopardy to patient health and safety if unmet;
(ii) Indicate any disparity between certification by the accreditation organization and certification by HCFA or its agent on standards that constitute immediate jeopardy to patient health and safety if unmet; or
(iii) Indicate that, irrespective of the rate of disparity, there are widespread or systematic problems in an organization's accreditation process such that accreditation no longer provides assurance that the Medicare requirements are met or exceeded.

(3) Onsite observation. HCFA may conduct an onsite inspection of the accreditation organization's operations and offices to verify the organization's representations and assess the organization's compliance with its own policies and procedures. The onsite inspection may include, but is not limited to, reviewing documents, auditing meetings concerning the accreditation process, evaluating survey results or the accreditation status decision making process, and interviewing the organization's staff.

(4) Notice of intent to withdraw approval. If an equivalency review, validation review, onsite observation, or HCFA’s daily experience with the accreditation organization suggests that the accreditation organization is not meeting the requirements of this subpart, HCFA gives the organization written notice of its intent to withdraw approval.

(5) Withdrawal of approval. HCFA may withdraw its approval of an accreditation organization at any time if HCFA determines that—
(i) Deeming based on accreditation no longer guarantees that the M+C organization meets the M+C requirements, and failure to meet those requirements could jeopardize the health or safety of Medicare enrollees and constitute a significant hazard to the public health; or
(ii) The accreditation organization has failed to meet its obligations under this section or under §422.156 or §422.158.

(6) Reconsideration of withdrawal of approval. An accreditation organization dissatisfied with a determination to withdraw HCFA approval may request a reconsideration of that determination in accordance with subpart D of part 488 of this chapter.


§422.158 Procedures for approval of accreditation as a basis for deeming compliance.

(a) Required information and materials. A private, national accreditation organization applying for approval must furnish to HCFA all of the following information and materials. (When reapplying for approval, the organization need furnish only the particular information and materials requested by HCFA.)

(1) The types of M+C plans that it would review as part of its accreditation process.

(2) A detailed comparison of the organization’s accreditation requirements and standards with the Medicare requirements (for example, a crosswalk).

(3) Detailed information about the organization’s survey process, including—
(i) Frequency of surveys and whether surveys are announced or unannounced.

(ii) Copies of survey forms, and guidelines and instructions to surveyors.

(iii) Descriptions of—
(A) The survey review process and the accreditation status decision making process;
(B) The procedures used to notify accredited M+C organizations of deficiencies and to monitor the correction of those deficiencies; and
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(C) The procedures used to enforce compliance with accreditation requirements.

(4) Detailed information about the individuals who perform surveys for the accreditation organization, including—
   (i) The size and composition of accreditation survey teams for each type of plan reviewed as part of the accreditation process;
   (ii) The education and experience requirements surveyors must meet;
   (iii) The content and frequency of the in-service training provided to survey personnel;
   (iv) The evaluation systems used to monitor the performance of individual surveyors and survey teams; and
   (v) The organization’s policies and procedures regarding coordination of these activities with appropriate licensing bodies and ombudsman programs.

(5) A description of the organization’s data management and analysis system with respect to its surveys and accreditation decisions, including the kinds of reports, tables, and other displays generated by that system.

(6) A description of the organization’s procedures for responding to and investigating complaints against accredited organizations, including policies and procedures regarding coordination of these activities with appropriate licensing bodies and ombudsman programs.

(7) A description of the organization’s policies and procedures with respect to the withholding or removal of accreditation for failure to meet the accreditation organization’s standards or requirements, and other actions the organization takes in response to noncompliance with its standards and requirements.

(8) A description of all types (for example, full, partial) and categories (for example, provisional, conditional, temporary) of accreditation offered by the organization, the duration of each type and category of accreditation and a statement identifying the types and categories that would serve as a basis for accreditation if HCFA approves the accreditation organization.

(9) A list of all currently accredited M+C organizations and the type, category, and expiration date of the accreditation held by each of them.

(10) A list of all full and partial accreditation surveys scheduled to be performed by the accreditation organization as requested by HCFA.

(11) The name and address of each person with an ownership or control interest in the accreditation organization.

(b) Required supporting documentation.

A private, national accreditation organization applying or reapplying for approval must also submit the following supporting documentation:

(1) A written presentation that demonstrates its ability to furnish HCFA with electronic data in HCFA compatible format.

(2) A resource analysis that demonstrates that its staffing, funding, and other resources are adequate to perform the required surveys and related activities.

(3) A statement acknowledging that, as a condition for approval, it agrees to comply with the ongoing responsibility requirements of §422.157(c).

(c) Additional information. If HCFA determines that it needs additional information for a determination to grant or deny the accreditation organization’s request for approval, it notifies the organization and allows time for the organization to provide the additional information.

(d) Onsite visit. HCFA may visit the accreditation organization’s offices to verify representations made by the organization in its application, including, but not limited to, review of documents, and interviews with the organization’s staff.

(e) Notice of determination. HCFA gives the accreditation organization, within 210 days of receipt of its completed application, a formal notice that—

(1) States whether the request for approval has been granted or denied;

(2) Gives the rationale for any denial; and

(3) Describes the reconsideration and reapplication procedures.

(f) Withdrawal. An accreditation organization may withdraw its application for approval at any time before it receives the formal notice specified in paragraph (e) of this section.
(g) Reconsideration of adverse determination. An accreditation organization that has received notice of denial of its request for approval may request reconsideration in accordance with subpart D of part 488 of this chapter.

(h) Request for approval following denial. (1) Except as provided in paragraph (h)(2) of this section, an accreditation organization that has received notice of denial of its request for approval may submit a new request if it—
   (i) Has revised its accreditation program to correct the deficiencies on which the denial was based;
   (ii) Can demonstrate that the M+C organizations that it has accredited meet or exceed applicable Medicare requirements; and
   (iii) Resubmits the application in its entirety.

   (2) An accreditation organization that has requested reconsideration of HCFA's denial of its request for approval may not submit a new request until the reconsideration is administratively final.


Subpart E—Relationships With Providers

SOURCE: 63 FR 35085, June 26, 1998, unless otherwise noted.

422.200 Basis and scope.

This subpart is based on sections 1852(a)(1), (a)(2), (b)(2), (c)(2)(D), (j), and (k) of the Act; section 1859(b)(2)(A) of the Act; and the general authority under 1856(b) of the Act requiring the establishment of standards. It sets forth the requirements and standards for the M+C organization's relationships with providers including physicians, other health care professionals, institutional providers and suppliers, under contracts or arrangements or deemed contracts under M+C private fee-for-service plans. This subpart also contains some requirements that apply to noncontracting providers.

§ 422.202 Participation procedures.

(a) Notice and appeal rights. An M+C organization that operates a coordinated care plan or network MSA plan must provide for the participation of individual physicians, and the management and members of groups of physicians, through reasonable procedures that include the following:

   (1) Written notice of rules of participation including terms of payment, credentialing, and other rules directly related to participation decisions.

   (2) Written notice of material changes in participation rules before the changes are put into effect.

   (3) Written notice of participation decisions that are adverse to physicians.

   (4) A process for appealing adverse participation decisions, including the right of physicians to present information and their views on the decision. In the case of a termination or suspension of a provider contract by the M+C organization, this process must conform to the rules in §422.204(c).

   (b) Consultation. The M+C organization must establish a formal mechanism to consult with the physicians who have agreed to provide services under the M+C plan offered by the organization, regarding the organization’s medical policy, quality assurance programs and medical management procedures and ensure that the following standards are met:

   (1) Practice guidelines and utilization management guidelines—
   (i) Are based on reasonable medical evidence or a consensus of health care professionals in the particular field;
   (ii) Consider the needs of the enrolled population;
   (iii) Are developed in consultation with contracting physicians; and
   (iv) Are reviewed and updated periodically.

   (2) The guidelines are communicated to providers and, as appropriate, to enrollees.

   (3) Decisions with respect to utilization management, enrollee education, coverage of services, and other areas in which the guidelines apply are consistent with the guidelines.

   (c) Subcontracted groups. An M+C organization that operates an M+C plan through subcontracted physician groups must provide that the participation procedures in this section apply equally to physicians within those subcontracted groups.
§ 422.204 Provider selection and credentialing.

(a) General rule. An M+C organization must have written policies and procedures for the selection and evaluation of providers. These policies must conform with the credential and recredentialing requirements set forth in paragraph (b) of this section and with the antidiscrimination provisions set forth in § 422.205.

(b) Basic requirements. An M+C organization must follow a documented process with respect to providers and suppliers who have signed contracts or participation agreements that—

(1) For providers (other than physicians and other health care professionals) requires determination, and redetermination at specified intervals, that each provider is—

(ⅰ) Licensed to operate in the State, and in compliance with any other applicable State or Federal requirements; and

(ⅱ) Reviewed and approved by an accrediting body, or meets the standards established by the organization itself;

(2) For physicians and other health care professionals, including members of physician groups, covers—

(ⅰ) Initial credentialing that includes written application, verification of licensure or certification from primary sources, disciplinary status, eligibility for payment under Medicare, and site visits as appropriate. The application must be signed and dated and include an attestation by the applicant of the correctness and completeness of the application and other information submitted in support of the application; and

(ⅱ) Recredentialing at least every 2 years that updates information obtained during initial credentialing and considers performance indicators such as those collected through quality assurance programs, utilization management systems, handling of grievances and appeals, enrollee satisfaction surveys, and other plan activities, and that includes an attestation of the correctness and completeness of the new information; and

(ⅲ) A process for consulting with contracting health care professionals with respect to criteria for credentialing and recredentialing.

(3) Specifies that basic benefits must be provided through, or payments must be made to, providers and suppliers that meet applicable requirements of title XVIII and part A of title XI of the Act. In the case of providers meeting the definition of “provider of services” in section 1861(u) of the Act, basic benefits may only be provided through these providers if they have a provider agreement with HCFA permitting them to provide services under original Medicare.

(4) Ensures compliance with the requirements at § 422.752(a)(8) that prohibit employment or contracts with individuals (or with an entity that employs or contracts with such an individual) excluded from participation.
under Medicare and with the requirements at §422.220 regarding physicians and practitioners who opt out of Medicare.

[65 FR 40324, June 29, 2000]

§ 422.205 Provider antidiscrimination rules.

(a) General rule. Consistent with the requirements of this section, the policies and procedures concerning provider selection and credentialing established under §422.204, and with the requirement under §422.100(c) that all Medicare-covered services be available to M+C plan enrollees, an M+C organization may select the practitioners that participate in its plan provider networks. In selecting these practitioners, an M+C organization may not discriminate, in terms of participation, reimbursement, or indemnification, against any health care professional who is acting within the scope of his or her license or certification under State law, solely on the basis of the license or certification. If an M+C organization declines to include a given provider or group of providers in its network, it must furnish written notice to the affected provider(s) of the reason for the decision.

(b) Construction. The prohibition in paragraph (a)(1) of this section does not preclude any of the following by the M+C organization:

(1) Refusal to grant participation to health care professionals in excess of the number necessary to meet the needs of the plan's enrollees (except for M+C private-fee-for-service plans, which may not refuse to contract on this basis).

(2) Use of different reimbursement amounts for different specialties or for different practitioners in the same specialty.

(3) Implementation of measures designed to maintain quality and control costs consistent with its responsibilities.

[65 FR 40324, June 29, 2000]

§ 422.206 Interference with health care professionals' advice to enrollees prohibited.

(a) General rule. (1) An M+C organization may not prohibit or otherwise restrict a health care professional, acting within the lawful scope of practice, from advising, or advocating on behalf of, an individual who is a patient and enrolled under an M+C plan about—

(i) The patient's health status, medical care, or treatment options (including any alternative treatments that may be self-administered), including the provision of sufficient information to the individual to provide an opportunity to decide among all relevant treatment options;

(ii) The risks, benefits, and consequences of treatment or non-treatment; or

(iii) The opportunity for the individual to refuse treatment and to express preferences about future treatment decisions.

(2) Health care professionals must provide information regarding treatment options in a culturally-competent manner, including the option of no treatment. Health care professionals must ensure that individuals with disabilities have effective communications with participants throughout the health system in making decisions regarding treatment options.

(b) Conscience protection. The general rule in paragraph (a) of this section does not require the M+C plan to cover, furnish, or pay for a particular counseling or referral service if the M+C organization that offers the plan—

(1) Objects to the provision of that service on moral or religious grounds; and

(2) Through appropriate written means, makes available information on these policies as follows:

(i) To HCFA, with its application for a Medicare contract, within 10 days of submitting its ACR proposal or, for policy changes, in accordance with §422.80 (concerning approval of marketing materials and election forms) and with §422.111.

(ii) To prospective enrollees, before or during enrollment.

(iii) With respect to current enrollees, the organization is eligible for the exception provided in paragraph (b)(1) of this section if it provides notice of such change within 90 days after adopting the policy at issue; however, under §422.111(d), notice of such a change must be given in advance.
§422.208 Physician incentive plans: requirements and limitations.

(a) Definitions. In this subpart, the following definitions apply:

Bonus means a payment made to a physician or physician group beyond any salary, fee-for-service payments, capitation, or returned withhold.

Capitation means a set dollar payment per patient per unit of time (usually per month) paid to a physician or physician group to cover a specified set of services and administrative costs without regard to the actual number of services provided. The services covered may include the physician’s own services, referral services, or all medical services.

Physician group means a partnership, association, corporation, individual practice association, or other group of physicians that distributes income from the practice among members. An individual practice association is defined as a physician group for this section only if it is composed of individual physicians and has no subcontracts with physician groups.

Physician incentive plan means any compensation arrangement to pay a physician or physician group that may directly or indirectly have the effect of reducing or limiting medically necessary services furnished to any particular enrollee.

Potential payments means the maximum payments possible to physicians or physician groups including payments for services they furnish directly, and additional payments based on use and costs of referral services, such as withholds, bonuses, capitation, or any other compensation to the physician or physician group. Bonuses and other compensation that are not based on use of referrals, such as quality of care furnished, patient satisfaction or committee participation, are not considered payments in the determination of substantial financial risk.

Referral services means any specialty, inpatient, outpatient, or laboratory services that a physician or physician group orders or arranges, but does not furnish directly.

Risk threshold means the maximum risk, if the risk is based on referral services, to which a physician or physician group may be exposed under a physician incentive plan without being at substantial financial risk. This is set at 25 percent risk.

Substantial financial risk, for purposes of this section, means risk for referral services that exceeds the risk threshold.

Withhold means a percentage of payments or set dollar amounts deducted from a physician’s service fee, capitation, or salary payment, and that may or may not be returned to the physician, depending on specific predetermined factors.

(b) Applicability. The requirements in this section apply to an M+C organization and any of its subcontracting arrangements that utilize a physician incentive plan in their payment arrangements with individual physicians or physician groups. Subcontracting arrangements may include an intermediate entity, which includes but is not limited to, an individual practice association that contracts with one or more physician groups or any other organized group such as those specified in §422.4.

(c) Basic requirements. Any physician incentive plan operated by an M+C organization must meet the following requirements:

(1) The M+C organization makes no specific payment, directly or indirectly, to a physician or physician group as an inducement to reduce or limit medically necessary services furnished to any particular enrollee. Indirect payments may include offerings of monetary value (such as stock options or waivers of debt) measured in the present or future.

(2) If the physician incentive plan places a physician or physician group...
at substantial financial risk (as deter-
mined under paragraph (d) of this sec-
tion) for services that the physician or
physician group does not furnish itself,
the M+C organization must assure that
all physicians and physician groups at
substantial financial risk have either
aggregate or per-patient stop-loss pro-
tection in accordance with paragraph
(f) of this section, and conduct periodic
surveys in accordance with paragraph
(h) of this section.
(3) For all physician incentive plans,
the M+C organization provides to
HCFA the information specified in
(d) Determination of substantial finan-
cial risk—(1) Basis. Substantial finan-
cial risk occurs when risk is based on
the use or costs of referral services,
and that risk exceeds the risk thresh-
hold. Payments based on other factors,
such as quality of care furnished, are
not considered in this determination.
(2) Risk threshold. The risk threshold
is 25 percent of potential payments.
(3) Arrangements that cause substantial
financial risk. The following incentive
arrangements cause substantial finan-
cial risk within the meaning of this
section, if the physician's or physician
group’s patient panel size is not great-
er than 25,000 patients, as shown in the
table at paragraph (f)(2)(iii) of this sec-
tion:
(i) Withholds greater than 25 percent
of potential payments.
(ii) Withholds less than 25 percent of
potential payments if the physician or
physician group is potentially liable
for amounts exceeding 25 percent of po-
tential payments.
(iii) Bonuses that are greater than 33
percent of potential payments minus
the bonus.
(iv) Withholds plus bonuses if the
withholds plus bonuses equal more
than 25 percent of potential payments.
The threshold bonus percentage for a
particular withhold percentage may be
calculated using the formula—With-
hold % = −0.75 (Bonus %) +25%.
(v) Capitation arrangements, if—
(A) The difference between the max-
imum potential payments and the min-
imum potential payments is more than
25 percent of the maximum potential
payments;
(B) The maximum and minimum po-
tential payments are not clearly ex-
plained in the contract with the physi-
cian or physician group.
(vi) Any other incentive arrange-
ments that have the potential to hold a
physician or physician group liable for
more than 25 percent of potential pay-
ments.
(e) Prohibition for private M+C fee-for-
service plans. An M+C fee-for-service
plan may not operate a physician in-
centive plan.
(f) Stop-loss protection requirements—
(1) Basic rule. The M+C organization
must assure that all physicians and
physician groups at substantial finan-
cial risk have either aggregate or per-
patient stop-loss protection in accord-
ance with the following requirements:
(2) Specific requirements. (i) Aggregate
stop-loss protection must cover 90 per-
cent of the costs of referral services
that exceed 25 percent of potential pay-
ments, as shown in the
panel size
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Separate institu-
Separate profes-
Payments if the physician
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1 None.
§ 422.210 Disclosure of physician incentive plans

(a) Disclosure to HCFA—(1) Basic requirement. Each M+C organization must provide to HCFA descriptive information about its physician incentive plan in sufficient detail to enable HCFA to determine whether that plan complies with the requirements of §422.208. Reporting should be on the HCFA PIP Disclosure Form (OMB No. 0938-0700).

(2) Content. The information must include at least the following:

(i) Whether services not furnished by the physician or physician group are covered by the incentive plan.

(ii) The type or types of incentive arrangements, such as, withholds, bonus, capitation.

(iii) The percent of any withhold or bonus the plan uses.

(iv) Assurance that the physicians or physician group has adequate stop-loss protection, and the amount and type of stop-loss protection.

(v) The patient panel size and, if the plan uses pooling, the pooling method.

(vi) If the M+C organization is required to conduct enrollee surveys, a summary of the survey results.

(b) Disclosure to Medicare beneficiaries—Basic requirement. An M+C organization must provide the following information to any Medicare beneficiary who requests it:

(i) Whether services not furnished by the physician or physician group are covered by the incentive plan.

(ii) The type or types of incentive arrangements, such as, withholds, bonus, capitation.

(iii) The percent of any withhold or bonus the plan uses.

(iv) Assurance that the physicians or physician group has adequate stop-loss protection, and the amount and type of stop-loss protection.

(v) The patient panel size and, if the plan uses pooling, the pooling method.

(vi) If the M+C organization is required to conduct enrollee surveys, a summary of the survey results.

(2) Be designed, implemented, and analyzed in accordance with commonly accepted principles of survey design and statistical analysis;

(3) Measure the degree of enrollees/disenrollees’ satisfaction with the quality of the services provided and the degree to which the enrollees/disenrollees have or had access to the services provided under the M+C organization; and

(4) Be conducted no later than 1 year after the effective date of the M+C organization’s contract and at least annually thereafter.

(i) Sanctions. An M+C organization that fails to comply with the requirements of this section is subject to intermediate sanctions under subpart O of this part.

§ 422.216 Special rules for M+C private fee-for-service plans.

(a) Payment to providers—(1) Payment rate. (i) The M+C organization must establish uniform payment rates for items and services that apply to all contracting providers, regardless of whether the contract is signed or deemed under paragraph (f) of this section.

(ii) Contracting providers must be reimbursed on a fee-for-service basis.

(iii) The M+C organization must make information on its payment rates available to providers that furnish services that may be covered under the M+C private fee-for-service plan.

(2) Payment to contract providers. For each service, the M+C organization pays a contract provider (including one deemed to have a contract) an amount that is equal to the payment rate under paragraph (a)(1) of this section minus any applicable cost-sharing.

(3) Noncontract providers. The organization pays for services of noncontract providers in accordance with §422.100(b)(2).

(4) Service furnished by providers of service. Any provider of services as defined in section 1861(u) of the Act that does not have in effect a contract establishing payment amounts for services furnished to a beneficiary enrolled in an M+C coordinated care plan or M+C private fee-for-service plan must accept as payment in full the amounts (less any payments under §§412.105(g) and 413.86(d) of this chapter) that it could collect if the beneficiary were enrolled in original Medicare. (Section 413.86(d) concerns calculating payment for direct graduate medical education costs.)

[63 FR 35085, June 26, 1998, as amended(2,6),(998,995)
charge enrollees no more than the cost-sharing and, subject to the limit in paragraph (b)(1)(ii) of this section, balance billing amounts that are permitted under the plan, and these amounts must be the same for "deemed" contract providers as for those that have signed contracts in effect.

(ii) The organization may permit balance billing no greater than 15 percent of the payment rate established under paragraph (a)(1) of this section.

(iii) The M+C organization must specify the amount of cost-sharing and balance billing in its contracts with providers and these amounts must be the same for "deemed" contract providers as for those that have signed contracts in effect.

(iv) The M+C organization is subject to intermediate sanctions under §422.752(a)(7), under the rules in subpart O of this part, if it fails to enforce the limit specified in paragraph (b)(1)(i) of this section.

(2) Noncontract providers. A noncontract provider may not collect from an enrollee more than the cost-sharing established by the M+C private fee-for-service plan as specified in §422.308(b), unless the provider has opted out of Medicare as described in part 405, subpart D of this chapter.

(c) Enforcement of limit—(1) Contract providers. An M+C organization that offers an M+C fee-for-service plan must enforce the limit specified in paragraph (b)(1) of this section.

(2) Noncontract providers. An M+C organization that offers an M+C private fee-for-service plan must monitor the amount collected by noncontract providers to ensure that those amounts do not exceed the amounts permitted to be collected under paragraph (b)(2) of this section, unless the provider has opted out of Medicare as described in part 405, subpart D of this chapter. The M+C organization must develop and document violations specified in instructions and must forward documented cases to HCFA.

(d) Information on enrollee liability—(1) General information. An M+C organization that offers an M+C fee-for-service plan must provide to plan enrollees, for each claim filed by the enrollee or the provider that furnished the service, an appropriate explanation of benefits. The explanation must include a clear statement of the enrollee’s liability for deductibles, coinsurance, copayment, and balance billing.

(2) Advance notice for hospital services. In its terms and conditions of payment to hospitals, the M+C organization must require the hospital, if it imposes balance billing, to provide to the enrollee, before furnishing any services for which balance billing could amount to not less than $500—

(i) Notice that balance billing is permitted for those services;

(ii) A good faith estimate of the likely amount of balance billing, based on the enrollee’s presenting condition; and

(iii) The amount of any deductible, coinsurance, and copayment that may be due in addition to the balance billing amount.

(e) Coverage determinations. The M+C organization must make coverage determinations in accordance with subpart M of this part.

(f) Rules describing deemed contract providers. Any provider furnishing health services, except for emergency services furnished in a hospital pursuant to §489.24 of this chapter, to an enrollee in an M+C private fee-for-service plan, and who has not previously entered into a contract or agreement to furnish services under the plan, is treated as having a contract in effect and is subject to the limitations of this section that apply to contract providers if the following conditions are met:

(1) The services are covered under the plan and are furnished—

(i) To an enrollee of an M+C fee-for-service plan; and

(ii) Provided by a provider including a provider of services (as defined in section 1861(u) of the Act) that does not have in effect a signed contract with the M+C organization.

(2) Before furnishing the services, the provider—

(i) Was informed of the individual’s enrollment in the plan; and

(ii) Was informed (or given a reasonable opportunity to obtain information) about the terms and conditions of payment under the plan, including the information described in §422.202(a)(1).
Health Care Financing Administration, HHS § 422.250

(3) The information was provided in a manner that was reasonably designed to effect informed agreement and met the requirements of paragraphs (g) and (h) of this section.

(g) Enrollment information. Enrollment information was provided by one of the following methods or a similar method:

(1) Presentation of an enrollment card or other document attesting to enrollment.

(2) Notice of enrollment from HCFA, a Medicare intermediary or carrier, or the M+C organization itself.

(h) Information on payment terms and conditions. Information on payment terms and conditions was made available through either of the following methods:

(1) The M+C organization used postal service, electronic mail, FAX, or telephone to communicate the information to one of the following:

(i) The provider.

(ii) The employer or billing agent of the provider.

(iii) A partnership of which the provider is a member.

(iv) Any party to which the provider makes assignment or reassigns benefits.

(2) The M+C organization has in effect a procedure under which—

(i) Any provider furnishing services to an enrollee in an M+C private fee-for-service plan, and who has not previously entered into a contract or agreement to furnish services under the plan, can receive instructions on how to request the payment information;

(ii) The organization responds to the request before the entity furnishes the service;

(iii) The information the organization provides includes the following:

(A) Billing procedures.

(B) The amount the organization will pay towards the service.

(C) The amount the provider is permitted to collect from the enrollee.

(D) The information described in §422.202(a)(1).

(3) Announcements in newspapers, journals, or magazines or on radio or television are not considered communication of the terms and conditions of payment.

(i) Provider credentialing requirements. Contracts with providers must provide that, in order to be paid to provide services to plan enrollees, providers must meet the requirements specified in §422.204(a)(1) and (a)(1)(iii).


§ 422.220 Exclusion of services furnished under a private contract.

An M+C organization may not pay, directly or indirectly, on any basis, for services (other than emergency or urgently needed services as defined in §422.2) furnished to a Medicare enrollee by a physician (as defined in section 1861(r)(1) of the Act) or other practitioner (as defined in section 1842(b)(18)(C) of the Act) who has filed with the Medicare carrier an affidavit promising to furnish Medicare-covered services to Medicare beneficiaries only through private contracts under section 1802(b) of the Act with the beneficiaries. An M+C organization must pay for emergency or urgently needed services furnished by a physician or practitioner who has not signed a private contract with the beneficiary.

Subpart F—Payments to Medicare+Choice Organizations

SOURCE: 63 FR 35090, June 26, 1998, unless otherwise noted.

§ 422.249 Terminology.

In this subpart—

(a) The terms “per capita rate” and “capitation rate” (see §422.252) are used interchangeably; and

(b) In the term “area-specific,” “area” refers to any of the payment areas described in §422.250(c).

§ 422.250 General provisions.

(a) Monthly payments—(1) General rule. Except as provided in paragraphs (a)(2) or (f) of this section, HCFA makes advance monthly payments equal to 1/12th of the annual M+C capitation rate for the payment area described in paragraph (c) of this section adjusted for such demographic risk factors as an individual’s age, disability status, sex, institutional status, and other such factors as it determines to
be appropriate to ensure actuarial equivalence. Effective January 1, 2000, HCFA adjusts for health status as provided in §422.256(c). When the new risk adjustment is implemented, ½th of the annual capitation rate for the payment area described in paragraph (c) of this section will be adjusted by the risk adjustment methodology under §422.256(d).

(2) Special rules—(i) Enrollees with end-stage renal disease. (A) For enrollees determined to have end-stage renal disease (ESRD), HCFA establishes special rates that are determined under an actuarially equivalent approach to that used in establishing the rates under original Medicare.

(B) HCFA reduces the payment rate for each renal dialysis treatment by the same amount that the Secretary is authorized to reduce the amount of each composite rate payment for each treatment as set forth in section 1881(b)(7) of the Act. These funds are to be used to help pay for the ESRD network program in the same manner as similar reductions are used in original Medicare.

(ii) MSA enrollees. For MSA enrollees, HCFA makes advanced monthly payments as described in paragraph (a)(1) less the amount (if any) identified in §422.262(c)(1)(i) to be deposited in the M+C MSA. In addition, HCFA deposits in the M+C MSA the lump sum amounts (if any) determined in accordance with §422.262(c).

(iii) RFB plan enrollees. For RFB plan enrollees, HCFA adjusts the capitation payments otherwise determined under this subpart to ensure that the payment level is appropriate for the actuarial characteristics and experience of these enrollees. Such adjustment can be made on an individual or organization basis.

(b) Adjustment of payments to reflect number of Medicare enrollees—(1) General rule. HCFA adjusts payments retroactively to take into account any difference between the actual number of Medicare enrollees and the number on which it based an advance monthly payment.

(2) Special rules for certain enrollees. (i) Subject to paragraph (b)(2)(ii) of this section, HCFA may make adjustments, for a period (not to exceed 90 days) that begins when a beneficiary elects a group health plan (as defined in §411.101 of this chapter) offered by an M+C organization, and ends when the beneficiary is enrolled in an M+C plan offered by the M+C organization.

(ii) HCFA does not make an adjustment unless the beneficiary certifies that, at the time of enrollment under the M+C plan, he or she received from the organization the disclosure statement specified in §422.111.

(c) Payment areas—(1) General rule. Except as provided in paragraph (e) of this section, the M+C payment area is a county or an equivalent geographic area specified by HCFA.

(2) Special rule for ESRD enrollees. For ESRD enrollees, the M+C payment area is a State or other geographic area specified by HCFA.

(d) Terminology. As used in paragraph (e) of this section, “metropolitan statistical area,” “consolidated metropolitan statistical area,” and “primary metropolitan statistical area” mean any areas so designated by the Secretary of Commerce.

(e) Geographic adjustment of payment areas. For contract years beginning after 1999—

(1) State request. A State’s chief executive may request, no later than February 1 of any year, a geographic adjustment of the State’s payment areas for the following calendar year. The chief executive may request any of the following adjustments to the payment area specified in paragraph (c)(1) of this section:

(i) A single Statewide M+C payment area.

(ii) A metropolitan-based system in which all nonmetropolitan areas within the State constitute a single payment area and any of the following constitutes a separate M+C payment area:

(A) All portions of each single metropolitan statistical area within the State.

(B) All portions of each primary metropolitan statistical area within each consolidated metropolitan statistical area within the State.

(iii) A consolidation of noncontiguous counties.
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(2) HCFA response. In response to the request, HCFA makes the payment adjustment requested by the chief executive.

(3) Budget neutrality adjustment for geographically adjusted payment areas. If HCFA adjusts a State's payment areas in accordance with paragraph (e)(2) of this section, HCFA at that time, and each year thereafter, adjusts the capitation rates so that the aggregate Medicare payments do not exceed the aggregate Medicare payments that would have been made to all the State's payments areas, absent the geographic adjustment.

(f) Determination and applicability of payment rates. (1) All payment rates are annual rates, determined and promulgated no later than March 1st, for the following calendar year.

(2) For purposes of paragraphs (b) and (c) of § 422.252, except as provided in § 422.254(e)(4), the "capitation payment rate for 1997" is the rate determined under section 1876(a)(1)(c) of the Act.

(g) Bonus payments. (1) HCFA provides bonus payments to the M+C organization(s) that first offers a plan in a previously unserved county on or after January 1, 2000 and no later than December 31, 2000. The bonus payment amounts equal—

(i) For the first 12 months after a plan is offered in a previously unserved county, 5 percent of the monthly capitation rate otherwise payable under this section; and

(ii) For the subsequent 12 months, 3 percent of the monthly capitation rate otherwise payable under this section.

(2) A previously unserved county is defined as—

(i) A county in which no M+C plan has been offered; or

(ii) A county in which an M+C plan or plans has been offered, but where any M+C organization offering an M+C plan notified HCFA by October 13, 1999, that it will no longer offer plans in the county as of January 1, 2000.

(3) A plan is considered to be offered when—

(i) The M+C organization sponsoring the plan has a contract in effect to serve beneficiaries in the previously unserved area; and

(ii) The M+C plan is open for enrollment.

§ 422.252 Annual capitation rates.

Subject to the adjustments specified in this subpart, the annual capitation rate for a particular payment area is equal to the largest of the following:

(a) Blended capitation rate. The blended capitation rate is the sum of—

(1) The area-specific percentage (specified in §422.254(a)) for the year multiplied by the annual area-specific capitation rate for the payment area as determined under §422.254(e) for the year, and

(2) The national percentage (specified in §422.254(a)) for the year multiplied by the national input-price-adjusted capitation rate for the payment area as determined under §422.254(g) for the year.

(3) Multiplied by the budget neutrality adjustment factor determined under §422.254(d).

(b) Minimum amount rate. (1) For 1998—

(i) For the 50 States and the District of Columbia, the minimum amount rate is 12 times $367.

(ii) For all other jurisdictions the minimum amount rate is the lesser of the rate described in (b)(1)(i) or 150 percent of the capitation payment rate for 1997.

(2) For each succeeding year, the minimum amount rate is the minimum amount rate for the preceding year, increased by the national per capita growth percentage (specified in §422.254(b)) for the year.

(c) Minimum percentage increase rate. (1) For 1998, the minimum percentage increase rate is 102 percent of the annual capitation rate for the preceding year, increased by the national per capita growth percentage (specified in §422.254(b)) for the year.

(2) For each succeeding year, the minimum percentage increase rate is 102 percent of the annual capitation rate for the preceding year.

§ 422.254 Calculation and adjustment factors.

The following are the factors used in calculating the per capita payment rates:
(a) Area-specific and national percentages. For purposes of §422.252(a)(1), the area-specific percentage and the national percentage, for each year, are as follows:

<table>
<thead>
<tr>
<th>Year</th>
<th>Area-specific</th>
<th>National</th>
</tr>
</thead>
<tbody>
<tr>
<td>1998</td>
<td>90</td>
<td>10</td>
</tr>
<tr>
<td>1999</td>
<td>82</td>
<td>18</td>
</tr>
<tr>
<td>2000</td>
<td>74</td>
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<td>2001</td>
<td>66</td>
<td>34</td>
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<tr>
<td>2002</td>
<td>58</td>
<td>42</td>
</tr>
<tr>
<td>After 2002</td>
<td>50</td>
<td>50</td>
</tr>
</tbody>
</table>

(b) National per capita growth percentage. For purposes of §422.252(a)(2),

1. The national per capita growth percentage for a year is HCFA’s estimate of the rate of growth in per capita expenditures, reduced by the percentage points specified in paragraph (b)(2) of this section for the year. HCFA may make separate estimates for aged enrollees, disabled enrollees, and enrollees who have ESRD.

2. The percentage points that HCFA uses to reduce its estimates are as follows:

   (i) For 1998, 0.8 percentage points.
   (ii) For years 1999 through 2001, 0.5 percentage points.
   (iii) For 2002, 0.3 percentage points.
   (iv) For years after 2002, 0 percentage points.

3. Medical education payment adjustments. For purposes of paragraph (e)(2) the medical education payment adjustments are amounts that HCFA estimates were payable to teaching hospitals during 1997 for—

   (1) the indirect costs of medical education under section 1886(d)(5)(B) of the Act; and
   (2) the direct costs of graduate medical education under section 1886(h) of the Act.

4. General budget neutrality factor. For each year, HCFA applies a budget neutrality factor to the blended capitation rates under §422.252(a) so that the estimated aggregate payments made under this part equal the estimated aggregate payments that would have been made if based entirely on area-specific capitation rates.

(e) Annual area-specific capitation rate

(1) Basic rule. Subject to the provisions of paragraphs (e)(2) and (e)(3) of this section, the annual area-specific capitation rate for a particular payment area is—

   (i) For 1998, subject to paragraph (e)(4) of this section, the per capita rate determined for that area for 1997 under section 1876(a)(1)(c) of the Act, increased by the national per capita growth percentage for 1998; and
   (ii) For a subsequent year, the area-specific capitation rate determined for the previous year, increased by the national per capita growth percentage for the year.

(2) Exclusion of medical education costs. In calculating the area-specific capitation rates, the following percentages of the amounts estimated by HCFA under §422.254(c) as medical education payment adjustments to hospitals, are excluded:

   (i) For 1998, 20 percent.
   (ii) For 1999, 40 percent.
   (iii) For 2000, 60 percent.
   (iv) For years after 2000, 100 percent.

(f) National standardized annual capitation rate. The national standardized annual capitation rate is equal to—

   (1) The sum, for all payment areas, of the products of—
      (i) The area-specific capitation rate and
      (ii) The average number of Medicare beneficiaries residing in the area multiplied by the average of the risk-factor weights used to adjust payments under §422.256(c); and
   (2) Divided by the sum, for all payment areas, of the products specified in
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paragraph (f)(1)(ii) of this section for all payment areas.

(g) The input-price-adjusted annual national capitation rate—(1) General rule. The input-price-adjusted annual national capitation rate for a M+C payment area for a year is equal to the sum, for all the types of Medicare services (as classified by HCFA), of the product (for each service) of—

(i) The national standardized annual M+C capitation rate (determined under paragraph (f) of this section) for the year;

(ii) The proportion of such rates for the year which is attributable to such type of services; and

(iii) An index that reflects (for that year and that type of services) the relative input price of such services in the area compared to the national average input price for such services.

(2) HCFA may, subject to the special rules for 1988, use indices that are used in applying or updating national payment rates for particular areas and localities.

(3) Special rules for 1988. In applying this paragraph for 1998—

(i) Medicare services are classified as Part A and Part B services;

(ii) The proportion attributable to Part A services is the ratio (expressed as a percentage) of the national average per capita rate of payment for Part A services for 1997 to the national average per capita rate of payment for Part A and Part B services for that year;

(iii) The proportion attributed to part B services is 100 percent minus the ratio described in paragraph (g)(3)(i) of this section;

(iv) For Part A services, 70 percent of the payments attributable to those services are adjusted by the index used under section 1886(d)(3)(E) of the Act to adjust payment rates for relative hospital wage levels for hospitals located in the particular payment area;

(v) For Part B services—

(A) 66 percent of payments attributable to those services are adjusted by the index of the geographic area factors under section 1848(e) of the Act used to adjust payment rates for physician services in the particular payment area; and

(B) Of the remaining 34 percent, 40 percent is adjusted by the index specified in paragraph (g)(3)(iv) of this section.

[63 FR 35090, June 26, 1998, as amended at 65 FR 40326, June 29, 2000]

§ 422.256 Adjustments to capitation rates and aggregate payments.

(a) Adjustment for over or under projection of national per capita growth percentages. (1) Beginning with rates for 1999, HCFA adjusts all area-specific and national capitation rates for the previous year to reflect any differences between the projected national per capita growth percentages for that year and previous years, and the current estimates of those percentages for such years.

(2) Beginning with rates for 2000, HCFA also adjusts the minimum amount rate (calculated under § 422.252(b)) in the same manner.

(b) Adjustment for national coverage determination (NCD) services. If HCFA determines that the cost of furnishing an NCD service is “significant,” HCFA adjusts capitation rates for the next calendar year to take account of the cost of that service. Until the new capitation rates are in effect, the M+C organization is paid for the “significant cost” service on a fee-for-service basis as provided under section 422.105(b).

(c) Risk adjustment: General rule. Capitation payments are adjusted for age, gender, institutional status, and other appropriate factors, including health status.

(d) Risk adjustment: Health status—(1) Data collection. To adjust for health status, HCFA applies a risk factor based on data obtained in accordance with § 422.257.

(2) Initial implementation. HCFA applies this adjustment factor to payments beginning January 1, 2000.

(3) Uniform application. Except as provided for M+C RFB plans under § 422.250(a)(2)(iii), HCFA applies this adjustment factor to all types of plans.

§ 422.257 Encounter data.

(a) Data collection: Basic rule. Each M+C organization must submit to HCFA (in accordance with HCFA instructions) all data necessary to characterize the context and purposes of
each encounter between a Medicare enrollee and a provider, supplier, physician, or other practitioner.

(b) Types of service and timing of submittal. M+C organizations must submit data as follows:

(1) Beginning on a date determined by HCFA, inpatient hospital care data for all discharges that occur on or after July 1, 1997.

(2) HCFA will provide advance notice to M+C organizations to collect and submit data for services that occur on or after July 1, 1999, as follows:
   (i) Physician, outpatient hospital, SNF, and HHA data beginning no earlier than October 1, 1999; and
   (ii) All other data HCFA deems necessary beginning no earlier than October 1, 2000.

(c) Sources and extent of data. (1) To the extent required by HCFA, the data must account for services covered under the original Medicare program, for Medicare covered services for which Medicare is not the primary payor, or for other additional or supplemental benefits that the M+C organization may provide.

(2) The data must account separately for each provider, supplier, physician, or other practitioner that would be permitted to bill separately under the Medicare fee-for-service program, even if they participate jointly in the same encounter.

(d) Other data requirements. (1) M+C organizations must submit data that conform to the requirements for equivalent data for Medicare fee-for-service when appropriate, and to all relevant national standards.

(2) The data must be submitted electronically to the appropriate HCFA contractor.

(3) M+C organizations must obtain the encounter data required by HCFA from the provider, supplier, physician, or other practitioner that rendered the services.

(4) M+C organizations may include in their contracts with providers, suppliers, physicians, and other practitioners, provisions that require submission of complete and accurate encounter data as required by HCFA. These provisions may include financial penalties for failure to submit complete data, or for failure to submit data that conform to the requirements for equivalent data for Medicare fee-for-service.

(e) Validation of data. M+C organizations and their providers and practitioners will be required to submit medical records for the validation of encounter data, as prescribed by HCFA.

(f) Use of data. HCFA uses the data obtained under this section to determine the risk adjustment factor that it applies to annual capitation rates under §422.256(c). HCFA may also use the data for other purposes.

(g) Deadlines for submission of encounter data. Risk adjustment factors for each payment year are based on encounter data submitted for services furnished during the 12 month period ending 6 months before to the payment year (for example, risk adjustment factors for CY 2000 are based on data for services furnished during the period July 1, 1998 through June 30, 1999).

   (1) The annual deadline for encounter data submission is September 10 for encounter data reflecting services furnished during the 12 month period ending the prior June 30 (for example, the deadline for submission of data for the period July 1, 1998 through June 30, 1999 is September 10, 1999).

   (2) HCFA allows a reconciliation process to account for late data submissions. HCFA continues to accept encounter data submitted after the September 10 deadline until June 30 of the payment year (for example, until June 30, 2000 for data from the period July 1, 1998 through June 30, 1999). After the payment year is completed, HCFA recalculates the risk factors for affected individuals to determine if adjustments to payments are necessary.

[63 FR 35090, June 26, 1998, as amended at 65 FR 40326, June 29, 2000]

§ 422.258 Announcement of annual capitation rates and methodology changes.

(a) Capitation rates. (1) No later than March 1 of each year, HCFA announces to M+C organizations and other interested parties the capitation rates for the following calendar year.

(2) HCFA includes in the announcement a description of the risk and other factors and explains the methodology in sufficient detail to enable...
M+C organizations to compute monthly adjusted capitation rates for individuals in each of its payment areas.

(b) Advance notice of changes in methodology. (1) No later than January 15 of each year, HCFA notifies M+C organizations of changes it proposes to make in the factors and the methodology it used in the previous determination of capitation rates.

(2) The M+C organizations have 15 days to comment on the proposed changes.

§ 422.262 Special rules for beneficiaries enrolled in M+C MSA plans.

(a) Establishment and designation of medical savings account (MSA). A beneficiary who elects coverage under an M+C MSA plan—

(1) Must establish an M+C MSA with a trustee that meets the requirements of paragraph (b) of this section; and

(2) If he or she has more than one M+C MSA, designate the particular account to which payments under the M+C MSA plan are to be made.

(b) Requirements for MSA trustees. An entity that acts as a trustee for an M+C MSA must—

(1) Register with HCFA;

(2) Certify that it is a licensed bank, insurance company, or other entity qualified, under sections 408(a)(2) or 408(h) of the IRS Code, to act as a trustee of individual retirement accounts;

(3) Agree to comply with the M+C MSA provisions of section 138 of the IRS Code of 1986; and

(4) Provide any other information that HCFA may require.

(c) Deposit in the M+C MSA. (1) The payment is calculated as follows:—

(i) The monthly M+C MSA premium is compared with 1/12 of the annual capitation rate for the area determined under §422.252.

(ii) If the monthly M+C MSA premium is less than 1/12 of the annual capitation rate, the difference is the amount to be deposited in the M+C MSA for each month for which the beneficiary is enrolled in the MSA plan.

(2) HCFA deposits the full amount to which a beneficiary is entitled under paragraphs (c)(i)(ii) of this section for the calendar year, beginning with the month in which M+C MSA coverage begins.

(3) If the beneficiary’s coverage under the M+C MSA plan ends before the end of the calendar year, HCFA recovers the amount that corresponds to the remaining months of that year.

§ 422.264 Special rules for coverage that begins or ends during an inpatient hospital stay.

(a) Applicability. This section applies to inpatient services in a “subsection (d) hospital” as defined in section 1886(d)(1)(B) of the Act.

(b) Coverage that begins during an inpatient hospital stay. If coverage under an M+C plan offered by an M+C organization begins while the beneficiary is an inpatient in a subsection (d) hospital—

(1) Payment for inpatient services until the date of the beneficiary’s discharge is made by the previous M+C organization or original Medicare, as appropriate.

(2) The M+C organization offering the newly-elected M+C plan is not responsible for the inpatient services until the date after the beneficiary’s discharge; and

(3) The M+C organization offering the newly-elected M+C plan is paid the full amount otherwise payable under this subpart.

(c) Coverage that ends during an inpatient hospital stay. If coverage under an M+C plan offered by an M+C organization ends while the beneficiary is an inpatient in a subsection (d) hospital—

(1) The M+C organization is responsible for the inpatient services until the date of the beneficiary’s discharge;

(2) Payment for those services during the remainder of the stay is not made by original Medicare or by any succeeding M+C organization offering a newly-elected M+C plan; and

(3) The M+C organization that no longer provides coverage receives no payment for the beneficiary for the period after coverage ends.

§ 422.266 Special rules for hospice care.

(a) Information. An M+C organization that has a contract under subpart K of this part must inform each Medicare enrollee eligible to elect hospice care
§ 422.268 Source of payment and effect of election of the M+C plan election on payment.

(a) Source of payments. Payments under this subpart, to M+C organizations or M+C MSAs, are made from the Federal Hospital Insurance Trust Fund or the Supplementary Medical Insurance Trust Fund. HCFA determines the proportions to reflect the relative weight that benefits under Part A, and benefits under Part B represents of the actuarial value of the total benefits under title XVIII of the Act.

(b) Payments to the M+C organization. Subject to §§422.109, 422.264, and 422.266, HCFA’s payments under a contract with an M+C organization (described in §422.250) with respect to an individual electing an M+C plan offered by the organization are instead of the amounts which (in the absence of the contract) would otherwise be payable under original Medicare for items and services furnished to the individual.

(c) Only the M+C organization entitled to payment. Subject to §§422.262, 422.264, 422.266, and 422.520 of this part and sections 1886(d)(11) and 1886(h)(3)(D) of the Act, only the M+C organization is entitled to receive payment from HCFA under title XVIII of the Act for items and services furnished to the individual.


Subpart G—Premiums and Cost-Sharing

SOURCE: 63 FR 35093, June 26, 1998, unless otherwise noted.

§ 422.300 Basis and scope.

(a) General. This subpart is based on section 1854 of the Act. It sets forth the requirements and limitations for payments by and on behalf of Medicare beneficiaries who elect an M+C plan.

(b) Transition period. For contract periods beginning before January 1, 2002, HCFA applies the following special rules.

(1) M+C organizations may, with HCFA’s agreement, modify an M+C plan offered prior to January 1, 2002 by—

(i) Adding benefits at no additional cost to the M+C plan enrollee; and

(ii) Lowering the premiums approved through the ACR process;

(iii) Lowering other cost-sharing amounts approved through the ACR process.

(2) For contracts beginning on a date other than January 1, 2002, M+C organizations may submit ACRs on a date other than July 1 approved by HCFA.

[63 FR 35093, June 26, 1998, as amended at 65 FR 40326, June 29, 2000]

§ 422.302 Terminology.

As used in this subpart, unless specified otherwise—

Additional revenues are revenues collected or expected to be collected from charges for M+C plans offered by an
M+C organization in excess of costs actually incurred or expected to be incurred. Additional revenues would include such things as revenues in excess of expenses of an M+C plan, profits, contribution to surplus, risk margins, contributions to risk reserves, assessments by a related entity that do not represent a direct medical or related administrative cost, and any other premium component not reflected in direct medical care costs and administrative costs.

APR stands for the M+C plan’s average per capita rates of payment. The APR is the average amount the M+C organization estimates HCFA will pay (without any needed offsets or reductions, such as, those required by §422.250(a)(2)(ii) for M+C MSA plan enrollees) for the period covered by the ACR for all of the Medicare beneficiaries electing the M+C plan.

M+C monthly basic beneficiary premium means, with respect to an M+C coordinated care plan, the amount authorized to be charged under §422.308(a)(1) for the plan, or, with respect to a M+C private fee-for-service plan, the amount filed under §422.306(d)(1).

M+C monthly supplemental beneficiary premium means, with respect to an M+C coordinated care plan, the amount authorized to be charged under §422.308(a)(2) for the M+C plan, or, with respect to an MSA or an M+C private fee-for-service plan, the amount filed under §422.306(c)(2) or §422.306(d)(2).

M+C monthly MSA premium means, with respect to an M+C plan, the amount of such premium filed under §422.306(c)(1).

§ 422.304 Rules governing premiums and cost-sharing.

(a) Monthly premiums. The monthly premium charged to the beneficiary is—

(1) For an individual enrolled in an M+C plan (other than an M+C MSA plan) offered by an M+C organization, the sum of the M+C monthly basic beneficiary premium plus the M+C monthly supplemental beneficiary premium (if any); or

(2) For an individual enrolled in an M+C MSA plan offered by an M+C organization, the M+C monthly supplemental beneficiary premium (if any).

(b) Uniformity.—(1) General rule. The M+C monthly basic beneficiary premium, the M+C monthly supplemental beneficiary premiums, and the M+C monthly MSA premium of an M+C organization may not vary among individuals enrolled in an M+C plan (or segment of the plan as provided under paragraph (b)(2) of this section). In addition, the M+C organization may not vary the level of cost-sharing charged for basic benefits or supplemental benefits (if any), among individuals enrolled in an M+C plan (or segment of the plan as provided under paragraph (b)(2) of this section).

(2) Segmented service area option. An M+C organization may apply the uniformity requirements in paragraph (b)(1) of this section to segments of an M+C plan service area (rather than to the entire service area) as long as any such segment is composed of one or more M+C payment areas, and the information specified under §422.306 is submitted separately, as provided in that section, for each such segment.

(c) Timing of payments. The M+C organization must permit payments of M+C monthly basic and supplemental beneficiary premium on a monthly basis and may not terminate coverage for failure to make timely payments except as provided in §422.74(b)(1).

(d) Monetary inducements prohibited. An M+C organization may not provide for cash or other monetary rebates as an inducement for enrollment or for any other reason or purpose.

§ 422.306 Submission of proposed premiums and related information.

(a) General rule. (1) Not later than July 1 of each year, each M+C organization and any organization intending to contract as an M+C organization in the subsequent year must submit to HCFA, in the manner and form prescribed by HCFA, for each M+C plan (or service area segment, under §422.304(b)(2)) it intends to offer in the following year—

(i) The information specified in paragraph (b), (c), or paragraph (d) of this section for the type of M+C plan involved; and
§ 422.308 Limits on premiums and cost sharing amounts.

(ii) The service area and enrollment capacity (if any).

(2) If the submission is not complete, timely, or accurate, HCFA has the authority to impose sanctions under subpart O of this part or may choose not to renew the contract.

(b) Information required for coordinated care plans—(1) Basic benefits. For basic benefits, the following information is required:

(i) The ACR as specified in §422.310.

(ii) The M+C monthly basic beneficiary premium.

(iii) A description of cost-sharing to be imposed under the plan, and its actuarial value.

(iv) A description of any additional benefits to be provided pursuant to §422.312 and the actuarial value determined for those benefits.

(v) Amounts collected in the previous contract period for basic benefits.

(2) Supplemental benefits. For supplemental benefits, the following information is required:

(i) The ACR.

(ii) The M+C monthly supplemental beneficiary premium.

(iii) A description of supplemental benefits being offered, the cost sharing to be imposed, and their actuarial value.

(iv) Amounts collected in the previous contract period for supplemental benefits.

(c) Information required for MSA plans.

(1) The monthly MSA premium for basic benefits.

(2) The M+C monthly supplementary beneficiary premium for supplemental benefits.

(3) A description of all benefits offered under the M+C MSA plan.

(4) The amount of the deductible imposed under the plan.

(5) Amounts collected in the previous contract period for supplemental benefits.

(d) Information required for M+C private fee-for-service plans.

(1) The information specified under paragraph (b)(1) of this section.

(2) The amount of M+C supplemental beneficiary premium.

(3) A description of all benefits offered under the plan.

(4) Amounts collected in the previous contract period for basic and supplemental benefits.

(e) HCFA review—(1) Basic rule. Except as specified in paragraph (e)(2) of this section, HCFA reviews and approves or disapproves the information submitted under this section.

(2) Exception. HCFA does not review or approve or disapprove the following information:

(i) Any amounts submitted with respect to M+C MSA plans.

(ii) The M+C monthly basic and supplemental beneficiary premiums for M+C private fee-for-service plans.

[63 FR 35093, June 26, 1998, as amended at 65 FR 40326, June 29, 2000]
§ 422.309 Incorrect collections of premiums and cost-sharing.

(a) Definitions. As used in this section—
   (1) Amounts incorrectly collected means amounts that:
      (i) Exceed the limits imposed by §422.308;
      (ii) Include amounts collected from an enrollee who was believed not entitled to Medicare benefits but was later found to be entitled.
   (2) Other amounts due are amounts due for services that were—
      (i) Emergency, urgently needed services, or other services obtained outside the M+C plan; or
      (ii) Initially denied but, upon appeal, found to be services the enrollee was entitled to have furnished by the M+C organization.

(b) Rule for M+C private fee-for-service plans. (1) The average actuarial value of the cost-sharing for basic benefits may not exceed the amounts approved in the ACR for those benefits, as determined under §422.310.
   (2) For supplemental benefits, the actuarial value of its cost-sharing may not exceed the amounts approved in the ACR for those benefits, as determined under §422.310 on an annual basis.

(c) Special rules for determination of actuarial value. If HCFA determines that adequate data are not available to determine actuarial value under paragraph (a) or (b) of this section, HCFA may make the determination with respect to all M+C eligible beneficiaries in the same geographic area or State or in the United States, or on the basis of other appropriate data.

§ 422.309 Incorrect collections of premiums and cost-sharing.

(3) Coverage of Part A services for Part B-only Medicare enrollees. If an M+C organization furnishes coverage of Medicare Part A-type services to a Medicare enrollee entitled to Part B only, the M+C plan's premium plus the actuarial value of its cost-sharing for these services may not exceed the lesser of—
   (i) The APR that is payable for these services for those beneficiaries entitled to Part A plus the actuarial value of Medicare deductibles and coinsurance for the services:
      (ii) or the ACR for such services.

(b) Rule for M+C private fee-for-service plans. (1) The average actuarial value of the cost-sharing for basic benefits may not exceed the actuarial value of the cost-sharing that would apply, on average, to beneficiaries entitled to Medicare Part A and enrolled in Medicare Part B if they were not enrolled in an M+C plan as determined in the ACR under §422.310.
   (2) For supplemental benefits, the actuarial value of its cost-sharing may not exceed the amounts approved in the ACR for those benefits, as determined under §422.310 on an annual basis.

(c) Special rules for determination of actuarial value. If HCFA determines that adequate data are not available to determine actuarial value under paragraph (a) or (b) of this section, HCFA may make the determination with respect to all M+C eligible individuals in the same geographic area or State or in the United States, or on the basis of other appropriate data.

§ 422.310 Adjusted community rate (ACR) approval process.

(a) General rule.

(1) Except with respect to M+C MSA plans, each M+C organization must compute a separate ACR for each M+C coordinated care or private fee-for-service plan offered to Medicare beneficiaries. In computing the ACR, the M+C organization calculates an initial rate (for years after 1999, using the methods described in paragraph (b), for 1999, under § 417.594(b)) that represents the “commercial premium” the M+C organization would charge its general non-Medicare eligible enrollment population for the basic benefits, and any mandatory supplemental benefits covered under the M+C plan. The M+C organization should also calculate a separate initial rate (using the same approach) for each optional supplemental benefit package it offers under an M+C plan. For years after 1999 the M+C organization then either adjusts that rate by the factors specified in paragraph (c) of this section or requests that HCFA adjust the rate in accordance with the procedures specified in paragraph (c)(6) of this section. For 1999, adjustments are made under section 417.594(c). All data submitted as part of the ACR process is subject to audit by HCFA or any person or organization designated by HCFA.

(2) To calculate the adjusted excess described in section 422.312, the M+C organization or HCFA further reduces the rate for Medicare-covered services by the actuarial value of applicable Medicare coinsurance and deductibles.

(3) Separate ACRs must be calculated for Part A and Part B enrollees and Part B-only enrollees for each M+C plan offered, and for each optional supplemental benefit option.

(4) In calculating its initial rate, the M+C organization must identify and take into account anticipated revenue collectible from other payers for those services for which Medicare is not the primary payer as described in § 422.108.

(b) Initial rate calculation for years after 1999.

(1) The M+C organization’s initial rate for each M+C plan is calculated on a 12-month basis for non-Medicare enrollees, using either, at the M+C organization’s election—

(i) A community rating system (as defined in section 1308(8) of the PHS Act, other than subparagraph (C)); or

(ii) A system, approved by HCFA, under which the M+C organization develops an aggregate premium for each M+C plan that is weighted by the size of the various enrolled groups and individuals that compose the M+C organization’s enrollment in that M+C plan. For purposes of this section, enrolled groups are defined as employee groups or other bodies of subscribers (including individual subscribers) that enroll in the M+C plan on a premium basis.

(2) Regardless of which method the M+C organization uses to calculate its initial rate, the initial rate must be equal to the premium the M+C organization would charge its non-Medicare enrollees on a yearly basis for services included in the M+C plan.

(3) Except as provided in paragraph (b)(4) of this section, the M+C organization must identify in its initial rate calculation for an M+C plan, the following components whose rates must be consistent with rates used by the M+C organization in calculating premiums for non-Medicare enrollees:

(i) Direct medical care.

(ii) Administration.

(iii) Additional Revenues.

(iv) Enrollee cost sharing (for example, deductibles, coinsurance, or copayments) for Medicare-covered services and for additional and supplemental benefits.
(4) An M+C organization that does not usually separate its premium components as described in paragraph (b)(3) of this section may calculate its initial rate with the methods it uses for its other enrolled groups if the M+C organization provides HCFA with the documentation necessary to support any adjustments the M+C organization makes to the initial rate in accordance with paragraph (c)(5) of this section.

(5) The initial rate calculation must not carry forward any losses experienced by the M+C organization during prior contract periods. The M+C organization must submit supporting documentation to assure HCFA that ACR values do not include past losses but only premiums for covered services, additional services, and supplemental benefits for the upcoming 12-month period.

(c) Adjustment factors for years after 1999. Adjustment factors are designed to adjust on a component basis the initial rate calculated under paragraph (b) of this section to reflect differences in utilization characteristics of the M+C organization's Medicare enrollees electing an M+C plan using a relative cost ratio. Adjustment factors are as follows:

(1) Direct medical care. The relative cost ratio for direct medical care for an M+C plan is determined by comparing the direct medical care costs actually incurred on an accrual basis during the most recently ended calendar year prior to submission of the ACR for Medicare enrollees that elected the M+C plan to the direct medical care costs of non-Medicare enrollees incurred over the same period. The non-Medicare enrollees included in this computation must be consistent with the non-Medicare enrollees included in the initial rate computation. When the relative cost ratio for total revenues is applied to the total initial rate, the value of additional revenues is the remaining value after removing the value of direct medical costs (as adjusted by paragraph (c)(1) of this section) and the value of Administration (as adjusted by paragraph (c)(2) of this section).

(2) Administration. The relative cost ratio for Administration for an M+C plan is determined by comparing the administrative costs actually incurred on an accrual basis during the most recently ended calendar year prior to submission of the ACR for Medicare enrollees that elected the M+C plan to the administrative costs of non-Medicare enrollees incurred over the same period. The non-Medicare enrollees included in this computation must be consistent with the non-Medicare enrollees included in the initial rate computation.

(3) Additional revenues. The relative cost ratio for total revenues for an M+C plan is determined by comparing the total revenues charged on an accrual basis during the most recently ended calendar year prior to submission of the ACR for Medicare enrollees (including payments from HCFA without any needed offsets or reductions, such as, those required by §422.250(a)(2)(i)(B) for ESRD enrollees) that elected the M+C plan to the total revenues charged for non-Medicare enrollees over the same period. The non-Medicare enrollees included in this computation must be consistent with the non-Medicare enrollees included in the initial rate computation. When the relative cost ratio for total revenues is applied to the total initial rate, the value of additional revenues is the remaining value after removing the value of direct medical costs (as adjusted by paragraph (c)(1) of this section) and the value of Administration (as adjusted by paragraph (c)(2) of this section).

(4) Additional adjustments. Additional adjustments may be necessary if the M+C organization, with agreement of HCFA, determines that the adjustment of the initial rate by the relative cost ratios does not represent an accurate ACR value of the initial rate component. In addition, adjustments will be allowed that are designed to reduce ACR values to equal the actuarial value of the M+C plan charge structure.

(5) Supporting documentation. All adjustments made by the M+C organization must be accompanied by adequate supporting data. If an M+C organization does not have sufficient enrollment experience to develop this data, it may, during its initial contract period use reasonable estimates acceptable to HCFA to establish its ACR values.

(6) Adjustment by HCFA. If it is determined that the M+C organization does not have adequate data to adjust the initial rate calculated under paragraph (b) of this section to reflect the utilization characteristics of Medicare enrollees, HCFA adjusts the initial rate.
§ 422.312 Requirement for additional benefits.

(a) Definitions. As used in this section—

(1) Excess amount is the amount by which the APR exceeds the actuarial value of the Medicare covered services required under §422.101(a), as determined on the basis of the ACR determined under §422.310, as reduced for the actuarial value of the cost-sharing under Medicare Parts A and B. A separate excess amount must be determined for Part B-only enrollees.

(2) Adjusted excess amount is the excess amount minus any amount withheld and reserved for the organization in a stabilization fund, as provided in paragraph (c) of this section.

(b) Requirement for additional benefits. If there is an adjusted excess amount for the plan it offers, the M+C organization must—

(1) Provide additional benefits with an actuarial value (less the actuarial value of any cost-sharing associated
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with the benefit) which HCFA determines is at least equal to the adjusted excess amount; and

(2) Provide those benefits uniformly for all Medicare enrollees electing the plan.

(c) Stabilization fund. (1) An M+C organization may request for part of an excess amount to be withheld and reserved, for a specified number of contract periods, in the Federal Hospital Insurance Trust Fund, or the Federal Supplementary Insurance Trust Fund in the proportions that HCFA determines to be appropriate.

(2) The reserved funds are to be used to stabilize and prevent undue fluctuations in the additional benefits that are required under this section and are provided during subsequent contract periods.

(3) Any amounts not provided as additional benefits during the period specified by the M+C organization for which the stabilization fund is established, reverts for the use of the trust funds.

(4) Establishment of a stabilization fund. An M+C organization’s request to have monies withheld in a stabilization fund for a specific M+C plan must be made when the M+C organization notifies HCFA under §422.306 of its proposed premiums, cost-sharing amounts, and related information in preparation for its next contract period.

(i) Limit per contract period. Except as provided in paragraph (c)(4)(iii) of this section, HCFA does not withhold in a stabilization fund more than 15 percent of the excess amount for a given contract period.

(ii) Cumulative limit. If HCFA has established a stabilization fund for an M+C plan, it does not approve a request for withholding made by that M+C organization for a subsequent contract period that would cause the total value of the stabilization fund to exceed 25 percent of the excess amount applicable to the M+C plan for that subsequent contract period.

(iii) Exception. HCFA may grant an exception to the limit described in paragraph (c)(3)(i) of this section if the M+C organization can demonstrate to HCFA’s satisfaction that the value of the additional benefits it provides to its Medicare enrollees electing this M+C plan fluctuates substantially in excess of 15 percent from one contract period to another.

(iv) Interest. The amounts withheld in a stabilization fund are accounted for by HCFA in accounts for which interest does not accrue to the M+C organization.

(5) Withdrawal from a stabilization fund. An M+C organization’s request to make a withdrawal from the stabilization fund established for an M+C plan must be made when the M+C organization notifies HCFA under §422.306 of its proposed premiums, cost-sharing amounts, and related information in preparation for its next contract period.

(i) Notification requirements. An M+C organization must—

(A) Indicate how it intends to use the withdrawn amounts;

(B) Justify the need for the withdrawal in terms of stabilizing the additional benefits it provides to Medicare enrollees;

(C) Document the M+C plan’s experience with fluctuations of revenue requirements relative to the additional benefits it provides to Medicare enrollees; and

(D) Document its experience during the contract period previous to the one for which it requests withdrawal to ensure that the M+C organization will not be using the withdrawn amounts to refinance losses suffered during that previous contract period.

(ii) Criteria for HCFA approval. HCFA approves a request for a withdrawal from a benefit stabilization fund for use during the next contract period only if—

(A) The average of the APR for the M+C plan’s next contract period of the M+C plan is less than that of the previous contract period;

(B) The M+C plan’s ACR for the next contract period is significantly higher than that of the previous contract period;

(C) The M+C plan’s revenue requirements for the next contract period for providing the additional benefits it provides during the previous contract period is significantly higher than the...
§ 422.350 Basis, scope, and definitions.

(a) Basis and scope. This subpart is based on sections 1851 and 1855 of the Act which, in part—

(1) Authorize provider sponsored organizations, (PSOs), to contract as a M+C plan;

(2) Require that a PSO meet certain qualifying requirements; and

(3) Provide for waiver of State licensure for PSOs under specified conditions.

(b) Definitions. As used in this subpart (unless otherwise specified)—

Capitation payment means a fixed per enrollee per month amount paid for contracted services without regard to the type, cost, or frequency of services furnished.

Cash equivalent means those assets excluding accounts receivable that can be exchanged on an equivalent basis as cash, or converted into cash within 90 days from their presentation for exchange.

Control means that an individual, group of individuals, or entity has the power, directly or indirectly, to direct or influence significantly the actions or policies of an organization or institution.

Current ratio means total current assets divided by total current liabilities.

Deferred acquisition costs are those costs incurred in starting or purchasing a business. These costs are capitalized as intangible assets and carried on the balance sheet as deferred charges since they benefit the business for periods after the period in which the costs were incurred.

Engaged in the delivery of health care services means—

(1) For an individual, that the individual directly furnishes health care services, or

(2) For an entity, that the entity is organized and operated primarily for the purpose of furnishing health care services directly or through its provider members or entities.

Generally accepted accounting principles (GAAP) means broad rules adopted by the accounting profession as guides in measuring, recording, and reporting the financial affairs and activities of a business to its owners, creditors and other interested parties.

Guarantor means an entity that—

(1) Has been approved by HCFA as meeting the requirements to be a guarantor; and

(2) Obligates its resources to a PSO to enable the PSO to meet the solvency requirements required to contract with HCFA as a M+C organization.

Health care delivery assets (HCDAs) means any tangible assets that are part of a PSO’s operation, including hospitals and other medical facilities and their ancillary equipment, and such property as may be reasonably required for the PSO’s principal office or for such other purposes as the PSO may need for transacting its business.

Insolvency means a condition in which the liabilities of the debtor exceed the fair valuation of its assets.
Net worth means the excess of total assets over total liabilities, excluding fully subordinated debt or subordinated liabilities.

Provider-sponsored organization (PSO) means a public or private entity that—
(1) Is established or organized, and operated, by a provider or group of affiliated providers;
(2) Provides a substantial proportion (as defined in §422.352) of the health care services under the M+C contract directly through the provider or affiliated group of providers; and
(3) When it is a group, is composed of affiliated providers who—
   (i) Share, directly or indirectly, substantial financial risk, as determined under §422.356, for the provision of services that are the obligation of the PSO under the M+C contract; and
   (ii) Have at least a majority financial interest in the PSO.

Qualified actuary means a member in good standing of the American Academy of Actuaries or a person recognized by the Academy as qualified for membership, or a person who has otherwise demonstrated competency in the field of actuarial determination and is satisfactory to HCFA.

Statutory accounting practices means those accounting principles or practices prescribed or permitted by the domiciliary State insurance department in the State that PSO operates.

Subordinated debt means an obligation that is owed by an organization, that the creditor of the obligation, by law, agreement, or otherwise, has a lower repayment rank in the hierarchy of creditors than another creditor. The creditor would be entitled to repayment only after all higher ranking creditors' claims have been satisfied. A debt is fully subordinated if it has a lower repayment rank than all other classes of creditors.

Subordinated liability means claims liabilities otherwise due to providers that are retained by the PSO to meet net worth requirements and are fully subordinated to all other creditors.

Uncovered expenditures means those expenditures for health care services that are the obligation of an organization, for which an enrollee may also be liable in the event of the organization's insolvency and for which no alternative arrangements have been made that are acceptable to HCFA. They include expenditures for health care services for which the organization is at risk, such as out-of-area services, referral services and hospital services. However, they do not include expenditures for services when a provider has agreed not to bill the enrollee.

§ 422.352 Basic requirements.

(a) General rule. An organization is considered a PSO for purposes of a M+C contract if the organization—
   (1) Has obtained a waiver of State licensure as provided for under §422.370; and
   (2) Meets the definition of a PSO set forth in §422.350 and other applicable requirements of this subpart; and
(3) Is effectively controlled by the provider or, in the case of a group, by one or more of the affiliated providers that established and operate the PSO.

(b) Provision of services. A PSO must demonstrate to HCFA's satisfaction that it is capable of delivering to Medicare enrollees the range of services required under a contract with HCFA. Each PSO must deliver a substantial proportion of those services directly through the provider or the affiliated providers responsible for operating the PSO. Substantial proportion means—
   (1) For a non-rural PSO, not less than 70% of Medicare services covered under the contract.
   (2) For a rural PSO, not less than 60% of Medicare services covered under the contract.

(c) Rural PSO. To qualify as a rural PSO, a PSO must—
   (1) Demonstrate to HCFA that—
      (i) It has available in the rural area, as defined in §412.62(f) of this chapter, routine services including but not limited to primary care, routine specialty care, and emergency services; and
      (ii) The level of use of providers outside the rural area is consistent with general referral patterns for the area; and
(2) Enroll Medicare beneficiaries, the majority of which reside in the rural area the PSO serves.


§ 422.354 Requirements for affiliated providers.

A PSO that consists of two or more providers must demonstrate to HCFA'S satisfaction that it meets the following requirements:

(a) The providers are affiliated. For purposes of this subpart, providers are affiliated if, through contract, ownership, or otherwise—

(1) One provider, directly or indirectly, controls, is controlled by, or is under common control with another;

(2) Each provider is part of a lawful combination under which each shares substantial financial risk in connection with the PSO's operations;

(3) Both, or all, providers are part of a controlled group of corporations under section 1563 of the Internal Revenue Code of 1986; or

(4) Both, or all, providers are part of an affiliated service group under section 414 of that Code.

(b) Each affiliated provider of the PSO shares, directly or indirectly, substantial financial risk for the furnishing of services the PSO is obligated to provide under the contract.

(c) Affiliated providers, as a whole or in part, have at least a majority financial interest in the PSO.

(d) For purposes of paragraph(a)(1) of this section, control is presumed to exist if one party, directly or indirectly, owns, controls, or holds the power to vote, or proxies for, not less than 51 percent of the voting rights or governance right of another.


§ 422.356 Determining substantial financial risk and majority financial interest.

(a) Determining substantial financial risk. The PSO must demonstrate to HCFA'S satisfaction that it apportions a significant part of the financial risk of the PSO enterprise under the M+C contract to each affiliated provider. The PSO must demonstrate that the financial arrangements among its affiliated providers constitute "substantial" risk in the PSO for each affiliated provider. The following mechanisms may constitute risk-sharing arrangements, and may have to be used in combination to demonstrate substantial financial risk in the PSO enterprise.

(1) Agreement by a provider to accept capitation payment for each Medicare enrollee.

(2) Agreement by a provider to accept as payment a predetermined percentage of the PSO premium or the PSO's revenue.

(3) The PSO's use of significant financial incentives for its affiliated providers, with the aim of achieving utilization management and cost containment goals. Permissible methods include the following:

(i) Affiliated providers agree to a withholding of a significant amount of the compensation due them, to be used for any of the following:

(A) To cover losses of the PSO.

(B) To cover losses of other affiliated providers.

(C) To be returned to the affiliated provider if the PSO meets its utilization management or cost containment goals for the specified time period.

(D) To be distributed among affiliated providers if the PSO meets its utilization management or cost-containment goals for the specified time period.

(ii) Affiliated providers agree to preestablished cost or utilization targets for the PSO and to subsequent significant financial rewards and penalties (which may include a reduction in payments to the provider) based on the PSO's performance in meeting the targets.

(4) Other mechanisms that demonstrate significant shared financial risk.

(b) Determining majority financial interest. Majority financial interest means maintaining effective control of the PSO.


§ 422.370 Waiver of State licensure.

For an organization that seeks to contract to offer an M+C plan under this subpart, HCFA may waive the...
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State licensure requirement of section 1855(a)(1) of the Act if—
(a) The organization requests a waiver no later than November 1, 2002; and
(b) HCFA determines there is a basis for a waiver under § 422.372.


§ 422.372 Basis for waiver of State licensure.

(a) General rule. Subject to this section and to paragraphs (a) and (e) of § 422.374, HCFA may waive the State licensure requirement if the organization has applied (except as provided in paragraph (b)(4) of this section) for the most closely appropriate State license or authority to conduct business as an M+C plan.

(b) Basis for waiver of State licensure. Any of the following may constitute a basis for HCFA’s waiver of State licensure:

(1) Failure to act timely on application. The State failed to complete action on the licensing application within 90 days of the date the State received a substantially complete application.

(2) Denial of application based on discriminatory treatment. The State has—
   (i) Denied the license application on the basis of material requirements, procedures, or standards (other than solvency requirements) not generally applied by the State to other entities engaged in a substantially similar business; or
   (ii) Required, as a condition of licensure, that the organization offer any product or plan other than an M+C plan.

(3) Denial of application based on different solvency requirements. (i) The State has denied the application, in whole or in part, on the basis of the organization’s failure to meet solvency requirements that are different from those set forth in §§ 422.380 through 422.390, or
   (ii) HCFA determines that the State has imposed, as a condition of licensure, any documentation or information requirements relating to solvency or other material requirements, procedures, or standards relating to solvency that are different from the requirements, procedures, or standards set forth by HCFA to implement, monitor, and enforce §§ 422.380 through 422.390.

(4) State declines to accept licensure application. The appropriate State licensing authority has given the organization written notice that it will not accept its licensure application.

[63 FR 35098, June 26, 1998]

§ 422.374 Waiver request and approval process.

(a) Substantially complete waiver request. The organization must submit a substantially complete waiver request that clearly demonstrates and documents its eligibility for a waiver under § 422.372.

(b) HCFA gives the organization written notice of granting or denial of waiver within 60 days of receipt of a substantially complete waiver request.

(c) Subsequent waiver requests. An organization that has had a waiver request denied, may submit subsequent waiver requests until November 1, 2002.

(d) Effective date. A waiver granted under § 422.370 will be effective on the effective date of the organization’s M+C contract.

(e) Consistency in application. HCFA reserves the right to revoke waiver eligibility if it subsequently determines that the organization’s M+C application is significantly different from the application submitted by the organization to the State licensing authority.

[63 FR 25377, May 7, 1998, as amended at 63 FR 35098, June 26, 1998]

§ 422.376 Conditions of the waiver.

A waiver granted under this section is subject to the following conditions:

(a) Limitation to State. The waiver is effective only for the particular State for which it is granted and does not apply to any other State. For each State in which the organization wishes to operate without a State license, it must submit a waiver request and receive a waiver.

(b) Limitation to 36-month period. The waiver is effective for 36 months or through the end of the calendar year in which the 36 month period ends unless it is revoked based on paragraph (c) of this section.

(c) Mid-period revocation. During the waiver period (set forth in paragraph
(b) of this section), the waiver is automatically revoked upon—
(1) Termination of the M+C contract;
(2) The organization's compliance with the State licensure requirement of section 1855(a)(1) of the Act; or
(3) The organization's failure to comply with §422.378.
[63 FR 25377, May 7, 1998]

§ 422.378 Relationship to State law.

(a) Preemption of State law. Any provisions of State law that relate to the licensing of the organization and that prohibit the organization from providing coverage under a contract as specified in this subpart, are superseded.

(b) Consumer protection and quality standards. (1) A waiver of State licensure granted under this subpart is conditioned upon the organization's compliance with all State consumer protection and quality standards that—
(i) Would apply to the organization if it were licensed under State law;
(ii) Generally apply to other M+C organizations and plans in the State; and
(iii) Are consistent with the standards established under this part.
(2) The standards specified in paragraph (b)(1) of this section do not include any standard preempted under section 1856(b)(3)(B) of the Act.
(c) Incorporation into contract. In contracting with an organization that has a waiver of State licensure, HCFA incorporates into the contract the requirements specified in paragraph (b) of this section.
(d) Enforcement. HCFA may enter into an agreement with a State for the State to monitor and enforce compliance with the requirements specified in paragraph (b) of this section by an organization that has obtained a waiver under this subpart.
[63 FR 25377, May 7, 1998]

§ 422.380 Solvency standards.

General rule. A PSO or the legal entity of which the PSO is a component that has been granted a waiver under §422.370 must have a fiscally sound operation that meets the requirements of §§422.382 through 422.390.
[63 FR 25377, May 7, 1998]

§ 422.382 Minimum net worth amount.

(a) At the time an organization applies to contract with HCFA as a PSO under this part, the organization must have a minimum net worth amount, as determined under paragraph (c) of this section, of:
(1) At least $1,500,000, except as provided in paragraph (a)(2) of this section.
(2) No less than $1,000,000 based on evidence from the organization's financial plan (under §422.384) demonstrating to HCFA's satisfaction that the organization has available to it an administrative infrastructure that HCFA considers appropriate to reduce, control or eliminate start-up administrative costs.
(b) After the effective date of a PSO's M+C contract, a PSO must maintain a minimum net worth amount equal to the greater of—
(1) One million dollars;
(2) Two percent of annual premium revenues as reported on the most recent annual financial statement filed with HCFA for up to and including the first $150,000,000 of annual premiums and 1 percent of annual premium revenues on premiums in excess of $150,000,000;
(3) An amount equal to the sum of three months of uncovered health care expenditures as reported on the most recent financial statement filed with HCFA; or
(4) Using the most recent financial statement filed with HCFA, an amount equal to the sum of—
(i) Eight percent of annual health care expenditures paid on a non-capitated basis to non-affiliated providers; and
(ii) Four percent of annual health care expenditures paid on a capitated basis to non-affiliated providers plus annual health care expenditures paid on a non-capitated basis to affiliated providers.
(iii) Annual health care expenditures that are paid on a capitated basis to affiliated providers are not included in the calculation of the net worth requirement (regardless of downstream arrangements from the affiliated provider) under paragraphs (a) and (b)(4) of this section.
[63 FR 25377, May 7, 1998]
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(c) Calculation of the minimum net worth amount—(1) Cash requirement. 
(i) At the time of application, the organization must maintain at least $750,000 of the minimum net worth amount in cash or cash equivalents.
(ii) After the effective date of a PSO's M+C contract, a PSO must maintain the greater of $750,000 or 40 percent of the minimum net worth amount in cash or cash equivalents.

(2) Intangible assets. An organization may include intangible assets, the value of which is based on Generally Accepted Accounting Principles (GAAP), in the minimum net worth amount calculation subject to the following limitations—
(i) At the time of application. (A) Up to 20 percent of the minimum net worth amount, provided at least $1,000,000 of the minimum net worth amount is met through cash or cash equivalents; or
(B) Up to 10 percent of the minimum net worth amount, if less than $1,000,000 of the minimum net worth amount is met through cash or cash equivalents, or if HCFA has used its discretion under paragraph (a)(2) of this section.
(ii) From the effective date of the contract. (A) Up to 20 percent of the minimum net worth amount if the greater of $1,000,000 or 67 percent of the minimum net worth amount is met by cash or cash equivalents; or
(B) Up to ten percent of the minimum net worth amount if the greater of $1,000,000 or 67 percent of the minimum net worth amount is not met by cash or cash equivalents.

(3) Health care delivery assets. Subject to the other provisions of this section, a PSO may apply 100 percent of the GAAP depreciated value of health care delivery assets (HCDAs) to satisfy the minimum net worth amount.

(4) Other assets. A PSO may apply other assets not used in the delivery of health care provided that those assets are valued according to statutory accounting practices (SAP) as defined by the State.

(5) Subordinated debts and subordinated liabilities. Fully subordinated debt and subordinated liabilities are excluded from the minimum net worth amount calculation.

(6) Deferred acquisition costs. Deferred acquisition costs are excluded from the calculation of the minimum net worth amount.

§ 422.384 Financial plan requirement.

(a) General rule. At the time of application, an organization must submit a financial plan acceptable to HCFA.

(b) Content of plan. A financial plan must include—
(1) A detailed marketing plan;
(2) Statements of revenue and expense on an accrual basis;
(3) Cash-flow statements;
(4) Balance sheets;
(5) Detailed justifications and assumptions in support of the financial plan including, where appropriate, certification of reserves and actuarial liabilities by a qualified actuary; and
(6) If applicable, statements of the availability of financial resources to meet projected losses.

(c) Period covered by the plan. A financial plan must—
(1) Cover the first 12 months after the estimated effective date of a PSO's M+C contract; or
(2) If the PSO is projecting losses, cover 12 months beyond the end of the period for which losses are projected.

(d) Funding for projected losses. Except for the use of guarantees, LOC, and other means as provided in §422.384(e), (f) and (g), an organization must have the resources for meeting projected losses on its balance sheet in cash or a form that is convertible to cash in a timely manner, in accordance with the PSO's financial plan.

(e) Guarantees and projected losses. Guarantees will be an acceptable resource to fund projected losses, provided that a PSO—
(1) Meets HCFA's requirements for guarantors and guarantee documents as specified in §422.390; and
(2) Obtains from the guarantor cash or cash equivalents to fund the projected losses timely, as follows—
(i) Prior to the effective date of a PSO's M+C contract, the amount of the projected losses for the first two quarters;
(ii) During the first quarter and prior to the beginning of the second quarter.
of a PSO's M+C contract, the amount of projected losses through the end of the third quarter; and

(iii) During the second quarter and prior to the beginning of the third quarter of a PSO's M+C contract, the amount of projected losses through the end of the fourth quarter.

(3) If the guarantor complies with the requirements in paragraph (e)(2) of this section, the PSO, in the third quarter, may notify HCFA of its intent to reduce the period of advance funding of projected losses. HCFA will notify the PSO within 60 days of receiving the PSO's request if the requested reduction in the period of advance funding will not be accepted.

(4) If the guarantee requirements in paragraph (e)(2) of this section are not met, HCFA may take appropriate action, such as requiring funding of projected losses through means other than a guarantee. HCFA retains discretion to require other methods or timing of funding, considering factors such as the financial condition of the guarantor and the accuracy of the financial plan.

(f) Letters of credit. Letters of credit are an acceptable resource to fund projected losses, provided they are irrevocable, unconditional, and satisfactory to HCFA. They must be capable of being promptly paid upon presentation of a sight draft under the letters of credit without further reference to any other agreement, document, or entity. HCFA retains discretion to require other methods or timing of funding, considering factors such as the financial condition of the guarantor and the accuracy of the financial plan.

(g) Other means. If satisfactory to HCFA, a PSO may use the following to fund projected losses—

(1) Lines of credit from regulated financial institutions;

(2) Legally binding agreements for capital contributions; or

(3) Legally binding agreements of a similar quality and reliability as permitted in paragraphs (g)(1) and (2) of this section.

(h) Application of guarantees. Letters of credit or other means of funding projected losses. Notwithstanding any other provision of this section, a PSO may use guarantees, letters of credit and, beginning one year after the effective date of a PSO's M+C contract, other means of funding projected losses, but only in a combination or sequence that HCFA considers appropriate.


§ 422.386 Liquidity.

(a) A PSO must have sufficient cash flow to meet its financial obligations as they become due and payable.

(b) To determine whether the PSO meets the requirement in paragraph (a) of this section, HCFA will examine the following—

(1) The PSO's timeliness in meeting current obligations;

(2) The extent to which the PSO's current ratio of assets to liabilities is maintained at 1:1 including whether there is a declining trend in the current ratio over time; and

(3) The availability of outside financial resources to the PSO.

(c) If HCFA determines that a PSO fails to meet the requirement in paragraph (b)(1) of this section, HCFA will require the PSO to initiate corrective action and pay all overdue obligations.

(d) If HCFA determines that a PSO fails to meet the requirement of paragraph (b)(2) of this section, HCFA may require the PSO to initiate corrective action to—

(1) Change the distribution of its assets;

(2) Reduce its liabilities; or

(3) Make alternative arrangements to secure additional funding to restore the PSO's current ratio to 1:1.

(e) If HCFA determines that there has been a change in the availability of outside financial resources as required by paragraph (b)(3) of this section, HCFA requires the PSO to obtain funding from alternative financial resources.


§ 422.388 Deposits.

(a) Insolvency deposit. (1) At the time of application, an organization must deposit $100,000 in cash or securities (or any combination thereof) into an account in a manner that is acceptable to HCFA.

(2) The deposit must be restricted to use in the event of insolvency to help assure continuation of services or pay...
§ 422.390 Guarantees.

(a) General policy. A PSO, or the legal entity of which the PSO is a component, may apply to HCFA to use the financial resources of a guarantor for the purpose of meeting the requirements in §422.384. HCFA has the discretion to approve or deny approval of the use of a guarantor.

(b) Request to use a guarantor. To apply to use the financial resources of a guarantor, a PSO must submit to HCFA—

(1) Documentation that the guarantor meets the requirements for a guarantor under paragraph (c) of this section; and

(2) The guarantor's independently audited financial statements for the current year-to-date and for the two most recent fiscal years. The financial statements must include the guarantor's balance sheets, profit and loss statements, and cash flow statements.

(c) Requirements for guarantor. To serve as a guarantor, an organization must meet the following requirements:

(1) Be a legal entity authorized to conduct business within a State of the United States.

(2) Not be under Federal or State bankruptcy or rehabilitation proceedings.

(3) Have a net worth (not including other guarantees, intangibles and restricted reserves) equal to three times the amount of the PSO guarantee.

(4) If the guarantor is regulated by a State insurance commissioner, or other State official with authority for risk-bearing entities, it must meet the net worth requirement in §422.390(c)(3) with all guarantees and all investments in and loans to organizations covered by guarantees excluded from its assets.

(5) If the guarantor is not regulated by a State insurance commissioner, or other similar State official, it must meet the net worth requirement in §422.390(c)(3) with all guarantees and

§ 422.390 Guarantees.

(a) General policy. A PSO, or the legal entity of which the PSO is a component, may apply to HCFA to use the financial resources of a guarantor for the purpose of meeting the requirements in §422.384. HCFA has the discretion to approve or deny approval of the use of a guarantor.

(b) Request to use a guarantor. To apply to use the financial resources of a guarantor, a PSO must submit to HCFA—

(1) Documentation that the guarantor meets the requirements for a guarantor under paragraph (c) of this section; and

(2) The guarantor's independently audited financial statements for the current year-to-date and for the two most recent fiscal years. The financial statements must include the guarantor's balance sheets, profit and loss statements, and cash flow statements.

(c) Requirements for guarantor. To serve as a guarantor, an organization must meet the following requirements:

(1) Be a legal entity authorized to conduct business within a State of the United States.

(2) Not be under Federal or State bankruptcy or rehabilitation proceedings.

(3) Have a net worth (not including other guarantees, intangibles and restricted reserves) equal to three times the amount of the PSO guarantee.

(4) If the guarantor is regulated by a State insurance commissioner, or other State official with authority for risk-bearing entities, it must meet the net worth requirement in §422.390(c)(3) with all guarantees and all investments in and loans to organizations covered by guarantees excluded from its assets.

(5) If the guarantor is not regulated by a State insurance commissioner, or other similar State official, it must meet the net worth requirement in §422.390(c)(3) with all guarantees and
all investments in and loans to organizations covered by a guarantee and to related parties (subsidiaries and affiliates) excluded from its assets.

(d) Guarantee document. If the guarantee request is approved, a PSO must submit to HCFA a written guarantee document signed by an appropriate authority of the guarantor. The guarantee document must—

(1) State the financial obligation covered by the guarantee;
(2) Agree to—
   (i) Unconditionally fulfill the financial obligation covered by the guarantor; and
   (ii) Not subordinate the guarantee to any other claim on the resources of the guarantor;
(3) Declare that the guarantor must act on a timely basis, in any case not more than 5 business days, to satisfy the financial obligation covered by the guarantee; and
(4) Meet other conditions as HCFA may establish from time to time.

(e) Reporting requirement. A PSO must submit to HCFA the current internal financial statements and annual audited financial statements of the guarantor according to the schedule, manner, and form that HCFA requests.

(f) Modification, substitution, and termination of a guarantee. A PSO cannot modify, substitute or terminate a guarantee unless the PSO—

(1) Requests HCFA’s approval at least 90 days before the proposed effective date of the modification, substitution, or termination;
(2) Demonstrates to HCFA’s satisfaction that the modification, substitution, or termination will not result in insolvency of the PSO; and
(3) Demonstrates how the PSO will meet the requirements of this section.

(g) Nullification. If at any time the guarantor or the guarantee ceases to meet the requirements of this section, HCFA will notify the PSO that it ceases to recognize the guarantee document. In the event of this nullification, a PSO must—

(1) Meet the applicable requirements of this section within 15 business days; and
(2) If required by HCFA, meet a portion of the applicable requirements in less than the time period granted in paragraph (g)(1) of this section.

[63 FR 25379, May 7, 1998]

Subpart I—Organization Compliance With State Law and Preemption by Federal Law

SOURCE: 63 FR 35099, June 26, 1998, unless otherwise noted.

§ 422.400 State licensure requirement.

Except in the case of a PSO granted a waiver under subpart H of this part, each M+C organization must—

(a) Be licensed under State law, or otherwise authorized to operate under State law, as a risk-bearing entity (as defined in §422.2) eligible to offer health insurance or health benefits coverage in each State in which it offers one or more M+C plans;
(b) If not commercially licensed, obtain certification from the State that the organization meets a level of financial solvency and such other standards as the State may require for it to operate as an M+C organization;
(c) Demonstrate to HCFA that—
   (1) The scope of its license or authority allows the organization to offer the type of M+C plan or plans that it intends to offer in the State; and
   (2) If applicable, it has obtained the State certification required under paragraph (b) of this section.

[63 FR 25379, May 7, 1998]

§ 422.402 Federal preemption of State law.

(a) General preemption. Except as provided in paragraph (b) of this section, the rules, contract requirements, and standards established under this part supersede any State laws, regulations, contract requirements, or other standards that would otherwise apply to M+C organizations and their M+C plans only to the extent that such State laws are inconsistent with the standards established under this part. This preemption of State laws and other standards applies only to coverage pursuant to an M+C contract, and does not extend to benefits outside of such contract or to individuals who are not M+C enrollees of an organization with an M+C contract.
(b) Specific preemption. As they might otherwise apply to the M+C plans of an M+C organization in a State, State laws and regulations pertaining to the following areas are specifically preempted by this part:

(1) Benefit requirements, such as mandating the inclusion in an M+C plan of a particular service, or specifying the scope or duration of a service (for example, length of hospital stay, number of home health visits). State cost-sharing standards with respect to any benefits are preempted only if they are inconsistent with this part, as provided for in paragraph (a) of this section.

(2) Requirements relating to inclusion or treatment of providers and suppliers.

(3) Coverage determinations (including related appeal and grievance processes for all benefits included under an M+C contract). Determinations on issues other than whether a service is covered under an M+C contract, and the extent of enrollee liability under the M+C plan for such a service, are not considered coverage determinations for purposes of this paragraph.

(c) Except as provided in paragraphs (a) and (b) of this section, nothing in this section may be construed to affect or modify the provisions of any other law or regulation that imposes or preempts a specific State authority.

§ 422.404 State premium taxes prohibited.

(a) Basic rule. No premium tax, fee, or other similar assessment may be imposed by any State, the District of Columbia, the Commonwealth of Puerto Rico, the Virgin Islands, Guam, and American Samoa, or any of their political subdivision or other governmental authorities with respect to any payment HCFA makes on behalf of M+C enrollees under subpart F of this part.

(b) Construction. Nothing in this section shall be construed to exempt any M+C organization from taxes, fees, or other monetary assessments related to the net income or profit that accrues to, or is realized by, the organization from business conducted under this part, if that tax, fee, or payment is applicable to a broad range of business activity.
§ 422.501  General provisions.

(a) Basic rule. In order to qualify as an M+C organization, enroll beneficiaries in any M+C plans it offers, and be paid on behalf of Medicare beneficiaries enrolled in those plans, an M+C organization must enter into a contract with HCFA.

(b) Conditions necessary to contract as an M+C organization. Any entity seeking to contract as an M+C organization must:

(1) Be licensed by the State as a risk bearing entity in each State in which it seeks to offer an M+C plan as defined in §422.2.

(2) Meet the minimum enrollment requirements of §422.514, unless waived under §422.514(b).

(3) Have administrative and management arrangements satisfactory to HCFA, as demonstrated by at least the following:

(i) A policy making body that exercises oversight and control over the M+C organization’s policies and personnel to ensure that management actions are in the best interest of the organization and its enrollees.

(ii) Personnel and systems sufficient for the M+C organization to organize, plan, control, and evaluate financial and marketing activities, the furnishing of services, the quality assurance program, and the administrative and management aspects of the organization.

(iii) At a minimum, an executive manager, whose appointment and removal are under the control of the policy making body.

(iv) A fidelity bond or bonds, procured and maintained by the M+C organization, in an amount fixed by its policymaking body but not less than $100,000 per individual, covering each officer and employee entrusted with the handling of its funds. The bond may have reasonable deductibles, based upon the financial strength of the M+C organization.

(v) Insurance policies or other arrangements, secured and maintained by the M+C organization and approved by HCFA to insure the M+C organization against losses arising from professional liability claims, fire, theft, fraud, embezzlement, and other casualty risks.

(vi) A compliance plan that consists of the following:

1. Any director, officer, partner, or employee responsible for management or administration of an M+C organization.
2. Any person who is directly or indirectly the beneficial owner of more than 5 percent of the organization’s equity; or the beneficial owner of a mortgage, deed of trust, note, or other interest secured by and valuing more than 5 percent of the organization.
3. In the case of an M+C organization organized as a nonprofit corporation, an incorporator or member of such corporation under applicable State corporation law.
4. Any entity in which a person described in paragraph (1), (2), or (3) of this definition:
   (i) Is an officer, director, or partner; or
   (ii) Has the kind of interest described in paragraphs (1), (2), or (3) of this definition.
5. Any person that directly or indirectly controls, is controlled by, or is under common control with, the M+C organization.
6. Any spouse, child, or parent of an individual described in paragraph (1), (2), or (3) of this definition.

Related entity means any entity that is related to the M+C organization by common ownership or control and—

1. Performs some of the M+C organization’s management functions under contract or delegation;
2. Furnishes services to Medicare enrollees under an oral or written agreement; or
3. Leases real property or sells materials to the M+C organization at a cost of more than $2,500 during a contract period.

Significant business transaction means any business transaction or series of transactions of the kind specified in the above definition of “business transaction” that, during any fiscal year of the M+C organization, have a total value that exceeds $25,000 or 5 percent of the M+C organization’s total operating expenses, whichever is less.

§ 422.502 Contract provisions.

The contract between the M+C organization and HCFA must contain the following provisions:

(a) Agreement to comply with regulations and instructions. The M+C organization agrees to comply with all the applicable requirements and conditions set forth in this part and in general instructions. An M+C organization's compliance with paragraphs (a)(1) through (a)(13) of this section is material to performance of the contract. The M+C organization agrees—

(A) Written policies, procedures, and standards of conduct that articulate the organization's commitment to comply with all applicable Federal and State standards.

(B) The designation of a compliance officer and compliance committee that are accountable to senior management.

(C) Effective training and education between the compliance officer and the organization's employees.

(D) Effective lines of communication between the compliance officer and the organization's employees.

(E) Enforcement of standards through well-publicized disciplinary guidelines.

(F) Provision for internal monitoring and auditing.

(G) Procedures for ensuring prompt response to detected offenses and development of corrective action initiatives relating to the organization's M+C contract.

(4) Not accept new enrollees under a section 1876 reasonable cost contract in any area in which it seeks to offer an M+C plan.

(5) The M+C organization's contract must not have been terminated by HCFA under § 422.510 within the past 2 years unless—

(i) During the 6-month period beginning on the date the organization notified HCFA of the intention to terminate the most recent previous contract, there was a change in the statute or regulations that had the effect of increasing M+C payments in the payment area or areas at issue; or

(ii) HCFA has otherwise determined that circumstances warrant special consideration.

(c) Contracting authority. Under the authority of section 1857(c)(5) of the Act, HCFA may enter into contracts under this part without regard to Federal and Departmental acquisition regulations set forth in title 48 of the CFR and provisions of law or other regulations relating to the making, performance, amendment, or modification of contracts of the United States if HCFA determines that those provisions are inconsistent with the efficient and effective administration of the Medicare program.

(d) Protection against fraud and beneficiary protections. (1) HCFA annually audits the financial records (including data relating to Medicare utilization, costs, and computation of the ACR) of at least one-third of the M+C organizations offering M+C plans. These auditing activities are subject to monitoring by the Comptroller General.

(2) Each contract under this section must provide that HCFA, or any person or organization designated by HCFA has the right to:

(i) Inspect or otherwise evaluate the quality, appropriateness, and timeliness of services performed under the M+C contract;

(ii) Inspect or otherwise evaluate the facilities of the organization when there is reasonable evidence of some need for such inspection; and

(iii) Audit and inspect any books, contracts, and records of the M+C organization that pertain to—

(A) The ability of the organization or its first tier or downstream providers to bear the risk of potential financial losses; or

(B) Services performed or determinations of amounts payable under the contract.

(e) Severability of contracts. The contract must provide that, upon HCFA's request—

(1) The contract will be amended to exclude any M+C plan or State-licensed entity specified by HCFA; and

(2) A separate contract for any such excluded plan or entity will be deemed to be in place when such a request is made.

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(1) To accept new enrollments, make enrollments effective, process voluntary disenrollments, and limit involuntary disenrollments, as provided in subpart B of this part.

(2) That it will comply with the prohibition in §422.110 on discrimination in beneficiary enrollment.

(3) To provide—

(i) The basic benefits as required under §422.101 and, to the extent applicable, supplemental benefits under §422.102; and

(ii) Access to benefits as required under subpart C of this part;

(iii) In a manner consistent with professionally recognized standards of health care, all benefits covered by Medicare.

(4) To disclose information to beneficiaries in the manner and the form prescribed by HCFA as required under §422.111;

(5) To operate a quality assurance and performance improvement program and have an agreement for external quality review as required under subpart D of this part;

(6) To comply with all applicable provider requirements in subpart E of this part, including provider certification requirements, anti-discrimination requirements, provider participation and consultation requirements, the prohibition on interference with provider advice, limits on provider indemnification, rules governing payments to providers, and limits on physician incentive plans;

(7) To comply with all requirements in subpart M of this part governing coverage determinations, grievances, and appeals;

(8) To comply with the reporting requirements in §422.516 and the requirements in §422.257 for submitting encounter data to HCFA;

(9) That it will be paid under the contract in accordance with the payment rules in subpart F of this part;

(10) To develop its annual ACR, and submit all required information on premiums, benefits, and cost-sharing by May 1, as provided in subpart G of this part;

(11) That its contract may not be renewed or may be terminated in accordance with this subpart and subpart N of this part.

(12) To comply with all requirements that are specific to a particular type of M+C plan, such as the special rules for private fee-for-service plans in §§422.114 and 422.216 and the MSA requirements in §§422.56, 422.103, and 422.262, and

(13) To comply with the confidentiality and enrollee record accuracy requirements in §422.118.

(14) An M+C organization’s compliance with paragraphs (a)(1) through (a)(13) and (c) of this section is material to performance of the contract.

(b) Communication with HCFA. The M+C organization must have the capacity to communicate with HCFA electronically.

(c) Prompt payment. The M+C organization must comply with the prompt payment provisions of §422.520 and with instructions issued by HCFA, as they apply to each type of plan included in the contract.

(d) Maintenance of records. The M+C organization agrees to maintain for 6 years books, records, documents, and other evidence of accounting procedures and practices that—

(1) Are sufficient to do the following:

(i) Accommodate periodic auditing of the financial records (including data related to Medicare utilization, costs, and computation of the ACR) of M+C organizations.

(ii) Enable HCFA to inspect or otherwise evaluate the quality, appropriateness and timeliness of services performed under the contract, and the facilities of the organization.

(iii) Enable HCFA to audit and inspect any books and records of the M+C organization that pertain to the ability of the organization to bear the risk of potential financial losses, or to services performed or determinations of amounts payable under the contract.

(iv) Properly reflect all direct and indirect costs claimed to have been incurred and used in the preparation of the ACR proposal.

(v) Establish component rates of the ACR for determining additional and supplementary benefits.

(vi) Determine the rates utilized in setting premiums for State insurance agency purposes and for other government and private purchasers; and

(2) Include at least records of the following:
(i) Ownership and operation of the M+C organization's financial, medical, and other record keeping systems.

(ii) Financial statements for the current contract period and six prior periods.

(iii) Federal income tax or information returns for the current contract period and six prior periods.

(iv) Asset acquisition, lease, sale, or other action.

(v) Agreements, contracts, and subcontracts.

(vi) Franchise, marketing, and management agreements.

(vii) Schedules of charges for the M+C organization's fee-for-service patients.

(viii) Matters pertaining to costs of operations.

(ix) Amounts of income received by source and payment.

(x) Cash flow statements.

(xi) Any financial reports filed with other Federal programs or State authorities.

(e) Access to facilities and records. The M+C organization agrees to the following:

(1) HHS, the Comptroller General, or their designee may evaluate, through inspection or other means—
   (i) The quality, appropriateness, and timeliness of services furnished to Medicare enrollees under the contract;
   (ii) The facilities of the M+C organization; and
   (iii) The enrollment and disenrollment records for the current contract period and six prior periods.

(2) HHS, the Comptroller General, or their designees may audit, evaluate, or inspect any books, contracts, medical records, patient care documentation, and other records of the M+C organization, related entity, contractor, subcontractor, or its transferee that pertain to any aspect of services performed, reconciliation of benefit liabilities, and determination of amounts payable under the contract, or as the Secretary may deem necessary to enforce the contract.

(3) The M+C organization agrees to make available, for the purposes specified in paragraph (d) of this section, its premises, physical facilities and equipment, records relating to its Medicare enrollees, and any additional relevant information that HCFA may require.

(4) HHS, the Comptroller General, or their designee's right to inspect, evaluate, and audit extends through 6 years from the final date of the contract period or completion of audit, whichever is later unless—

(i) HCFA determines there is a special need to retain a particular record or group of records for a longer period and notifies the M+C organization at least 30 days before the normal disposition date;

(ii) There has been a termination, dispute, or fraud or similar fault by the M+C organization, in which case the retention may be extended to 6 years from the date of any resulting final resolution of the termination, dispute, or fraud or similar fault; or

(iii) HCFA determines that there is a reasonable possibility of fraud, in which case it may inspect, evaluate, and audit the M+C organization at any time.

(f) Disclosure of information. The M+C organization agrees to submit—

(1) To HCFA, certified financial information that must include the following:
   (i) Such information as HCFA may require demonstrating that the organization has a fiscally sound operation.
   (ii) Such information as HCFA may require pertaining to the disclosure of ownership and control of the M+C organization.

(2) To HCFA, all information that is necessary for HCFA to administer and evaluate the program and to simultaneously establish and facilitate a process for current and prospective beneficiaries to exercise choice in obtaining Medicare services. This information includes, but is not limited to:
   (i) The benefits covered under an M+C plan;
   (ii) The M+C monthly basic beneficiary premium and M+C monthly supplemental beneficiary premium, if any, for the plan or in the case of an MSA plan, the M+C monthly MSA premium.
   (iii) The service area and continuation area, if any, of each plan and the enrollment capacity of each plan;
   (iv) Plan quality and performance indicators for the benefits under the plan including —
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(A) Disenrollment rates for Medicare enrollees electing to receive benefits through the plan for the previous 2 years;
(B) Information on Medicare enrollee satisfaction;
(C) Information on health outcomes;
(D) The recent record regarding compliance of the plan with requirements of this part, as determined by HCFA; and
(E) Other information determined by HCFA to be necessary to assist beneficiaries in making an informed choice among M+C plans and traditional Medicare;

(v) Information about beneficiary appeals and their disposition;
(vi) Information regarding all formal actions, reviews, findings, or other similar actions by States, other regulatory bodies, or any other certifying or accrediting organization;
(vii) For M+C organizations offering an MSA plan, information specified by HCFA for HCFA’s use in preparing its report to the Congress on the MSA demonstration, including data specified by HCFA in the areas of selection, use of preventative care, and access to services.
(viii) To HCFA, any other information deemed necessary by HCFA for the administration or evaluation of the Medicare program.

(3) To its enrollees all informational requirements under §422.64 and, upon an enrollee’s request, the financial disclosure information required under §422.516.

(g) Beneficiary financial protections. The M+C organization agrees to comply with the following requirements:

(1) Each M+C organization must adopt and maintain arrangements satisfactory to HCFA to protect its enrollees from incurring liability (for example, as a result of an organization’s insolvency or other financial difficulties) for payment of any fees that are the legal obligation of the M+C organization. To meet this requirement, the M+C organization must—
(i) Indemnify the beneficiary enrollee for payment of any fees that are the legal obligation of the M+C organization for services furnished by providers that do not contract, or that have not otherwise entered into an agreement with the M+C organization, to provide services to the organization’s beneficiary enrollees.
(ii) For all enrollees, for the duration of the contract period for which HCFA payments have been made; and
(iii) For enrollees who are hospitalized on the date its contract with HCFA terminates, or, in the event of an insolvency, through discharge.

(3) In meeting the requirements of this paragraph, other than the provider contract requirements specified in paragraph (g)(1)(i) of this section, the M+C organization may use—
(i) Contractual arrangements;
(ii) Insurance acceptable to HCFA;
(iii) Financial reserves acceptable to HCFA; or
(iv) Any other arrangement acceptable to HCFA.

(h) Requirements of other laws and regulations. (1) The M+C organization agrees to comply with—
(i) Title VI of the Civil Rights Act of 1964 as implemented by regulations at 45 CFR part 84;
(ii) The Age Discrimination Act of 1975 as implemented by regulations at 45 CFR part 91;
(iii) The Rehabilitation Act of 1973;
(iv) The Americans With Disabilities Act;
(v) Other laws applicable to recipients of Federal funds; and
(vi) All other applicable laws and rules.

(2) M+C organizations receiving Federal payments under M+C contracts, and related entities, contractors, and subcontractors paid by an M+C organization to fulfill its obligations under its M+C contract are subject to certain laws that are applicable to individuals and entities receiving Federal funds. M+C organizations must inform all related entities, contractors and subcontractors that payments that they receive are, in whole or in part, from Federal funds.
(i) M+C organization relationship with related entities, contractors, and subcontractors. (1) Notwithstanding any relationship(s) that the M+C organization may have with related entities, contractors, or subcontractors, the M+C organization maintains ultimate responsibility for adhering to and otherwise fully complying with all terms and conditions of its contract with HCFA.

(2) The M+C organization agrees to require all related entities, contractors, or subcontractors to agree that—
   (i) HHS, the Comptroller General, or their designees have the right to inspect, evaluate, and audit any pertinent contracts, books, documents, papers, and records of the related entity(ies), contractor(s), or subcontractor(s) involving transactions related to the M+C contract; and
   (ii) HHS', the Comptroller General's, or their designee's right to inspect, evaluate, and audit any pertinent information for any particular contract period will exist through 6 years from the final date of the contract period or from the date of completion of any audit, whichever is later.

(3) All contracts or written arrangements between M+C organizations and providers, related entities, contractors, subcontractors, first tier and downstream entities must contain the following:
   (i) Enrollee protection provisions that provide, consistent with paragraph (g)(1) of this section, arrangements that prohibit providers from holding an enrollee liable for payment of any fees that are the obligation of the M+C organization.
   (ii) Accountability provisions that indicate that—
      (A) The M+C organization oversees and is accountable to HCFA for any functions or responsibilities that are described in these standards; and
      (B) The M+C organization may only delegate activities or functions to a provider, related entity, contractor, or subcontractor in a manner consistent with requirements set forth at paragraph (i)(4) of this section.
   (iii) A provision requiring that any services or other activity performed by a related entity, contractor, subcontractor, or first-tier or downstream entity in accordance with a contract or written agreement are consistent and comply with the M+C organization's contractual obligations.

(4) If any of the M+C organizations' activities or responsibilities under its contract with HCFA are delegated to other parties, the following requirements apply to any related entity, contractor, subcontractor, or provider:
   (i) Written arrangements must specify delegated activities and reporting responsibilities.
   (ii) Written arrangements must either provide for revocation of the delegation activities and reporting requirements or specify other remedies in instances where HCFA or the M+C organization determine that such parties have not performed satisfactorily.
   (iii) Written arrangements must specify that the performance of the parties is monitored by the M+C organization on an ongoing basis.
   (iv) Written arrangements must specify that either—
      (A) The credentials of medical professionals affiliated with the party or parties will be either reviewed by the M+C organization; or
      (B) The credentialing process will be reviewed and approved by the M+C organization and the M+C organization must audit the credentialing process on an ongoing basis.
   (v) All contracts or written arrangements must specify that the related entity, contractor, or subcontractor must comply with all applicable Medicare laws, regulations, and HCFA instructions.

(5) If the M+C organization delegates selection of the providers, contractors, or subcontractor to another organization, the M+C organization's written arrangements with that organization must state that the HCFA-contracting M+C organization retains the right to approve, suspend, or terminate any such arrangement.

(j) Additional contract terms. The M+C organization agrees to include in the contract such other terms and conditions as HCFA may find necessary and appropriate in order to implement requirements in this part.

(k) Severability of contracts. The contract must provide that, upon HCFA's request—
§ 422.504

(1) The contract will be amended to exclude any M+C plan or State-licensed entity specified by HCFA; and

(2) A separate contract for any such excluded plan or entity will be deemed to be in place when such a request is made.

(1) Certification of data that determine payment. As a condition for receiving a monthly payment under subpart F of this part, the M+C organization agrees that its chief executive officer (CEO), chief financial officer (CFO), or an individual delegated the authority to sign on behalf of one of these officers, and who reports directly to such officer, must request payment under the contract on a document that certifies (based on best knowledge, information, and belief) the accuracy, completeness, and truthfulness of relevant data that HCFA requests. Such data include specified enrollment information, encounter data, and other information that HCFA may specify.

(1) The CEO, CFO, or an individual delegated the authority to sign on behalf of one of these officers, who reports directly to such officer, must certify that each enrollee for whom the organization is requesting payment is validly enrolled in an M+C plan offered by the organization and the information relied upon by HCFA in determining payment (based on best knowledge, information, and belief) is accurate, complete, and truthful.

(2) The CEO, CFO, or an individual delegated with the authority to sign on behalf of one of these officers, must certify (based on best knowledge, information, and belief) that the encounter data it submits under § 422.257 are accurate, complete, and truthful.

(3) If such encounter data are generated by a related entity, contractor, or subcontractor of an M+C organization, such entity, contractor, or subcontractor must similarly certify (based on best knowledge, information, and belief) the accuracy, completeness, and truthfulness of the data.

(4) The CEO, CFO, or an individual delegated the authority to sign on behalf of one of these officers, and who reports directly to such officer, must certify (based on best knowledge, information, and belief) that the information in its ACR submission is accurate, complete, and truthful and fully conforms to the requirements in § 422.310.


§ 422.504 Effective date and term of contract.

(a) Effective date. The contract is effective on the date specified in the contract between the M+C organization and HCFA and, for a contract that provides for coverage under an MSA plan, not earlier than January 1999.

(b) Term of contract. Each contract is for a period of at least 12 months.

(c) Renewal of contract. In accordance with § 422.506, contracts are renewed annually only if—

(1) HCFA informs the M+C organization that it authorizes a renewal; and

(2) The M+C organization has not provided HCFA with a notice of intention not to renew.


§ 422.506 Nonrenewal of contract.

(a) Nonrenewal by an M+C organization. (1) An M+C organization may elect not to renew its contract with HCFA as of the end of the term of the contract for any reason provided it meets the timeframes for doing so set forth in paragraphs (a)(2) and (a)(3) of this section.

(2) If an M+C organization does not intend to renew its contract, it must notify—

(i) HCFA in writing, by July 1 of the year in which the contract would end;

(ii) Each Medicare enrollee, at least 90 days before the date on which the nonrenewal is effective. This notice must include a written description of alternatives available for obtaining Medicare services within the service area, including alternative M+C plans, Medigap options, and original Medicare and must receive HCFA approval.

(iii) The general public, at least 90 days before the end of the current calendar year, by publishing a notice in one or more newspapers of general circulation in each community located in the M+C organization’s service area.

(3) HCFA may accept a nonrenewal notice submitted after July 1 if—
§ 422.508 Modification or termination of contract by mutual consent.

(a) A contract may be modified or terminated at any time by written mutual consent.

(1) If the contract is terminated by mutual consent, except as provided in paragraph (b) of this section, the M+C organization must provide notice to its Medicare enrollees and the general public as provided in §422.512(b)(2) and (b)(3).

(2) If the contract is modified by mutual consent, the M+C organization must notify its Medicare enrollees of any changes that HCFA determines are appropriate for notification within timeframes specified by HCFA.

(b) If the contract terminated by mutual consent is replaced the day following such termination by a new M+C contract, the M+C organization is not required to provide the notice specified in paragraph (a)(1) of this section.

§ 422.510 Termination of contract by HCFA.

(a) Termination by HCFA. HCFA may terminate a contract for any of the following reasons:

(1) The M+C organization has failed substantially to carry out the terms of its contract with HCFA.

(2) The M+C organization is carrying out its contract with HCFA in a manner that is inconsistent with the effective and efficient implementation of this part.

(3) HCFA determines that the M+C organization no longer meets the requirements of this part for being a contracting organization.

(4) The M+C organization commits or participates in fraudulent or abusive activities affecting the Medicare program, including submission of fraudulent data.

(5) The M+C organization experiences financial difficulties so severe that its ability to make necessary health services available is impaired to the point of posing an imminent and serious risk to the health of its enrollees, or otherwise fails to make services available to the extent that such a risk to health exists.

§422.512 Termination of contract by the M+C organization.

(a) Cause for termination. The M+C organization may terminate the M+C contract if HCFA fails to substantially carry out the terms of the contract.

(b) Notice. The M+C organization must give advance notice as follows:

(1) To HCFA, at least 90 days before the intended date of termination. This notice must specify the reasons why the M+C organization is requesting contract termination.
§ 422.514 Minimum enrollment requirements.

(a) Basic rule. Except as provided in paragraph (b) of this section, HCFA does not enter into a contract under this subpart unless the organization meets the following minimum enrollment requirement—

(1) At least 5,000 individuals (or 1,500 individuals if the organization is a PSO) are enrolled for the purpose of receiving health benefits from the organization; or

(2) At least 1,500 individuals (or 500 individuals if the organization is a PSO) are enrolled for purposes of receiving health benefits from the organization and the organization primarily serves individuals residing outside of urbanized areas as defined in §412.62(f) (or, in the case of a PSO, the PSO meets the requirements in §422.352(c)).

(3) Except as provided for in paragraph (b) of this section, an M+C organization must maintain a minimum enrollment as defined in paragraphs (a)(1) and (a)(2) of this section for the duration of its contract.

(b) Minimum enrollment waiver. (1) For a contract applicant or M+C organization that does not meet the applicable requirement of paragraph (a) of this section at application for an M+C contract or during the first 3 years of the contract, HCFA may waive the minimum enrollment requirement as provided for below. To receive a waiver, a contract applicant or M+C organization must demonstrate to HCFA’s satisfaction that it is capable of administering and managing an M+C contract and is able to manage the level of risk required under the contract. Factors that HCFA takes into consideration in making this evaluation include the extent to which—

(i) The contract applicant or M+C organization’s management and providers have previous experience in managing and providing health care services under a risk-based payment arrangement to at least as many individuals as the applicable minimum enrollment for the entity as described in paragraph (a) of this section, or

(ii) The contract applicant or M+C organization has the financial ability to bear financial risk under an M+C contract. In determining whether an organization is capable of bearing risk, HCFA considers factors such as the organization’s management experience as described in paragraph (b)(1)(i) of this section and stop-loss insurance that is adequate and acceptable to HCFA; and

(iii) The contract applicant or M+C organization is able to establish a marketing and enrollment process that allows it to meet the applicable enrollment requirement specified in paragraph (a) of this section before completion of the third contract year.

(2) If an M+C organization fails to meet the enrollment requirement in the first year, HCFA may waive the minimum requirements for another year provided that the organization—

(i) Requests an additional minimum enrollment waiver no later than 320 days before the end of the first year;
§422.516 Reporting requirements.

(a) Required information. Each M+C organization must have an effective procedure to develop, compile, evaluate, and report to HCFA, to its enrollees, and to the general public, at the times and in the manner that HCFA requires, and while safeguarding the confidentiality of the doctor-patient relationship, statistics and other information with respect to the following:

(1) The cost of its operations.
(2) The patterns of utilization of its services.
(3) The availability, accessibility, and acceptability of its services.
(4) To the extent practical, developments in the health status of its enrollees.
(5) Information demonstrating that the M+C organization has a fiscally sound operation.
(6) Other matters that HCFA may require.

(b) Significant business transactions. Each M+C organization must report to HCFA annually, within 120 days of the end of its fiscal year (unless for good cause shown, HCFA authorizes an extension of time), the following:

(1) A description of significant business transactions (as defined in §422.500) between the M+C organization and a party in interest.
(2) With respect to those transactions—
   (i) A showing that the costs of the transactions listed in paragraph (c) of this section do not exceed the costs that would be incurred if these transactions were with someone who is not a party in interest; or
   (ii) If they do exceed, a justification that the higher costs are consistent with prudent management and fiscal soundness requirements.

(c) Failure to meet enrollment requirements. HCFA may elect not to renew its contract with an M+C organization that fails to meet the applicable enrollment requirement in paragraph (a) of this section.

(d) Reporting and disclosure under ERISA. (1) For any employees’ health benefits plan that includes an M+C organization in its offerings, the M+C organization must furnish, upon request, the information the plan needs to fulfill its reporting and disclosure obligations (with respect to the particular M+C organization) under the Employee Retirement Income Security Act of 1974 (ERISA).

(2) The M+C organization must furnish the information to the employer or the employer’s designee, or to the
§ 422.550 General provisions.

(a) What constitutes change of ownership—(1) Partnership. The removal, addition, or substitution of a partner, unless the partners expressly agree otherwise as permitted by applicable State law, constitutes a change of ownership.

(2) Corporation. (i) The merger of the M+C organization’s corporation into another corporation or the consolidation of the M+C organization with one or more other corporations, resulting in a new corporate body, constitutes a change of ownership.

(ii) Transfer of corporate stock or the merger of another corporation into the M+C organization’s corporation, with the M+C organization surviving, does not ordinarily constitute change of ownership.

(b) Advance notice requirement. (1) An M+C organization that has a Medicare contract in effect and is considering or negotiating a change in ownership must notify HCFA at least 60 days before the anticipated effective date of the change. The M+C organization

§ 422.524 Special rules for RFB societies.

In order to participate as an M+C organization, an RFB society—

(a) May not impose any limitation on membership based on any factor related to health status; and

(b) Must offer, in addition to the M+C RFB plan, health coverage to individuals who are members of the church or convention or group of churches with which the society is affiliated, but who are not entitled to receive benefits from the Medicare program.

Subpart L—Effect of Change of Ownership or Leasing of Facilities During Term of Contract

Source: 63 FR 35067, June 26, 1998, unless otherwise noted.

Editorial Note: Nomenclature changes to subpart L appear at 63 FR 35106, June 26, 1998.

§ 422.520 Prompt payment by M+C organization.

(a) Contract between HCFA and the M+C organization.

(1) The contract between HCFA and the M+C organization must provide that the M+C organization will pay 95 percent of the “clean claims” within 30 days of receipt if they are submitted by, or on behalf of, an enrollee of an M+C private fee-for-service plan or are claims for services that are not furnished under a written agreement between the organization and the provider.

(2) The M+C organization must pay interest on clean claims that are not paid within 30 days in accordance with sections 1816(c)(2)(B) and 1842(c)(2)(B).

(3) All other claims must be paid or denied within 60 calendar days from the date of the request.

(b) Contracts between M+C organizations and providers and suppliers. Contracts or other written agreements between M+C organizations and providers must contain a prompt payment provision, the terms of which are developed and agreed to by both the M+C organization and the relevant provider.

(c) Failure to comply. If HCFA determines, after giving notice and opportunity for hearing, that an M+C organization has failed to make payments in accordance with paragraph (a) of this section, HCFA may provide—

(1) For direct payment of the sums owed to providers, or M+C private fee-for-service plan enrollees; and

(2) For appropriate reduction in the amounts that would otherwise be paid to the organization, to reflect the amounts of the direct payments and the cost of making those payments.

must also provide updated financial information and a discussion of the financial and solvency impact of the change of ownership on the surviving organization.

(2) If the M+C organization fails to give HCFA the required notice timely, it continues to be liable for capitation payments that HCFA makes to it on behalf of Medicare enrollees after the date of change of ownership.

(c) Novation agreement defined. A novation agreement is an agreement among the current owner of the M+C organization, the prospective new owner, and HCFA—

(1) That is embodied in a document executed and signed by all three parties;

(2) That meets the requirements of § 422.552; and

(3) Under which HCFA recognizes the new owner as the successor in interest to the current owner's Medicare contract.

(d) Effect of change of ownership without novation agreement. Except to the extent provided in paragraph (b)(2) of this section, the effect of a change of ownership without a novation agreement is that—

(1) The existing contract becomes invalid; and

(2) If the new owner wishes to participate in the Medicare program, it must apply for, and enter into, a contract in accordance with subpart K of this part.

(e) Effect of change of ownership with novation agreement. If the M+C organization submits a novation agreement that meets the requirements of § 422.552, and HCFA signs it, the new owner becomes the successor in interest to the current owner's Medicare contract.


§ 422.552 Novation agreement requirements.

(a) Conditions for HCFA approval of a novation agreement. HCFA approves a novation agreement if the following conditions are met:

(1) Advance notification. The M+C organization notifies HCFA at least 60 days before the date of the proposed change of ownership. The M+C organization also provides HCFA with updated financial information and a discussion of the financial and solvency impact of the change of ownership on the surviving organization.

(2) Advance submittal of agreement. The M+C organization submits to HCFA, at least 30 days before the proposed change of ownership date, three signed copies of the novation agreement containing the provisions specified in paragraph (b) of this section, and one copy of other relevant documents required by HCFA.

(3) HCFA's determination. HCFA determines that—

(i) The proposed new owner is in fact a successor in interest to the contract;

(ii) Recognition of the new owner as a successor in interest to the contract is in the best interest of the Medicare program; and

(iii) The successor organization meets the requirements to qualify as an M+C organization under subpart J of this part.

(b) Provisions of a novation agreement.

(1) Assumption of contract obligations. The new owner must assume all obligations under the contract.

(2) Waiver of right to reimbursement. The previous owner must waive its rights to reimbursement for covered services furnished during the rest of the current contract period.

(3) Guarantee of performance. (i) The previous owner must guarantee performance of the contract by the new owner during the contract period; or

(ii) The new owner must post a performance bond that is satisfactory to HCFA.

(4) Records access. The previous owner must agree to make its books and records and other necessary information available to the new owner and to HCFA to permit an accurate determination of costs for the final settlement of the contract period.

§ 422.553 Effect of leasing of an M+C organization's facilities.

(a) General effect of leasing. If an M+C organization leases all or part of its facilities to another entity, the other entity does not acquire M+C organization status under section 1876 of the Act.

(b) Effect of lease of all facilities. (1) If an M+C organization leases all of its facilities to another entity, the contract terminates.

(2) If the other entity wishes to participate in Medicare as an M+C organization, it must apply for and enter into a contract in accordance with subpart L of this part.

(c) Effect of partial lease of facilities. If the M+C organization leases part of its facilities to another entity, its contract with HCFA remains in effect while HCFA surveys the M+C organization to determine whether it continues to be in compliance with the applicable requirements and qualifying conditions specified in subpart K of this part.


Subpart M—Grievances, Organization Determinations and Appeals

SOURCE: 63 FR 35107, June 26, 1998, unless otherwise noted.

§ 422.560 Basis and scope.

(a) Statutory basis. (1) Section 1852(f) of the Act provides that an M+C organization must establish meaningful grievance procedures.

(2) Section 1852(g) of the Act establishes requirements that an M+C organization must meet concerning organization determinations and appeals.

(b) Scope. This subpart sets forth—

(1) Requirements for M+C organizations with respect to grievance procedures, organization determinations, and appeal procedures.

(2) The rights of M+C enrollees with respect to organization determinations, and grievance and appeal procedures.

(3) The rules concerning notice of noncoverage of inpatient hospital care.

(4) The rules that apply when an M+C enrollee requests immediate PRO review of a determination that he or she no longer needs inpatient hospital care.

§ 422.561 Definitions.

As used in this subpart, unless the context indicates otherwise—

Appeal means any of the procedures that deal with the review of adverse organization determinations on the health care services the enrollee believes he or she is entitled to receive, including delay in providing, arranging for, or approving the health care services (such that a delay would adversely affect the health of the enrollee), or on any amounts the enrollee must pay for a service, as defined under §422.566(b).

These procedures include reconsiderations by the M+C organization, and if necessary, an independent review entity, hearings before ALJs, review by the Departmental Appeals Board (DAB), and judicial review.

Authorized representative means an individual authorized by an enrollee, or under State law, to act on his or her behalf in obtaining an organization determination or in dealing with any of the levels of the appeal process, subject to the rules described in 20 CFR part 404, subpart R, unless otherwise stated in this subpart.

Enrollee means an M+C eligible individual who has elected an M+C plan offered by an M+C organization, or his or her authorized representative.

Grievance means any complaint or dispute other than one involving an organization determination, as defined in §422.566(b).

Physician has the meaning given the term in section 1861(r) of the Act.

[63 FR 35067, June 26, 1998, as amended at 65 FR 40328, June 29, 2000]

§ 422.562 General provisions.

(a) Responsibilities of the M+C organization. (1) An M+C organization, with respect to each M+C plan that it offers, must establish and maintain—

(i) A grievance procedure as described in §422.564 for addressing issues that do not involve organization determinations;

(ii) A procedure for making timely organization determinations;

...
§ 422.564 Grievance procedures.

(a) General rules. (1) Each M+C organization must provide meaningful procedures for timely hearing and resolution of grievances between enrollees and the organization or any other entity or individual through which the organization provides health care services, the M+C organization is ultimately responsible for ensuring that the entity or individual satisfies the relevant requirements of this subpart.

(b) Rights of M+C enrollees. In accordance with the provisions of this subpart, enrollees have the following rights:

(1) The right to have grievances between the enrollee and the M+C organization heard and resolved, as described in §422.564.

(2) The right to a timely organization determination, as provided under §422.566.

(3) The right to request an expedited organization determination, as provided under §422.570.

(4) If dissatisfied with any part of an organization determination, the following appeal rights:

(i) The right to a reconsideration of the adverse organization determination by the M+C organization, as provided under §422.578.

(ii) The right to request an expedited reconsideration, as provided under §422.584.

(iii) If, as a result of a reconsideration, an M+C organization affirms, in whole or in part, its adverse organization determination, the right to an automatic reconsidered determination made by an independent, outside entity contracted by HCFA, as provided in §422.592.

(iv) The right to an ALJ hearing if the amount in controversy is $100 or more, as provided in §422.600.

(v) The right to request DAB review of the ALJ hearing decision, as provided in §422.608.

(vi) The right to judicial review of the hearing decision if the amount in controversy is $1000 or more, as provided in §422.612.

(c) Limits on when this subpart applies. (1) If an enrollee receives immediate PRO review (as provided in §422.622) of a determination of noncoverage of inpatient hospital care—

(i) The enrollee is not entitled to review of that issue by the M+C organization; and

(ii) The PRO review decision is subject only to the appeal procedures set forth in part 473 of this chapter.

(2) If an enrollee has no further liability to pay for services that were furnished by an M+C organization, a determination regarding these services is not subject to appeal.

(d) When other regulations apply. Unless this subpart provides otherwise, the regulations in 20 CFR, part 404, subparts J and R (covering, respectively, the administrative review and hearing process and representation of parties under title II of the Act), apply under this subpart to the extent they are appropriate.

[63 FR 35067, June 26, 1998, as amended at 65 FR 40329, June 29, 2000]
§ 422.566 Organization determinations.

(a) Responsibilities of the M+C organization. Each M+C organization must have a procedure for making timely organization determinations (in accordance with the requirements of this subpart) regarding the benefits an enrollee is entitled to receive under an M+C plan, including basic benefits as described under §422.100(c)(1) and mandatory and optional supplemental benefits as described under §422.102, and the amount, if any, that the enrollee is required to pay for a health service. The M+C organization must have a standard procedure for making determinations, in accordance with §422.568, and an expedited procedure for situations in which applying the standard procedure could seriously jeopardize the enrollee's life, health, or ability to regain maximum function, in accordance with §§422.570 and 422.572.

(b) Actions that are organization determinations. An organization determination is any determination made by an M+C organization with respect to any of the following:

(1) Payment for temporarily out of the area renal dialysis services, emergency services, post-stabilization care, or urgently needed services.

(2) Payment for any other health services furnished by a provider other than the M+C organization that the enrollee believes—

(i) Are covered under Medicare; or

(ii) If not covered under Medicare, should have been furnished, arranged for, or reimbursed by the M+C organization.

(3) The M+C organization's refusal to provide or pay for services, in whole or in part, including the type or level of services, that the enrollee believes should be furnished or arranged for by the M+C organization.

(4) Discontinuation of a service if the enrollee believes that continuation of the services is medically necessary.

(5) Failure of the M+C organization to approve, furnish, arrange for, or provide payment for health care services in a timely manner, or to provide the enrollee with timely notice of an adverse determination, such that a delay would adversely affect the health of the enrollee.

(c) Who can request an organization determination. Any of the parties listed in §422.574 can request an organization determination, with the exception that only the parties listed in §422.570(a) can request an expedited determination.

[63 FR 35067, June 26, 1998, as amended at 65 FR 40329, June 29, 2000]

§ 422.568 Standard timeframes and notice requirements for organization determinations.

(a) Timeframe for requests for service. When a party has made a request for a service, the M+C organization must notify the enrollee of its determination as expeditiously as the enrollee's health condition requires, but no later than 14 calendar days after the date the organization receives the request for a standard organization determination. The M+C organization may extend the timeframe by up to 14 calendar days if the enrollee requests the extension or if the organization justifies a need for additional information and how the delay is in the interest of the enrollee (for example, the receipt of additional medical evidence from noncontract providers may change an M+C organization's decision to deny). When the M+C organization extends the timeframe, it must notify the enrollee in writing of the reasons for the delay, and inform the enrollee of the right to file a grievance if he or she disagrees with the M+C organization's decision to grant an extension. The M+C organization must notify the enrollee of its determination as expeditiously as the enrollee's health condition requires, but no later than upon expiration of the extension.

(b) Timeframe for requests for payment. The M+C organization must process requests for payment according to the "prompt payment" provisions set forth in §422.520.

(c) Written notification by practitioners. At each patient encounter with an M+C enrollee, a practitioner must notify the enrollee of his or her right to receive, upon request, a detailed written notice from the M+C organization regarding the enrollee's services, consistent with paragraph (d) of this section. The practitioner's notification must—
§ 422.570 Expediting certain organization determinations.

(a) Request for expedited determination. An enrollee or a physician (regardless of whether the physician is affiliated with the M+C organization) may request that an M+C organization expedite an organization determination involving the issues described in § 422.566(b)(3) and (b)(4). (This does not include requests for payment of services already furnished.)

(b) How to make a request. (1) To ask for an expedited determination, an enrollee or a physician must submit an oral or written request directly to the M+C organization or, if applicable, to the entity responsible for making the determination, as directed by the M+C organization.

(2) A physician may provide oral or written support for a request for an expedited determination.

(c) How the M+C organization must process requests. The M+C organization must establish and maintain the following procedures for processing requests for expedited determinations:

(i) Establish an efficient and convenient means for individuals to submit oral or written requests. The M+C organization must document all oral requests in writing and maintain the documentation in the case file.

(ii) Promptly decide whether to expedite a determination, based on the following requirements:

(a) For a request made by an enrollee, the M+C organization must provide an expedited determination if it determines that applying the standard timeframe for making a determination could seriously jeopardize the life or health of the enrollee.

(b) For a request made or supported by a physician, the M+C organization must provide an expedited determination if the physician indicates that applying the standard timeframe for making a determination could seriously jeopardize the life or health of the enrollee.

(d) Actions following denial. If an M+C organization denies a request for expedited determination, it must take the following actions:

(i) Automatically transfer a request to the standard timeframe and make the determination within the 14-day timeframe established in § 422.568 for a standard determination. The 14-day period begins with the day the M+C organization receives the request for expedited determination.

(ii) Give the enrollee prompt oral notice of the denial and subsequently deliver, within 3 calendar days, a written letter that—

(a) Explains that the M+C organization will process the request using the...
§ 422.572 Timeframes and notice requirements for expedited organization determinations.

(a) Timeframe. Except as provided in paragraph (b) of this section, an M+C organization that approves a request for expedited determination must make its determination and notify the enrollee (and the physician involved, as appropriate) of its decision within 72 hours after receiving the request.

(b) Extensions. The M+C organization may extend the 72-hour deadline by up to 14 calendar days if the enrollee requests the extension or if the organization justifies a need for additional information and how the delay is in the interest of the enrollee (for example, the receipt of additional medical evidence from noncontract providers may change an M+C organization’s decision to deny). When the M+C organization extends the deadline, it must notify the enrollee in writing of the reasons for the delay and inform the enrollee of the right to file a grievance if he or she disagrees with the M+C organization’s decision to grant an extension. The M+C organization must notify the enrollee of its determination as expeditiously as the enrollee’s health condition requires, but no later than upon expiration of the extension.

(c) Confirmation of oral notice. If the M+C organization first notifies an enrollee of its expedited determination orally, it must mail written confirmation to the enrollee within 3 calendar days of the oral notification.

(d) How the M+C organization must request information from noncontract providers. If the M+C organization must receive medical information from noncontract providers, the M+C organization must request the necessary information from the noncontract provider within 24 hours of the initial request for an expedited organization determination. Noncontract providers must make reasonable and diligent efforts to expeditiously gather and forward all necessary information to assist the M+C organization in meeting the required timeframe. Regardless of whether the M+C organization must request information from noncontract providers, the M+C organization is responsible for meeting the timeframe and notice requirements of this section.

(e) Content of the notice of expedited determination. (1) The notice of any expedited determination must state the specific reasons for the determination in understandable language.

(ii) Describe both the standard and expedited reconsideration processes, including the enrollee’s right to request, and conditions for obtaining, an expedited reconsideration, and the reasons for the delay and inform the enrollee of the right to file a grievance if he or she disagrees with the M+C organization’s decision to grant an extension. The M+C organization must notify the enrollee of its determination as expeditiously as the enrollee’s health condition requires, but no later than upon expiration of the extension.

(c) Confirmation of oral notice. If the M+C organization first notifies an enrollee of its expedited determination orally, it must mail written confirmation to the enrollee within 3 calendar days of the oral notification.

(d) How the M+C organization must request information from noncontract providers. If the M+C organization must receive medical information from noncontract providers, the M+C organization must request the necessary information from the noncontract provider within 24 hours of the initial request for an expedited organization determination. Noncontract providers must make reasonable and diligent efforts to expeditiously gather and forward all necessary information to assist the M+C organization in meeting the required timeframe. Regardless of whether the M+C organization must request information from noncontract providers, the M+C organization is responsible for meeting the timeframe and notice requirements of this section.

(e) Content of the notice of expedited determination. (1) The notice of any expedited determination must state the specific reasons for the determination in understandable language.

(ii) Describe both the standard and expedited reconsideration processes, including the enrollee’s right to request, and conditions for obtaining, an expedited reconsideration, and the rest of the appeal process; and

(iii) Comply with any other requirements specified by HCFA.

(f) Effect of failure to provide a timely notice. If the M+C organization fails to provide the enrollee with timely notice of an expedited organization determination as specified in this section, this failure itself constitutes an adverse organization determination and may be appealed.

[63 FR 35107, June 26, 1998, as amended at 65 FR 40329, June 29, 2000]
§ 422.574 Parties to the organization determination.

The parties to the organization determination are—
(a) The enrollee (including his or her authorized representative);
(b) An assignee of the enrollee (that is, a physician or other provider who has furnished a service to the enrollee and formally agrees to waive any right to payment from the enrollee for that service);
(c) The legal representative of a deceased enrollee’s estate; or
(d) Any other provider or entity (other than the M+C organization) determined to have an appealable interest in the proceeding.

§ 422.576 Effect of an organization determination.

The organization determination is binding on all parties unless it is reconsidered under §§ 422.578 through 422.596 or is reopened and revised under § 422.616.

§ 422.578 Right to a reconsideration.

Any party to an organization determination (including one that has been reopened and revised as described in § 422.616) may request that the determination be reconsidered under the procedures described in § 422.582, which address requests for a standard reconsideration. An enrollee or physician (acting on behalf of an enrollee) may request an expedited reconsideration as described in § 422.584.

§ 422.580 Reconsideration defined.

A reconsideration consists of a review of an adverse organization determination, the evidence and findings upon which it was based, and any other evidence the parties submit or the M+C organization or HCFA obtains.

§ 422.582 Request for a standard reconsideration.

(a) Method and place for filing a request. A party to an organization determination must ask for a reconsideration of the determination by filing a written request with—
(1) The M+C organization that made the organization determination; or
(2) An SSA office; or
(3) In the case of a qualified railroad retirement beneficiary, an RRB office.
(b) Timeframe for filing a request. Except as provided in paragraph (c) of this section, a party must file a request for a reconsideration within 60 calendar days from the date of the notice of the organization determination. If the SSA or RRB receives a request, it forwards the request to the M+C organization for its reconsideration. The timeframe within which the organization must conduct its review begins when it receives the request.
(c) Extending the time for filing a request. (1) General rule. If a party shows good cause, the M+C organization may extend the timeframe for filing a request for reconsideration.
(2) How to request an extension of timeframe. If the 60-day period in which to file a request for a reconsideration has expired, a party to the organization determination may file a request for reconsideration with the M+C organization, SSA, or an RRB office. If SSA or RRB receives a request, it forwards the request to the M+C organization for its reconsideration. The request for reconsideration and to extend the timeframe must—
(i) Be in writing; and
(ii) State why the request for reconsideration was not filed on time.
(d) Parties to the reconsideration. The parties to the reconsideration are the parties to the organization determination, as described in § 422.574, and any other provider or entity (other than the M+C organization) whose rights with respect to the organization determination may be affected by the reconsideration, as determined by the entity that conducts the reconsideration.
(e) Withdrawing a request. The party who files a request for reconsideration may withdraw it by filing a written request for withdrawal at one of the places listed in paragraph (a) of this section.

§ 422.584 Expediting certain reconsiderations.

(a) Who may request an expedited reconsideration. An enrollee or a physician (regardless of whether he or she is affiliated with the M+C organization) may request that an M+C organization
§ 422.590 Expedited reconsideration.

(a) Health Care Financing Administration, HHS § 422.590 expedite a reconsideration of a determination that involves the issues described in §422.566(b)(3) and (b)(4). (This does not include requests for payment of services already furnished.)

(b) How to make a request. (1) To make an expedited reconsideration, an enrollee or a physician acting on behalf of an enrollee must submit an oral or written request directly to the M+C organization or, if applicable, to the entity responsible for making the reconsideration, as directed by the M+C organization.

(2) A physician may provide oral or written support for a request for an expedited reconsideration.

(c) How the M+C organization must process requests. The M+C organization must establish and maintain the following procedures for processing requests for expedited reconsiderations:

(1) Handling of requests. The M+C organization must establish an efficient and convenient means for individuals to submit oral or written requests, document all oral requests in writing, and maintain the documentation in the case file.

(2) Prompt decision. Promptly decide on whether to expedite the reconsideration or follow the timeframe for standard reconsiderations:

(i) For a request made by an enrollee, the M+C organization must provide an expedited reconsideration if it determines that applying the standard timeframe for reconsidering a determination could seriously jeopardize the life or health of the enrollee or the enrollee's ability to regain maximum function.

(ii) For a request made or supported by a physician, the M+C organization must provide an expedited reconsideration if the physician indicates that applying the standard timeframe for conducting a reconsideration could seriously jeopardize the life or health of the enrollee or the enrollee's ability to regain maximum function.

(d) Actions following denial. If an M+C organization denies a request for expedited reconsideration, it must take the following actions:

(i) Automatically transfer a request to the standard timeframe and make the determination within the 30-day timeframe established in §422.590(a).

The 30-day period begins the day the M+C organization receives the request for expedited reconsideration.

(2) Give the enrollee prompt oral notice, and subsequently deliver, within 3 calendar days, a written letter that—

(i) Explains that the M+C organization will process the enrollee's request using the 30-day timeframe for standard reconsiderations;

(ii) Informs the enrollee of the right to file a grievance if he or she disagrees with the organization's decision not to expedite;

(iii) Informs the enrollee of the right to resubmit a request for an expedited reconsideration with any physician's support; and

(iv) Provides instructions about the grievance process and its timeframes.

(e) Action following acceptance of a request. If an M+C organization grants a request for expedited reconsideration, it must conduct the reconsideration and give notice in accordance with §422.590(d).

(f) Prohibition of punitive action. An M+C organization may not take or threaten to take any punitive action against a physician acting on behalf or in support of an enrollee in requesting an expedited reconsideration.

[63 FR 35107, June 26, 1998, as amended at 65 FR 40330, June 29, 2000]

§ 422.586 Opportunity to submit evidence.

The M+C organization must provide the parties to the reconsideration with a reasonable opportunity to present evidence and allegations of fact or law, related to the issue in dispute, in person as well as in writing. In the case of an expedited reconsideration, the opportunity to present evidence is limited by the short timeframe for making a decision. Therefore, the M+C organization must inform the parties of the conditions for submitting the evidence.

§ 422.590 Timeframes and responsibility for reconsiderations.

(a) Standard reconsideration: Request for services. (1) If the M+C organization makes a reconsidered determination that is completely favorable to the enrollee, the M+C organization must issue the determination (and effectuate
§422.590 42 CFR Ch. IV (10–1–00 Edition)

it in accordance with §422.618(a)) as expeditiously as the enrollee’s health condition requires, but no later than 30 calendar days from the date it receives the request for a standard reconsideration. The M+C organization may extend the timeframe by up to 14 calendar days if the enrollee requests the extension or if the organization justifies a need for additional information and how the delay is in the interest of the enrollee (for example, the receipt of additional medical evidence from noncontract providers may change an M+C organization’s decision to deny).

When the M+C organization extends the timeframe, it must notify the enrollee in writing of the reasons for the delay, and inform the enrollee of the right to file a grievance if he or she disagrees with the M+C organization’s decision to grant an extension. For extensions, the M+C organization must issue and effectuate its determination as expeditiously as the enrollee’s health condition requires, but no later than upon expiration of the extension.

(2) If the M+C organization makes a reconsidered determination that affirms, in whole or in part, its adverse organization determination, it must prepare a written explanation and send the case file to the independent entity contracted by HCFA as expeditiously as the enrollee’s health condition requires, but no later than 60 calendar days from the date it receives the request for a standard reconsideration.

(b) Standard reconsideration: Request for payment.

(1) If the M+C organization makes a reconsidered determination that is completely favorable to the enrollee, the M+C organization must issue its reconsidered determination to the enrollee (and effectuate it in accordance with §422.618(a)) no later than 60 calendar days from the date it receives the request for a standard reconsideration.

(2) If the M+C organization affirms, in whole or in part, its adverse organization determination, it must prepare a written explanation and send the case file to the independent entity contracted by HCFA no later than 60 calendar days from the date it receives the request for a standard reconsideration. The organization must make reasonable and diligent efforts to assist in gathering and forwarding information to the independent entity.

(c) Effect of failure to meet timeframe for standard reconsideration. If the M+C organization fails to provide the enrollee with a reconsidered determination within the timeframes specified in paragraph (a) or paragraph (b) of this section, this failure constitutes an affirmation of its adverse organization determination, and the M+C organization must submit the file to the independent entity in the same manner as described under paragraphs (a)(2) and (b)(2) of this section.

(d) Expedited reconsideration—(1) Timeframe. Except as provided in paragraph (d)(2) of this section, an M+C organization that approves a request for expedited reconsideration must complete its reconsideration and give the enrollee (and the physician involved, as appropriate) notice of its decision as expeditiously as the enrollee’s health condition requires but no later than 72 hours after receiving the request.

(2) Extensions. The M+C organization may extend the 72-hour deadline by up to 14 calendar days if the enrollee requests the extension or if the organization justifies a need for additional information and how the delay is in the interest of the enrollee (for example, the receipt of additional medical evidence from noncontract providers may change an M+C organization’s decision to deny). When the M+C organization extends the timeframe, it must notify the enrollee in writing of the reasons for the delay, and inform the enrollee of the right to file a grievance if he or she disagrees with the M+C organization’s decision to grant an extension.

The M+C organization must notify the enrollee of its determination as expeditiously as the enrollee’s health condition requires but no later than upon expiration of the extension.

(3) Confirmation of oral notice. If the M+C organization first notifies an enrollee of a completely favorable expedited reconsideration, it must mail
written confirmation to the enrollee within 3 calendar days.

(4) How the M+C organization must request information from noncontract providers. If the M+C organization must receive medical information from noncontract providers, the M+C organization must request the necessary information from the noncontract provider within 24 hours of the initial request for an expedited reconsideration. Noncontract providers must make reasonable and diligent efforts to expeditiously gather and forward all necessary information to assist the M+C organization in meeting the required timeframe. Regardless of whether the M+C organization must request information from noncontract providers, the M+C organization is responsible for meeting the timeframe and notice requirements.

(5) Affirmation of an adverse expedited organization determination. If, as a result of its reconsideration, the M+C organization affirms, in whole or in part, its adverse expedited organization determination, the M+C organization must submit a written explanation and the case file to the independent entity contracted by HCFA as expeditiously as the enrollee’s health condition requires, but not later than within 24 hours of its affirmation. The organization must make reasonable and diligent efforts to assist in gathering and forwarding information to the independent entity.

(e) Notification of enrollee. If the M+C organization refers the matter to the independent entity as described under this paragraph, it must concurrently notify the enrollee of that action.

(f) Failure to meet timeframe for expedited reconsideration. If the M+C organization fails to provide the enrollee with the results of its reconsideration within the timeframe described in paragraph (d) of this section, this failure constitutes an adverse reconsidered determination, and the M+C organization must submit the file to the independent entity within 24 hours of expiration of the timeframe set forth in paragraph (d) of this section.

(g) Who must reconsider an adverse organization determination. (1) A person or persons who were not involved in making the organization determination must conduct the reconsideration.

(2) When the issue is the M+C organization’s denial of coverage based on a lack of medical necessity (or any substantively equivalent term used to describe the concept of medical necessity), the reconsidered determination must be made by a physician with expertise in the field of medicine that is appropriate for the services at issue. The physician making the reconsidered determination need not, in all cases, be of the same specialty or subspecialty as the treating physician.

§ 422.592 Reconsideration by an independent entity.

(a) When the M+C organization affirms, in whole or in part, its adverse organization determination, the issues that remain in dispute must be reviewed and resolved by an independent, outside entity that contracts with HCFA.

(b) The independent outside entity must conduct the review as expeditiously as the enrollee’s health condition requires but must not exceed the deadlines specified in the contract.

(c) When the independent entity conducts a reconsideration, the parties to the reconsideration are the same parties listed in § 422.582(d) who qualified during the M+C organization’s reconsideration, with the addition of the M+C organization.

§ 422.594 Notice of reconsidered determination by the independent entity.

(a) Responsibility for the notice. When the independent entity makes the reconsidered determination, it is responsible for mailing a notice of its reconsidered determination to the parties and for sending a copy to HCFA.

(b) Content of the notice. The notice must—

(1) State the specific reasons for the entity’s decisions in understandable language;

(2) If the reconsidered determination is adverse (that is, does not completely reverse the M+C organization’s adverse organization determination), inform the parties of their right to an ALJ
hearing if the amount in controversy is $100 or more;
(3) Describe the procedures that a party must follow to obtain an ALJ hearing; and
(4) Comply with any other requirements specified by HCFA.

[63 FR 35107, June 26, 1998, as amended at 65 FR 40330, June 29, 2000]

§ 422.596 Effect of a reconsidered determination.

A reconsidered determination is final and binding on all parties unless a party other than the M+C organization files a request for a hearing under the provisions of § 422.602, or unless the reconsidered determination is revised under § 422.616.

[65 FR 40331, June 29, 2000]

§ 422.600 Right to a hearing.

(a) If the amount remaining in controversy is $100 or more, any party to the reconsideration (except the M+C organization) who is dissatisfied with the reconsidered determination has a right to a hearing before an ALJ. The M+C organization does not have the right to request a hearing before an ALJ.

(b) The amount remaining in controversy, which can include any combination of Part A and Part B services, is computed in accordance with § 405.740 of this chapter for Part A services and § 405.817 of this chapter for Part B services.

(c) If the basis for the appeal is the M+C organization’s refusal to provide services, HCFA uses the projected value of those services to compute the amount remaining in controversy.

§ 422.602 Request for an ALJ hearing.

(a) How and where to file a request. A party must file a written request for a hearing at one of the places listed in § 422.582(a) or with the independent, outside entity. The organizations listed in § 422.582(a) forward the request to the independent, outside entity, which is responsible for transferring the case to the appropriate ALJ hearing office.

(b) When to file a request. Except when an ALJ extends the timeframe as provided in 20 CFR 404.939(c), a party must file a request for a hearing within 60 days of the date of the notice of a reconsidered determination.

(c) Parties to a hearing. The parties to a hearing are the parties to the reconsideration, the M+C organization, and any other person or entity whose rights with respect to the reconsideration may be affected by the hearing, as determined by the ALJ.

(d) When the amount in controversy is less than $100. (1) If a request for a hearing clearly shows that the amount in controversy is less than $100, the ALJ dismisses the request.

(2) If, after a hearing is initiated, the ALJ finds that the amount in controversy is less than $100, he or she discontinues the hearing and does not rule on the substantive issues raised in the appeal.

§ 422.608 Departmental Appeals Board (the Board) review.

Any party to the hearing, including the M+C organization, who is dissatisfied with the ALJ hearing decision, may request that the Board review the ALJ’s decision or dismissal. Regulations located at 20 CFR 404.967 through 404.984 regarding SSA Appeals Council Review apply to Board review for matters addressed by this subpart.


§ 422.612 Judicial review.

(a) Review of ALJ’s decision. Any party, including the M+C organization, may request judicial review (upon notifying the other parties) of an ALJ’s decision if—

(1) The Board denied the party’s request for review; and

(2) The amount in controversy is $1,000 or more.

(b) Review of Board decision. Any party, including the M+C organization, may request judicial review (upon notifying the other parties) of the Board decision if it is the final decision of HCFA and the amount in controversy is $1,000 or more.

(c) How to request judicial review. A party must file a civil action in a district court of the United States in accordance with section 205(g) of the Act (see 20 CFR 422.210 for a description of
§ 422.616 Reopening and revising determination and decisions.

(a) An organization or reconsidered determination made by an M+C organization, a reconsidered determination made by the independent entity described in §422.592, or the decision of an ALJ or the Board that is otherwise final and binding may be reopened and revised by the entity that made the determination or decision, under the rules in §405.750 of this chapter.

(b) Reopening may be at the instigation of any party.

(c) The filing of a request for reopening does not relieve the M+C organization of its obligation to make payment or provide services as specified in §422.618.

(d) Once an entity issues a revised determination or decision, any party may file an appeal.


§ 422.618 How an M+C organization must effectuate standard reconsidered determinations or decisions.

(a) Reversals by the M+C organization—(1) Requests for service. If, on reconsideration of a request for service, the M+C organization completely reverses its organization determination, the organization must authorize or provide the service under dispute as expeditiously as the enrollee's health condition requires, but no later than 30 calendar days after the date the M+C organization receives the request for reconsideration (or no later than upon expiration of an extension described in §422.590(a)(1)).

(2) Requests for payment. If, on reconsideration of a request for payment, the M+C organization completely reverses its organization determination, the M+C organization must pay for the service no later than 60 calendar days after the date the M+C organization receives the request for reconsideration.

(b) Reversals by the independent outside entity. (1) Requests for service. If, on reconsideration of a request for service, the M+C organization's determination is reversed in whole or in part by the independent outside entity, the M+C organization must authorize the service under dispute within 72 hours from the date it receives notice reversing the determination, or provide the service under dispute as expeditiously as the enrollee's health condition requires, but no later than 14 calendar days from that date. The M+C organization must inform the independent outside entity that the organization has effectuated the decision.

(2) Requests for payment. If, on reconsideration of a request for payment, the M+C organization's determination is reversed in whole or in part by the independent outside entity, the M+C organization must pay for the service no later than 30 calendar days from the date it receives notice reversing the organization determination. The M+C organization must inform the independent outside entity that the organization has effectuated the decision.

(c) Reversals other than by the M+C organization or the independent outside entity. If the independent outside entity's determination is reversed in whole or in part by the ALJ, or at a higher level of appeal, the M+C organization must pay for, authorize, or provide the service under dispute as expeditiously as the enrollee's health condition requires, but no later than 60 calendar days from the date it receives notice reversing the determination. The M+C organization must inform the independent outside entity that the organization has effectuated the decision.

[63 FR 35107, June 26, 1998; as amended at 65 FR 40331, June 29, 2000]

§ 422.619 How an M+C organization must effectuate expedited reconsidered determinations.

(a) Reversals by the M+C organization. If on reconsideration of an expedited request for service, the M+C organization completely reverses its organization determination, the M+C organization must authorize or provide the service under dispute as expeditiously as the enrollee's health condition requires, but no later than 72 hours after the date the M+C organization receives the request for reconsideration (or no
§ 422.620 How enrollees of M+C organizations must be notified of noncoverage of inpatient hospital care.

(a) Enrollee's entitlement. Where an M+C organization has authorized coverage of the inpatient admission of an enrollee, either directly or by delegation (or the admission constitutes emergency or urgently needed care, as described in §§ 422.2 and 422.113), written notice of noncoverage under paragraph (c) of this section must be provided to each enrollee. An enrollee is entitled to coverage until at least noon the day after such notice is provided. If PRO review is requested under § 422.622, coverage is extended as provided in that section.

(b) Physician concurrence required. Before notice of noncoverage is provided as described in paragraph (c) of this section, the entity that makes the noncoverage/discharge determination (that is, the hospital by delegation or the M+C organization) must obtain the concurrence of the physician who is responsible for the enrollee's hospital care.

(c) Notice to the enrollee. In all cases in which a determination is made that inpatient hospital care is no longer necessary, no later than the day before hospital coverage ends, written notice must be provided to the enrollee that includes the following elements:

(1) The reason why inpatient hospital care is no longer needed.

(2) The effective date and time of the enrollee's liability for continued inpatient care.

(3) The enrollee's appeal rights.

(4) Additional information specified by HCFA.

[65 FR 40331, June 29, 2000]

§ 422.622 Requesting immediate PRO review of noncoverage of inpatient hospital care.

(a) Enrollee's right to review or reconsideration. (1) An enrollee who wishes to appeal a determination by an M+C organization or hospital that inpatient care is no longer necessary must request immediate PRO review of the determination in accordance with paragraph (b) of this section. An enrollee who requests immediate PRO review may remain in the hospital with no additional financial liability as specified in paragraph (c) of this section.

(2) An enrollee who fails to request immediate PRO review in accordance with the procedures in paragraph (b) of this section may request expedited reconsideration by the M+C organization as described in § 422.584, but the financial liability rules of paragraph (c) of this section do not apply.

(b) Procedures enrollee must follow. For the immediate PRO review process, the following rules apply:

(1) The enrollee must submit the request for immediate review—

(i) To the PRO that has an agreement with the hospital under § 466.78 of this chapter;

(ii) In writing or by telephone; and

(iii) By noon of the first working day after he or she receives written notice that the M+C organization or hospital has determined that the hospital stay is no longer necessary.

(2) On the date it receives the enrollee's request, the PRO must notify the
§ 422.644 Notice of contract determination.

(a) When HCFA makes a contract determination, it gives the M+C organization written notice.

(b) The notice specifies—

(1) The reasons for the determination; and

(2) The M+C organization's right to request reconsideration.

(c) For HCFA-initiated terminations, HCFA mails notice 90 days before the anticipated effective date of the termination. For terminations based on initial determinations described at

§ 422.643 Liability for hospital costs.

(a) When the M+C organization determines that hospital services are not, or are no longer, covered.

(i) Except as provided in paragraph (c)(1)(ii) of this section, if the M+C organization authorized coverage of the inpatient admission directly or by delegation (or the admission constitutes emergency or urgently needed care, as described in §§422.2 and 422.112(c)), the organization continues to be financially responsible for the costs of the hospital stay when a timely appeal is filed under paragraph (a)(1) of this section until noon of the calendar day following the day the PRO notifies the enrollee of its review determination.

(ii) The hospital may not charge the M+C organization (or the enrollee) if—

(A) It was the hospital (acting on behalf of the enrollee) that filed the request for immediate PRO review; and

(B) The PRO upholds the noncoverage determination made by the M+C organization.

(2) When the hospital determines that hospital services are no longer required. If the hospital determines that inpatient hospital services are no longer necessary, and the enrollee could not reasonably be expected to know that the services would not be covered, the hospital may not charge the enrollee for inpatient services received before noon of the calendar day following the day the PRO notifies the enrollee of its review determination.
§ 422.646

§ 422.510(a)(5), HCFA immediately notifies the M+C organization of its decision to terminate the organization’s M+C contract.

(d) When HCFA determines that it will not authorize a contract renewal, HCFA mails the notice to the M+C organization by May 1 of the current contract year.

§ 422.646 Effect of contract determination.

The contract determination is final and binding unless—

(a) The determination is reconsidered in accordance with §§422.648 through 422.658;

(b) A timely request for a hearing is filed under §422.662; or

(c) The reconsideration decision is revised as a result of a reopening under §422.696.

§ 422.648 Reconsideration: Applicability.

(a) Reconsideration is the first step for appealing a contract determination specified in §422.641.

(b) HCFA reconsiders the specified determinations if the contract applicant or the M+C organization files a written request in accordance with §422.650.

[63 FR 35113, June 26, 1998, as amended at 65 FR 40331, June 29, 2000]

§ 422.650 Request for reconsideration.

(a) Method and place for filing a request. A request for reconsideration must be made in writing and filed with any HCFA office.

(b) Time for filing a request. The request for reconsideration must be filed within 15 days from the date of the notice of the initial determination.

(c) Proper party to file a request. Only an authorized official of the contract applicant or M+C organization that was the subject of a contract determination may file the request for reconsideration.

(d) Withdrawal of a request. The M+C organization or contract applicant who filed the request for a reconsideration may withdraw it at any time before the notice of the reconsidered determination is mailed. The request for withdrawal must be in writing and filed with HCFA.

[63 FR 35113, June 26, 1998, as amended at 65 FR 40331, June 29, 2000]

§ 422.652 Opportunity to submit evidence.

HCFA provides the M+C organization or contract applicant and the HCFA official or officials who made the contract determination reasonable opportunity, not to exceed the timeframe in which an M+C organization could choose to request a hearing as described at §422.662, to present as evidence any documents or written statements that are relevant and material to the matters at issue.

[65 FR 40332, June 29, 2000]

§ 422.654 Reconsidered determination.

A reconsidered determination is a new determination that—

(a) Is based on a review of the contract determination, the evidence and findings upon which that was based, and any other written evidence submitted before notice of the reconsidered determination is mailed, including facts relating to the status of the M+C organization subsequent to the contract determination; and

(b) Affirms, reverses, or modifies the initial determination.

[63 FR 35113, June 26, 1998, as amended at 65 FR 40331, June 29, 2000]

§ 422.656 Notice of reconsidered determination.

(a) HCFA gives the M+C organization or contract applicant written notice of the reconsidered determination.

(b) The notice—

(1) Contains findings with respect to the contract applicant’s qualifications to enter into, or the M+C organization’s qualifications to remain under, a contract with HCFA under Part C of title XVIII of the Act;

(2) States the specific reasons for the reconsidered determination; and

(3) Informs the M+C organization or contract applicant of its right to a hearing if it is dissatisfied with the determination.

[63 FR 35113, June 26, 1998, as amended at 65 FR 40332, June 29, 2000]
§ 422.650 Right to a hearing.

The following parties are entitled to a hearing:

(a) A contract applicant that has been determined in a reconsidered determination to be unqualified to enter into a contract with HCFA under Part C of title XVIII of the Act.

(b) An M+C organization whose contract with HCFA has been terminated or has not been renewed as a result of a contract determination as provided in § 422.641.

[63 FR 35113, June 26, 1998, as amended at 65 FR 40332, June 29, 2000]

§ 422.662 Request for hearing.

(a) Method and place for filing a request. A request for a hearing must be made in writing and filed by an authorized official of the contract applicant or M+C organization that was the party to the determination under appeal. The request for a hearing must be filed with any HCFA office.

(b) Time for filing a request. A request for a hearing must be filed within 15 days after the date of the reconsidered determination.

(c) Parties to a hearing. The parties to a hearing must be—

(1) The parties described in § 422.660;

(2) At the discretion of the hearing officer, any interested parties who make a showing that their rights may be prejudiced by the decision to be rendered at the hearing; and

(3) HCFA.

[63 FR 35113, June 26, 1998, as amended at 65 FR 40332, June 29, 2000]

§ 422.664 Postponement of effective date of a contract determination when a request for a hearing with respect to a contract determination is filed timely.

(a) HCFA postpones the proposed effective date of the contract determination to terminate a contract with an M+C organization until a hearing decision is reached and affirmed by the Administrator following review under § 422.692 in instances where an M+C organization requests review by the Administrator; and

(b) HCFA extends the current contract at the end of the contract period (in the case of a determination not to renew) only—

(1) If HCFA finds that an extension of the contract will be consistent with the purpose of this part; and

(2) For such period as HCFA and the M+C organization agree.

(c) Exception: A contract terminated in accordance with § 422.510(a)(5) will be immediately terminated and will not be postponed if a hearing is requested.

§ 422.666 Designation of hearing officer.

HCFA designates a hearing officer to conduct the hearing. The hearing officer need not be an ALJ.

§ 422.668 Disqualification of hearing officer.

(a) A hearing officer may not conduct a hearing in a case in which he or she is prejudiced or partial to any party or has any interest in the matter pending for decision.

(b) A party to the hearing who objects to the designated hearing officer must notify that officer in writing at the earliest opportunity.

(c) The hearing officer must consider the objections, and may, at his or her discretion, either proceed with the hearing or withdraw.

(1) If the hearing officer withdraws, HCFA designates another hearing officer to conduct the hearing.

(2) If the hearing officer does not withdraw, the objecting party may, after the hearing, present objections and request that the officer's decision be revised or a new hearing be held before another hearing officer. The objections must be submitted in writing to HCFA.

§ 422.670 Time and place of hearing.

(a) The hearing officer fixes a time and place for the hearing, which is not to exceed 30 days from the receipt of the request for the hearing, and sends
§ 422.672 Written notice to the parties. The notice also informs the parties of the general and specific issues to be resolved and information about the hearing procedure.

(b) The hearing officer may, on his or her own motion, or at the request of a party, change the time and place for the hearing. The hearing officer may adjourn or postpone the hearing.

(c) The hearing officer will give the parties reasonable notice of any change in time or place of hearing, or of adjournment or postponement.

§ 422.672 Appointment of representatives.

A party may appoint as its representative at the hearing anyone not disqualified or suspended from acting as a representative before the Secretary or otherwise prohibited by law.

§ 422.674 Authority of representatives.

(a) A representative appointed and qualified in accordance with §422.672 may, on behalf of the represented party—

(1) Gives or accepts any notice or request pertinent to the proceedings set forth in this subpart;

(2) Presents evidence and allegations as to facts and law in any proceedings affecting that party; and

(3) Obtains information to the same extent as the party.

(b) A notice or request sent to the representative has the same force and effect as if it had been sent to the party.

§ 422.676 Conduct of hearing.

(a) The hearing is open to the parties and to the public.

(b) The hearing officer inquires fully into all the matters at issue and receives in evidence the testimony of witnesses and any documents that are relevant and material.

(c) The hearing officer provides the parties an opportunity to enter any objection to the inclusion of any document.

(d) The hearing officer decides the order in which the evidence and the arguments of the parties are presented and the conduct of the hearing.

§ 422.678 Evidence.

The hearing officer rules on the admissibility of evidence and may admit evidence that would be inadmissible under rules applicable to court procedures.

§ 422.680 Witnesses.

(a) The hearing officer may examine the witnesses.

(b) The parties or their representatives are permitted to examine their witnesses and cross-examine witnesses of other parties.

§ 422.682 Discovery.

(a) Prehearing discovery is permitted upon timely request of a party.

(b) A request is timely if it is made before the beginning of the hearing.

(c) A reasonable time for inspection and reproduction of documents is provided by order of the hearing officer.

(d) The hearing officer's order on all discovery matters is final.

§ 422.684 Prehearing.

The hearing officer may schedule a prehearing conference if he or she believes that a conference would more clearly define the issues.

§ 422.686 Record of hearing.

(a) A complete record of the proceedings at the hearing is made and transcribed and made available to all parties upon request.

(b) The record may not be closed until a hearing decision has been issued.

§ 422.688 Authority of hearing officer.

In exercising his or her authority, the hearing officer must comply with the provisions of title XVIII and related provisions of the Act, the regulations issued by the Secretary, and general instructions issued by HCFA in implementing the Act.

§ 422.690 Notice and effect of hearing decision.

(a) As soon as practical after the close of the hearing, the hearing officer issues a written decision that—

(1) Is based upon the evidence of record; and

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§ 422.752 Basis for imposing sanctions.

(a) All intermediate sanctions. For the violations listed below, HCFA may impose any of the sanctions specified in §422.750 on any M+C organization that has a contract in effect. The M+C organization may also be subject to other

Subpart O—Intermediate Sanctions

SOURCE: 63 FR 35115, June 26, 1998, unless otherwise noted.

§ 422.750 Kinds of sanctions.

(a) The following intermediate sanctions and civil money penalties may be imposed:

(1) Civil money penalties ranging from $10,000 to $100,000 depending upon the violation.

(2) Suspension of enrollment of Medicare beneficiaries.

(3) Suspension of payment to the M+C organization for Medicare beneficiaries who enroll.

(4) Require the M+C organization to suspend all marketing activities to Medicare beneficiaries for the M+C plan subject to the intermediate sanctions.

(b) The enrollment, payment, and marketing sanctions continue in effect until HCFA is satisfied that the deficiency on which the determination was based has been corrected and is not likely to recur.

§ 422.752 Basis for imposing sanctions.
§ 422.756适用的补救措施

(1) Fails substantially to provide, to an M+C enrollee, medically necessary services that the organization is required to provide (under law or under the contract) to an M+C enrollee, and that failure adversely affects (or is substantially likely to adversely affect) the enrollee.

(2) Imposes on M+C enrollees premiums in excess of the monthly basic and supplemental beneficiary premiums permitted under section 1854 of the Act and subpart G of this part.

(3) Expels or refuses to reenroll a beneficiary in violation of the provisions of this part.

(4) Engages in any practice that could reasonably be expected to have the effect of denying or discouraging enrollment of individuals whose medical condition or history indicates a need for substantial future medical services.

(5) Misrepresents or falsifies information that it furnishes—
   (i) To HCFA; or
   (ii) To an individual or to any other entity.

(6) Fails to comply with the requirements of §422.206, which prohibits interference with practitioners' advice to enrollees.

(7) Fails to comply with §422.216, which requires the organization to enforce the limit on balance billing under a private fee-for-service plan.

(8) Employs or contracts with an individual who is excluded from participation in Medicare under section 1128 or 1128A of the Act (or with an entity that employs or contracts with such an individual) for the provision of any of the following:
   (i) Health care.
   (ii) Utilization review.
   (iii) Medical social work.
   (iv) Administrative services.

(b) Suspension of enrollment and marketing. If HCFA makes a determination under §422.510(a), HCFA may impose the intermediate sanctions in §§422.756(c)(1) and (c)(3).

§ 422.756 Procedures for imposing sanctions.

(a) Notice of sanction and opportunity to respond—(1) Notice of sanction. Before imposing the intermediate sanctions specified in paragraph (c) of this section HCFA—
   (i) Sends a written notice to the M+C organization stating the nature and basis of the proposed sanction; and
   (ii) Sends the OIG a copy of the notice.

(2) Opportunity to respond. HCFA allows the M+C organization 15 days from receipt of the notice to provide evidence that it has not committed an act or failed to comply with the requirements described in §422.752, as applicable. HCFA may allow a 15-day addition to the original 15 days upon receipt of a written request from the M+C organization. To be approved, the request must provide a credible explanation of why additional time is necessary and be received by HCFA before the end of the 15-day period following the date of receipt of the sanction notice. HCFA does not grant an extension if it determines that the M+C organization's conduct poses a threat to an enrollee's health and safety.

(b) Informal reconsideration. If, consistent with paragraph (a)(2) of this section the M+C organization submits a timely response to HCFA's notice of sanction, HCFA conducts an informal reconsideration that:

(1) Consists of a review of the evidence by an HCFA official who did not participate in the initial decision to impose a sanction; and

(2) Gives the M+C organization a concise written decision setting forth the factual and legal basis for the decision that affirms or rescinds the original determination.

(c) Specific sanctions. If HCFA determines that an M+C organization has acted or failed to act as specified in §422.752 and affirms this determination in accordance with paragraph (b) of this section, HCFA may—

(1) Require the M+C organization to suspend acceptance of applications made by Medicare beneficiaries for enrollment in the sanctioned M+C plan during the sanction period;

(2) In the case of a violation under §422.752(a), suspend payments to the
M+C organization for Medicare beneficiaries enrolled in the sanctioned M+C plan during the sanction period; and

(3) Require the M+C organization to suspend all marketing activities for the sanctioned M+C plan to Medicare enrollees.

(d) Effective date and duration of sanctions—(1) Effective date. Except as provided in paragraph (d)(2) of this section, a sanction is effective 15 days after the date that the organization is notified of the decision to impose the sanction or, if the M+C organization timely seeks reconsideration under paragraph (b) of this section, on the date specified in the notice of HCFA’s reconsidered determination.

(2) Exception. If HCFA determines that the M+C organization’s conduct poses a serious threat to an enrollee’s health and safety, HCFA may make the sanction effective on a date before issuance of HCFA’s reconsidered determination.

(3) Duration of sanction. The sanction remains in effect until HCFA notifies the M+C organization that HCFA is satisfied that the basis for imposing the sanction has been corrected and is not likely to recur.

(e) Termination by HCFA. In addition to or as an alternative to the sanctions described in paragraph (c) of this section, HCFA may decline to authorize the renewal of an organization’s contract in accordance with §422.506(b)(2) and (b)(3), or terminate the contract in accordance with §422.510.

(f) Civil money penalties. (1) If HCFA determines that an M+C organization has committed an act or failed to comply with a requirement described in §422.752, HCFA notifies the OIG of this determination, and also notifies OIG when HCFA reverses or terminates a sanction imposed under this part.

(2) In the case of a violation described in paragraph (a) of §422.752, or a determination under paragraph (b) of §422.752 based upon a violation under §422.510(a)(4) (involving fraudulent or abusive activities), in accordance with the provisions of 42 CFR parts 1003 and 1005, HCFA may impose civil money penalties on the M+C organization in the amounts specified in §422.758 in addition to, or in place of, the sanctions that HCFA may impose under paragraph (c) of this section.

(3) In the case of a determination under paragraph (b) of §422.752 other than a determination based upon a violation under §422.510(a)(4), in accordance with the provisions of 42 CFR parts 1003 and 1005, HCFA may impose civil money penalties on the M+C organization in the amounts specified in §422.758 in addition to, or in place of, the sanctions that HCFA may impose under paragraph (c) of this section.

§422.758 Maximum amount of civil money penalties imposed by HCFA.

If HCFA makes a determination under §422.752(b), based on any determination under §422.510(a) except a determination under §422.510(a)(4), HCFA may impose civil money penalties in the following amounts:

(a) If the deficiency on which the determination is based has directly adversely affected (or has the substantial likelihood of adversely affecting) one or more M+C enrollees—$25,000 for each determination.

(b) For each week that a deficiency remains uncorrected after the week in which the M+C organization receives HCFA’s notice of the determination—$10,000.

§422.760 Other applicable provisions.

The provisions of section 1128A of the Act (except subsections (a) and (b)) apply to civil money penalties under this subpart to the same extent that they apply to a civil money penalty or procedure under section 1128A of the Act.

PART 424—CONDITIONS FOR MEDICARE PAYMENT

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§ 424.5 Basic conditions.

(a) As a basis for Medicare payment, the following conditions must be met:

(1) Types of services. The services must be—

(i) Covered services, as specified in part 409 or part 410 of this chapter; or

(ii) Services excluded from coverage as custodial care or services not reasonable and necessary, but reimbursable in accordance with §§405.332 through 405.334 of this chapter, pertaining to limitation of liability.

(2) Sources of services. The services must have been furnished by a provider, nonparticipating hospital, or supplier that was, at the time it furnished the services, qualified to have payment made for them.

(3) Recipient of services. Except as provided in §409.68 of this chapter, the services must have been furnished while the individual was eligible to have payment made for them. (Section 409.68 provides for payment of inpatient hospital services furnished before the hospital is notified that the beneficiary...

§ 424.7 General limitations.

(a) Utilization review finding on medical necessity. When a PRO or a UR committee notifies a hospital or SNF of its finding that further services are not medically necessary, the following rules apply:

(1) Hospitals subject to PPS. Payment may not be made for inpatient hospital services furnished by a PPS hospital after the second day after the day on which the hospital received the notice.

(2) Hospitals not subject to PPS and SNFs—(i) Basic rule. Except as provided in paragraph (a)(2)(ii) of this section, payment may not be made for inpatient hospital services or posthospital SNF care furnished after the day on which the hospital or SNF received the notice.

(ii) Exception. Payment may be made for 1 or 2 additional days if the PRO or UR committee approves them as necessary for planning for post-discharge care.

(b) Failure to make timely utilization review. Payment may not be made for inpatient hospital services or posthospital SNF care furnished, after the 20th consecutive day of a stay, to an individual who is admitted to the hospital or SNF after HCFA has determined that the hospital or SNF has failed to make timely utilization review in long stay cases. (This provision does not apply to a hospital or SNF for which a PRO has assumed binding review.)

[53 FR 6635, Mar. 2, 1988; 53 FR 12945, Apr. 20, 1988]

Subpart B—Certification and Plan of Treatment Requirements

§ 424.10 Purpose and scope.

(a) Purpose. The physician has a major role in determining utilization of health services furnished by providers. The physician decides upon admissions, orders tests, drugs, and treatments, and determines the length of stay. Accordingly, sections 1814(a)(2) and 1835(a)(2) of the Act establish as a condition for Medicare payment that a physician certify the necessity of the services and, in some instances, recertify the continued need for those services.

Section 1814(a)(2) of the Act also permits nurse practitioners or clinical nurse specialists to certify and recertify the need for post-hospital extended care services.

(b) Scope. This subpart sets forth the timing, content, and signature requirements for certification and recertification with respect to certain Medicare services furnished by providers.

[60 FR 38271, July 26, 1995]

§ 424.11 General procedures.

(a) Responsibility of the provider. The provider must—

(1) Obtain the required certification and recertification statements;

(2) Keep them on file for verification by the intermediary, if necessary; and

(3) Certify, on the appropriate billing form, that the statements have been obtained and are on file.

(b) Obtaining the certification and recertification statements. No specific procedures or forms are required for certification and recertification statements. The provider may adopt any method that permits verification. The certification and recertification statements may be entered on forms, notes, or records that the appropriate individual signs, or on a special separate
§ 424.13 Requirements for inpatient services of hospitals other than psychiatric hospitals.

(a) Content of certification and recertification. Medicare Part A pays for inpatient hospital services of hospitals...

(i) Be a registered professional nurse who is currently licensed to practice nursing in the State where he or she practices; be authorized to perform the services of a nurse practitioner in accordance with State law; and have a master’s degree in nursing;

(ii) Be certified as a nurse practitioner by a professional association recognized by HCFA that has, at a minimum, eligibility requirements that meet the standards in paragraph (e)(5)(i) of this section; or

(iii) Meet the requirements for a nurse practitioner set forth in paragraph (e)(5)(i) of this section, except for the master’s degree requirement, and have received before August 25, 1998 a certificate of completion from a formal advanced practice program that prepares registered nurses to perform an expanded role in the delivery of primary care.

(5) For purposes of this section, to qualify as a clinical nurse specialist, an individual must—

(i) Be a registered professional nurse who is currently licensed to practice nursing in the State where he or she practices; be authorized to perform the services of a clinical nurse specialist in accordance with State law; and have a master’s degree in a defined clinical area of nursing;

(ii) Be certified as a clinical nurse specialist by a professional association recognized by HCFA that has, at a minimum, eligibility requirements that meet the standards in paragraph (e)(6)(i) of this section; or

(iii) Meet the requirements for a clinical nurse specialist set forth in paragraph (e)(6)(i) of this section, except for the master’s degree requirement, and have received before August 25, 1998 a certificate of completion from a formal advanced practice program that prepares registered nurses to perform an expanded role in the delivery of primary care.
other than psychiatric hospitals only if a physician certifies and recertifies the following:

(1) The reasons for either—
   (i) Continued hospitalization of the patient for medical treatment or medically required inpatient diagnostic study; or
   (ii) Special or unusual services for cost outlier cases (under the prospective payment system set forth in subpart F of part 412 of this chapter).
(2) The estimated time the patient will need to remain in the hospital.
(3) The plans for posthospital care, if appropriate.

(b) Certification of need for hospitalization when a SNF bed is not available.

(1) A physician may certify or recertify need for continued hospitalization if the physician finds that the patient could receive proper treatment in a SNF but no bed is available in a participating SNF.

(2) If this is the basis for the physician’s certification or recertification, the required statement must so indicate; and the physician is expected to continue efforts to place the patient in a participating SNF as soon as a bed becomes available.

(c) Signatures.

(1) Basic rule. Except as specified in paragraph (c)(2) of this section, certifications and recertifications must be signed by the physician responsible for the case, or by another physician who has knowledge of the case and who is authorized to do so by the responsible physician or by the hospital's medical staff.

(2) Exception. If the intermediary requests certification of the need to admit a patient in connection with dental procedures, because his or her underlying medical condition and clinical status or the severity of the dental procedures require hospitalization, that certification may be signed by the dentist caring for the patient.

(d) Timing of certifications and recertifications. Cases not subject to the prospective payment system (PPS).

(1) For cases that are not subject to PPS, certification is required no later than as of the 12th day of hospitalization.

(2) The first recertification is required no later than as of the 18th day of hospitalization.

(3) Subsequent recertifications are required at intervals established by the UR committee (on a case-by-case basis if it so chooses), but no less frequently than every 30 days.

(e) Timing of certification and recertification: Cases subject to PPS. For cases subject to PPS, certification is required as follows:

(1) For day-outlier cases, certification is required no later than one day after the hospital reasonably assumes that the case meets the outlier criteria, established in accordance with §412.80(a)(1)(i) of this chapter, or no later than 20 days into the hospital stay, whichever is earlier. The first and subsequent recertifications are required at intervals established by the UR committee (on a case-by-case basis if it so chooses) but not less frequently than every 30 days.

(2) For cost-outlier cases, certification is required no later than the date on which the hospital requests cost outlier payment or 20 days into the hospital stay, whichever is earlier. If possible, certification must be made before the hospital incurs costs for which it will seek cost outlier payment. In cost outlier cases, the first and subsequent recertifications are required at intervals established by the UR committee (on a case-by-case basis if it so chooses).

(f) Recertification requirement fulfilled by utilization review.

(1) At the hospital’s option, extended stay review by its UR committee may take the place of the second and subsequent physician recertifications required for cases not subject to PPS and for PPS day-outlier cases.

(2) A utilization review that is used to fulfill the recertification requirement is considered timely if performed no later than the seventh day after the day the physician recertification would have been required. The next physician recertification would need to be made no later than the 30th day following such review; if review by the UR committee took the place of this physician recertification, the review could be
performed as late as the seventh day following the 30th day.

(g) Description of procedures. The hospital must have available on file a written description that specifies the time schedule for certifications and recertifications, and indicates whether utilization review of long-stay cases fulfills the requirement for second and subsequent recertifications of all cases not subject to PPS and of PPS day outlier cases.

§ 424.14 Requirements for inpatient services of psychiatric hospitals.

(a) Content of certification and recertification: General considerations. The content requirements differ from those for other hospitals because the care furnished in psychiatric hospitals is often purely custodial and thus not covered under Medicare. The purpose of the statements, therefore, is to help ensure that Medicare pays only for services of the type appropriate for Medicare coverage. Accordingly, Medicare Part A pays for inpatient care in a psychiatric hospital only if a physician certifies and recertifies the need for services consistent with the content of paragraphs (b) or (c) of this section, as appropriate.

(b) Content of certification. Inpatient psychiatric services were required—

(1) For treatment that could reasonably be expected to improve the patient’s condition; or

(2) For diagnostic study.

(c) Content of recertification. (1) Inpatient services furnished since the previous certification or recertification were, and continue to be, required—

(i) For treatment that could reasonably be expected to improve the patient’s condition; or

(ii) For diagnostic study; and

(2) The hospital records show that the services furnished were—

(i) Intensive treatment services;

(ii) Admission and related services necessary for diagnostic study; or

(iii) Equivalent services.

(d) Timing of certification and recertification. (1) Certification is required at the time of admission or as soon thereafter as is reasonable and practicable.

(2) The first recertification is required as of the 18th day of hospitalization. Subsequent recertifications are required at intervals established by the UR committee (on a case-by-case basis if it so chooses), but no less frequently than every 30 days.

(e) Other requirements. Psychiatric hospitals must also meet the requirements set forth in §424.13(b), (c), (f), and (g).

§ 424.15 Requirements for inpatient CAH services.

(a) Content of certification. Medicare Part A pays for inpatient CAH services only if a physician certifies that the individual may reasonably be expected to be discharged or transferred to a hospital within 96 hours after admission to the CAH.

(b) Timing of certification. Certification is required no later than 1 day before the date on which the claim for payment for the inpatient CAH services is submitted.

§ 424.16 Timing of certification for individual admitted to a hospital before entitlement to Medicare benefits.

(a) Basic rule. If an individual is admitted to a hospital before becoming entitled to Medicare benefits (for instance, before attaining age 65), the day of entitlement (instead of the day of admission) is the starting point for the time limits specified in §424.13(e) for certification and recertification.

(b) Example. (Hospital that is not a psychiatric hospital and is not subject to PPS). For a patient who is admitted on August 15 and becomes entitled on September 1—

(1) The certification is required no later than September 12;

(2) The first recertification is required no later than September 18; and

(3) Subsequent recertifications are required at least every 30 days after September 18.

§ 424.20 Requirements for posthospital SNF care.

Medicare Part A pays for posthospital SNF care furnished by an SNF, or a hospital or CAH with a
§ 424.20 swing-bed approval, only if the certification and recertification for services are consistent with the content of paragraph (a) or (c) of this section, as appropriate.

(a) Content of certification—(1) General requirements. Posthospital SNF care is or was required because—
   (i) The individual needs or needed on a daily basis skilled nursing care (furnished directly by or requiring the supervision of skilled nursing personnel) or other skilled rehabilitation services that, as a practical matter, can only be provided in an SNF or a swing-bed hospital on an inpatient basis, and the SNF care is or was needed for a condition for which the individual received inpatient care in a participating hospital or a qualified hospital, as defined in §409.3 of this chapter; or
   (ii) The individual has been correctly assigned to one of the Resource Utilization Groups designated as representing the required level of care, as provided in §409.30 of this chapter.

(2) Special requirement: A swing-bed hospital with more than 49 beds (but fewer than 100) that does not transfer a swing-bed patient to a SNF within 5 days of the availability date. Transfer of the extended care patient to the SNF is not medically appropriate.

(b) Timing of certification. (1) General rule. The certification must be obtained at the time of admission or as soon thereafter as is reasonable and practicable.

(2) Special rules for certain swing-bed hospitals. For swing-bed hospitals with more than 49 beds that are approved after March 31, 1988, the extended care patient’s physician has 5 days (excluding weekends and holidays) beginning on the availability date as defined in §413.114(b), to certify that the transfer of the extended care patient to the SNF is not medically appropriate.

(b) Timing of certification. (1) General rule. The certification must be obtained at the time of admission or as soon thereafter as is reasonable and practicable.

(2) Special rules for certain swing-bed hospitals. For swing-bed hospitals with more than 49 beds that are approved after March 31, 1988, the extended care patient’s physician has 5 days (excluding weekends and holidays) beginning on the availability date as defined in §413.114(b), to certify that the transfer of the extended care patient to the SNF is not medically appropriate.

(c) Content of recertifications. (1) The reasons for the continued need for posthospital SNF care;

(2) The estimated time the individual will need to remain in the SNF;

(3) Plans for home care, if any; and

(4) If appropriate, the fact that continued services are needed for a condition that arose after admission to the SNF and while the individual was still under treatment for the condition for which he or she had received inpatient hospital services.

(d) Timing of recertifications. (1) The first recertification is required no later than the 14th day of posthospital SNF care.

(2) Subsequent recertifications are required at least every 30 days after the first recertification.

(e) Signature. Certification and recertification statements may be signed by—

(1) The physician responsible for the case or, with his or her authorization, by a physician on the SNF staff or a physician who is available in case of an emergency and has knowledge of the case; or

(2) A nurse practitioner or clinical nurse specialist, neither of whom has a direct or indirect employment relationship with the facility but who is working in collaboration with a physician. For purposes of this section, collaboration means a process whereby a nurse practitioner or clinical nurse specialist works with a doctor of medicine or osteopathy to deliver health care services. The services are delivered within the scope of the nurse’s professional expertise, with medical direction and appropriate supervision as provided for in guidelines jointly developed by the nurse and the physician or other mechanisms defined by Federal regulations and the law of the State in which the services are performed.

(f) Recertification requirement fulfilled by utilization review. A SNF may substitute utilization review of extended stay cases for the second and subsequent recertifications, if it includes this procedure in its utilization review plan.

(g) Description of procedures. The SNF must have available on file a written description that specifies the certification and recertification time schedule and indicates whether utilization review is used as an alternative to the second and subsequent recertifications.
§ 424.22 Requirements for home health services.

Medicare Part A or Part B pays for home health services only if a physician certifies and recertifies the content specified in paragraphs (a)(1) and (b)(2) of this section, as appropriate.

(a) Certification—(1) Content of certification. As a condition for payment of home health services under Medicare Part A or Medicare Part B, a physician must certify as follows:

(i) The individual needs or needed intermittent skilled nursing care, or physical or speech therapy, or (for the period from July through November 30, 1981) occupational therapy.

(ii) Home health services were required because the individual was confined to the home except when receiving outpatient services.

(iii) A plan for furnishing the services has been established and is periodically reviewed by a physician who is a doctor of medicine, osteopathy, or podiatric medicine, and who is not precluded from performing this function under paragraph (d) of this section. (A doctor of podiatric medicine may perform only plan of treatment functions that are consistent with the functions he or she is authorized to perform under State law.)

(iv) The services were furnished while the individual was under the care of a physician who is a doctor of medicine, osteopathy, or podiatric medicine.

(2) Timing and signature. The certification of need for home health services must be obtained at the time the plan of treatment is established or as soon thereafter as possible and must be signed by the physician who establishes the plan.

(b) Recertification. (1) Timing and signature of recertification. Recertification is required at least every 60 days, preferably at the time the plan is reviewed, and must be signed by the physician who reviews the plan of care. The recertification is required at least every 60 days when there is a—

(i) Beneficiary elected transfer; or
(ii) Discharge and return to the same HHA during the 60-day episode.

(2) Content and basis of recertification. The recertification statement must indicate the continuing need for services and estimate how much longer the services will be required. Need for occupational therapy may be the basis for continuing services that were initiated because the individual needed skilled nursing care or physical or speech therapy.

(c) [Reserved]

(d) Limitations on the performance of certification and plan of treatment functions. (1) Basic rule. Beginning November 26, 1982, and except as provided in paragraph (e) of this section, need for home health services to be provided by an HHA may not be certified or recertified, and a plan of treatment may not be established and reviewed, by any physician who has a significant ownership interest in, or a significant financial or contractual relationship with, that HHA.

(2) Significant ownership interest. A physician is considered to have a significant ownership interest in an HHA if he or she—

(i) Has a direct or indirect ownership interest of 5 percent or more in the capital, the stock, or the profits of the home health agency; or
(ii) Has an ownership interest of 5 percent or more in any mortgage, deed of trust, note, or other obligation that is secured by the agency, if that interest equals 5 percent or more of the agency's assets.

(3) Significant financial or contractual relationship. Beginning November 26, 1982, a physician is considered to have a significant financial or contractual relationship with an HHA if he or she—

(i) Receives any compensation as an officer or director of the HHA; or
(ii) Has direct or indirect business transactions with the HHA that, in any fiscal year, amount to more than $25,000 or 5 percent of the agency's total operating expenses, whichever is less. Business transactions means contracts, agreements, purchase orders, or leases to obtain services, supplies, equipment, and space and, after August 29, 1986, salaried employment.

1As a condition of Medicare Part A payment for home health services furnished before July 1981, the physician was also required to certify that the services were needed for a condition for which the individual had received inpatient hospital or SNF services.
§ 424.24  Requirements for medical and other health services furnished by providers under Medicare Part B.

(a) Exempted services. Certification is not required for the following: (1) Hospital services and supplies incident to physicians’ services furnished to outpatients. The exemption applies to drugs and biologicals that cannot be self-administered, but not to partial hospitalization services, as set forth in paragraph (e) of this section.

(2) Outpatient hospital diagnostic services, including necessary drugs and biologicals, ordinarily furnished or arranged for by a hospital for the purpose of diagnostic study.

(b) General rule. Medicare Part B pays for medical and other health services furnished by providers (and not exempted under paragraph (a) of this section) only if a physician certifies the content specified in paragraph (c)(1), (c)(4) or (e)(1) of this section, as appropriate.

(c) Outpatient physical therapy and speech-language pathology services—(1) Content of certification. (i) The individual needs, or needed, physical therapy or speech pathology services.

(ii) The services were furnished while the individual was under the care of a physician, nurse practitioner, clinical nurse specialist, or physician assistant.

(iii) The services were furnished under a plan of treatment that meets the requirements of §410.61 of this chapter.

(2) Timing. The certification statement must be obtained at the time the plan of treatment is established, or as soon thereafter as possible.

(3) Signature. (i) If the plan of treatment is established by a physician, nurse practitioner, clinical nurse specialist, or physician assistant, the certification must be signed by that physician or nonphysician practitioner.

(ii) If the plan of treatment is established by a physical therapist or speech-language pathologist, the certification must be signed by a physician or by a nurse practitioner, clinical nurse specialist, or physician assistant who has knowledge of the case.

(4) Recertification—(i) Timing. Recertification statements are required at least every 30 days and must be signed by the physician, nurse practitioner, clinical nurse specialist, or physician assistant who reviews the plan of treatment.

(ii) Content. The recertification statement must indicate the continuing need for physical therapy or speech-language pathology services and an estimate of how much longer the services will be needed.
(iii) Signature. Recertifications must be signed by the physician, nurse practitioner, clinical nurse specialist, or physician assistant who reviews the plan of treatment.

(d) [Reserved]

(e) Partial hospitalization services: Content of certification and plan of treatment requirements—

(i) The individual would require inpatient psychiatric care if the partial hospitalization services were not provided.

(ii) The services are or were furnished while the individual was under the care of a physician.

(iii) The services were furnished under a written plan of treatment that meets the requirements of paragraph (e)(2) of this section.

(2) Plan of treatment requirements. (i) The plan is an individualized plan that is established and is periodically reviewed by a physician in consultation with appropriate staff participating in the program, and that sets forth—

(A) The physician’s diagnosis;

(B) The type, amount, duration, and frequency of the services; and

(C) The treatment goals under the plan.

(ii) The physician determines the frequency and duration of the services taking into account accepted norms of medical practice and a reasonable expectation of improvement in the patient’s condition.

(3) Recertification requirements.

(i) Signature. The physician recertification must be signed by a physician who is treating the patient and has knowledge of the patient’s response to treatment.

(ii) Timing. The first recertification is required as of the 18th day of partial hospitalization services. Subsequent recertifications are required at intervals established by the provider, but no less frequently than every 30 days.

(iii) Content. The recertification must specify that the patient would otherwise require inpatient psychiatric care in the absence of continued stay in the partial hospitalization program and describe the following:

(A) The patient’s response to the therapeutic interventions provided by the partial hospitalization program.

(B) The patient’s psychiatric symptoms that continue to place the patient at risk of hospitalization.

(C) Treatment goals for coordination of services to facilitate discharge from the partial hospitalization program.

(f) All other covered medical and other health services furnished by providers—

(1) Content of certification. The services were medically necessary.

(2) Signature. The certificate must be signed by a physician, nurse practitioner, clinical nurse specialist, or physician assistant who has knowledge of the case.

(3) Timing. The physician, nurse practitioner, clinical nurse specialist, or physician assistant may provide certification at the time the services are furnished or, if services are provided on a continuing basis, either at the beginning or at the end of a series of visits.

(4) Recertification. Recertification of continued need for services is not required.


§ 424.27 Requirements for comprehensive outpatient rehabilitation facility (CORF) services.

Medicare Part B pays for CORF services only if a physician certifies, and the facility physician recertifies, the content specified in paragraphs (a) and (b)(2) of this section, as appropriate.

(a) Certification. Content. (1) The services were required because the individual needed skilled rehabilitation services;

(2) The services were furnished while the individual was under the care of a physician; and

(3) A written plan of treatment has been established and is reviewed periodically by a physician.

(b) Recertification—(1) Timing. Recertification is required at least every 60 days, based on review by a facility physician who, when appropriate, consults with the professional personnel who furnish the services.

(2) Content. (i) The plan is being followed;

(ii) The patient is making progress in attaining the rehabilitation goals; and,

(iii) The treatment is not having any harmful effect on the patient.
§ 424.30 Scope.

This subpart sets forth the requirements, procedures, and time limits for claiming Medicare payments. Claims must be filed in all cases except when services are furnished on a prepaid capitation basis by a health maintenance organization (HMO), a competitive medical plan (CMP), or a health care prepayment plan (HCPP). Special procedures for claiming payment after the beneficiary has died and for certain bills paid by organizations are set forth in subpart E of this part.

§ 424.32 Basic requirements for all claims.

(a) A claim must meet the following requirements:
   (1) A claim must be filed with the appropriate intermediary or carrier on a form prescribed by HCFA in accordance with HCFA instructions.
   (2) A claim for physician services, clinical psychologist services, or clinical social worker services must include appropriate diagnostic coding for those services using ICD-9-CM, and a claim for physician services furnished to an SNF resident under §411.15(p)(2) of this chapter must also include the SNF’s Medicare provider number.
   (3) A claim must be signed by the beneficiary or the beneficiary’s representative (in accordance with §424.36(b)).
   (4) A claim must be filed within the time limits specified in §424.44.
   (5) A Part B claim filed by an SNF must include appropriate HCPCS coding.

(b) The prescribed forms for claims are the following:
   HCFA-1450—Uniform Institutional Provider Bill. (This form is for institutional provider billing for Medicare inpatient, outpatient and home health services.)
   HCFA-1490—Request for Medicare payment. (For use by a patient to request payment for medical expenses.)
   HCFA-1490U—Request for Medicare Payment by Organization. (For use by an organization requesting payment for medical services.)
   HCFA-1500—Health Insurance Claim Form. (For use by physicians and other suppliers to request payment for medical services.)
   HCFA-1660—Request for Information-Medicare Payment for Services to a Patient now Deceased. (For use in requesting amounts payable under title XVIII to a deceased beneficiary.)
   (c) Where claims forms are available. Excluding forms HCFA-1450 and HCFA-1500, all claims forms prescribed for use in the Medicare program are distributed free-of-charge to the public, institutions, or organizations. The HCFA-1490S is also available at local Social Security Offices.

§ 424.33 Additional requirements: Claims for services of providers and claims by suppliers and nonparticipating hospitals.

All claims for services of providers and all claims by suppliers and nonparticipating hospitals must be—
   (a) Filed by the provider, supplier, or hospital; and
   (b) Signed by the provider, supplier, or hospital unless HCFA instructions waive this requirement.

§ 424.34 Additional requirements: Beneficiary’s claim for direct payment.

(a) Basic rule. A beneficiary’s claim for direct payment for services furnished by a supplier, or by a nonparticipating hospital that has not elected to claim payment for emergency services, must include an itemized bill or a “report of services”, as specified in paragraphs (b) and (c) of this section.
   (b) Itemized bill from the hospital or supplier. The itemized bill for the services, which may be receipted or unpaid, must include all of the following information:
      (1) The name and address of—
§ 424.37 Evidence of authority to sign on behalf of the beneficiary.

(a) Beneficiary incapable. When a party specified in § 424.36(b) signs a claim or request for payment statement, he or she must also submit a brief statement that—

(1) Describes his or her relationship to the beneficiary; and

(2) Explains the circumstances that make it impractical for the beneficiary to sign the claim or statement.

(b) Beneficiary not present for services. When a representative of the provider, nonparticipating hospital, or supplier signs a claim or request for payment statement under § 424.36(c), he or she must explain why it was not possible to obtain the beneficiary’s signature. (For example: “Patient not physically present for test.”)
§ 424.40 Request for payment effective for more than one claim.

(a) Basic procedure. A separate request for payment statement prescribed by HCFA and signed by the beneficiary (or by his or her representative) may be included in claims by reference, in the circumstances specified in paragraphs (b) through (d) of this section.

(b) Claims filed by a provider or non-participating hospital—(1) Inpatient services. A signed request for payment statement, included in the first claim for Part A services furnished by a facility (a participating hospital or SNF, or a nonparticipating hospital that has elected to claim payment) during a beneficiary's period of confinement, may be effective for all claims for Part A services the facility furnishes that beneficiary during that confinement.

(2) Home health services and outpatient physical therapy or speech pathology services. A signed request for payment statement, included in the first claim for home health services or outpatient physical therapy or speech pathology services furnished by a provider under a plan of treatment, may be effective for all claims for home health services or outpatient physical therapy or speech pathology services furnished by the provider under that plan of treatment.

(c) Signed statement in the provider record—(1) Services to inpatients. A signed request for payment statement in the files of a participating hospital or SNF may be effective for all claims for services furnished to the beneficiary during a single inpatient stay in that facility—

(i) By the hospital or SNF;

(ii) By physicians, if their services are billed by the hospital or SNF in its name; or

(iii) By physicians who bill separately, if the services were furnished in the hospital or SNF.

(2) Services to outpatients: Providers and renal dialysis facilities. A signed request for payment statement retained in the provider's or facility's files may be effective indefinitely, for all claims for services furnished to that beneficiary on an outpatient basis—

(i) By the provider or facility;

(ii) By physicians whose services are billed by the provider or facility in its name; or

(iii) By physicians who bill separately, if the services were furnished in the provider or facility.

(3) Services to outpatients: Independent rural health clinics and Federally qualified health centers. A signed request for payment statement retained in the clinic's or center's files may be effective indefinitely for all claims for services furnished to that beneficiary by the clinic.

(d) Signed statement in the supplier's record. A signed request for payment statement retained in the supplier's file may be effective indefinitely subject to the following restrictions:

(1) This policy does not apply to unassigned claims for rental of durable medical equipment (DME).

(2) With respect to assigned claims for rental or purchase of DME, a new statement is required if another item of equipment is rented or purchased.

[53 FR 6634, Mar. 2, 1988, as amended at 57 FR 24982, June 12, 1992]

§ 424.44 Time limits for filing claims.

(a) Basic limits. Except as provided in paragraph (b) of this section, the claim must be mailed or delivered to the intermediary or carrier, as appropriate—

(1) On or before December 31 of the following year for services that were furnished during the first 9 months of a calendar year; and

(2) On or before December 31 of the second following year for services that were furnished during the last 3 months of the calendar year.

(b) Extension of filing time because of error or misrepresentation. (1) The time for filing a claim will be extended if failure to meet the deadline in paragraph (a) of this section was caused by error or misrepresentation of an employee, intermediary, carrier, or agent of the Department that was performing Medicare functions and acting within the scope of its authority.

(2) The time will be extended through the last day of the 6th calendar month following the month in which the error or misrepresentation is corrected.
(c) Extension of period ending on a nonworkday. If the last day of the period allowed under paragraph (a) or (b) of this section falls on a Federal nonworkday (a Saturday, Sunday, legal holiday, or a day which by statute or Executive Order is declared to be a nonworkday for Federal employees), the time is extended to the next succeeding workday.

§ 424.45 What constitutes a claim for purposes of meeting the time limits.

A written statement of intent to claim Medicare benefits constitutes a claim if—

(a) The statement is filed with HCFA or any carrier or intermediary within the time limits specified in §424.44;

(b) The statement indicates the intent to claim Medicare payment for specified services furnished to an identified beneficiary; and

(c) A claim that meets the requirements of §424.32(a) is filed within 6 months after the month in which the intermediary or carrier, as appropriate, advises the claimant to file that claim.

Subpart D—To Whom Payment Is Ordinarily Made

§ 424.50 Scope.

(a) This subpart specifies to whom Medicare payment is ordinarily made for different kinds of services.

(b) Subpart E of this part sets forth provisions applicable in special situations.

(c) Subpart F of this part specifies the exceptional circumstances under which payment may be made to an assignee or reassigenee.

§ 424.51 Payment to the provider.

(a) Basic rule. Except as specified in paragraph (b) of this section, Medicare pays the provider for services furnished by a provider.

(b) Exception. Medicare pays the beneficiary for outpatient hospital services if the hospital has collected an amount in excess of the unmet deductible and coinsurance, as specified in §489.30(b)(4) of this chapter.

§ 424.52 Payment to a nonparticipating hospital.

Medicare pays a nonparticipating hospital for the following services, if covered, in the specified circumstances:

(a) Emergency inpatient and outpatient services furnished by a U.S. hospital, if the hospital has in effect an election to claim payment in accordance with subpart G of this part.

(b) Certain medical and other health services covered under Medicare Part B and furnished by a U.S. hospital, if the hospital meets the requirements of §424.55 for payment as a supplier.

(c) Emergency or nonemergency inpatient services furnished by a foreign hospital if the hospital has in effect an election to claim payment in accordance with subpart G of this part.

§ 424.53 Payment to the beneficiary.

Medicare pays the beneficiary for the following services, if covered, in the specified circumstances:

(a) Emergency inpatient and outpatient services furnished by a nonparticipating U.S. hospital that has not elected to claim payment in accordance with subpart G of this part.

(b) Certain medical and other health services covered under Medicare Part B and furnished by a nonparticipating U.S. hospital, if the hospital does not receive assigned payment as a supplier under §424.55.

(c) Emergency or nonemergency services furnished by a foreign hospital if the hospital does not have in effect an election to claim payment in accordance with subpart H of this part.

(d) Physician and ambulance services furnished outside the United States.

(e) Services furnished by a supplier if the claim has not been assigned to the supplier.

§ 424.54 Payment to the beneficiary's legal guardian or representative payee.

Medicare may pay amounts due a beneficiary to the beneficiary's legal guardian or representative payee.

§ 424.55 Payment to the supplier.

(a) Medicare pays the supplier for covered services if the beneficiary (or
§ 424.56 Payment to a beneficiary and to a supplier.

(a) Conditions for split payment. If the beneficiary assigns the claim after paying part of the bill, payment may be made partly to the beneficiary and partly to the supplier.

(b) Payment to the supplier. Payment to the supplier who submits the assigned claim is for whichever of the following amounts is less:

1. The reasonable charge minus the amount the beneficiary had already paid to the supplier; or
2. The full Part B benefit due for the services furnished.

(c) Payment to the beneficiary. Any part of the Part B benefit which, on the basis of paragraph (b) of this section, is not payable to the supplier, is paid to the beneficiary.

(d) Examples.

Example 1. An assigned bill of $300 on which partial payment of $100 has been made is submitted to the carrier. The carrier determines that $300 is the reasonable charge for the service furnished. Total payment due is 80 percent of $300 or $240. Of this amount, $200 (the difference between the $100 partial payment and the $300 reasonable charge) is paid to the supplier. The remaining $40 is paid to the beneficiary.

Example 2. An assigned bill of $275 on which partial payment of $275 has been made is submitted to the carrier. The carrier determines that $275 is the reasonable charge for the services. Total payment due is 80 percent of $275 or $220. The $220 is paid to the beneficiary, since any payment to the supplier, when added to the $275 partial payment would exceed the reasonable charge for the services furnished.


§ 424.57 Special payment rules for items furnished by DMEPOS suppliers and issuance of DMEPOS supplier billing numbers.

(a) Definitions. As used in this section “DMEPOS” is the acronym for durable medical equipment, prosthetics, orthotics and supplies. A “supplier” is an entity or individual, including a physician or part A provider, which sells or rents part B covered items to Medicare beneficiaries and which meets the standards in paragraph (c) of this section.

(b) Medicare pays for items furnished by a supplier with a billing number to the—

1. Supplier if the beneficiary (or the person authorized to request payment on the beneficiary’s behalf) assigns the claim to the supplier and the supplier accepts assignment;
2. Beneficiary, if the supplier does not accept assignment; or
3. Partly to the beneficiary and partly to the supplier, if the supplier accepts assignment of the bill, as described in §424.56.

(c) Medicare does not issue a billing number to a supplier that submits claims for items listed in §421.210(b) of this subchapter until that supplier
meets, and certifies that it meets, the following standards. The supplier—
(1) In response to orders which it receives, fills those orders from its own inventory or inventory in other companies with which it has contracted to fill such orders or fabricates or fits items for sale from supplies it buys under a contract;
(2) Is responsible for delivery of Medicare covered items to Medicare beneficiaries;
(3) Honors all warranties express and implied under applicable State law;
(4) Answers any questions or complaints a beneficiary has about the item or use of the item that was sold or rented to him or her, and refers beneficiaries with Medicare questions to the appropriate carrier;
(5) Maintains and repairs directly or through a service contract with another company, items it has rented to beneficiaries;
(6) Accepts returns of substandard (less than full quality for the particular item) or unsuitable items (inappropriate for the beneficiary at the time it was fitted and/or sold) from beneficiaries;
(7) Discloses consumer information to each beneficiary with whom it does business which consists of the supplier standards to which it must conform;
(8) Complies with the disclosure provisions in §420.206;
(9) Complies with all applicable State and Federal licensure and regulatory requirements;
(10) Maintains a physical facility on an appropriate site; and
(11) Has proof of appropriate liability insurance.
(d) If a supplier is found not to meet the standards in paragraph (c) of this section, its billing number is revoked, effective 15 days after the entity is sent notice of the revocation. A billing number may be issued, with the concurrence of HCFA, when a supplier has successfully completed a corrective action plan rectifying past violations of the supplier standards and provided sufficient assurance that it will comply with the supplier standards in the future. Corrective action includes repayment of monies due to beneficiaries and Medicare, and honoring applicable warranties.
(e) Suppliers must renew their applications for a billing number 3 years after the billing numbers are first re-issued, except for the first reissuance process, as follows: suppliers must renew applications for supplier numbers 2 years after initial issuance of billing numbers for one third of all suppliers. Another one third of suppliers must reapply 3 years after initial issuance. The last third of suppliers must reapply 4 years after initial issuance. Thereafter, each supplier must reapply 3 years after its last number is issued, unless no claim for an item furnished by a supplier has been submitted for four consecutive quarters, in which case the supplier must submit a new request for another billing number.
(f) Suppliers are required to have complaint resolution protocols to address beneficiary complaints which relate to the supplier standards in paragraph (c) of this section and to keep written complaints and related correspondence, and any notes of actions taken in response to written or oral complaints. Failure to maintain such information may be considered evidence that supplier standards have not been met. If a carrier determines that a supplier is not satisfactorily responding to one or more beneficiary complaints, the carrier may require that a supplier maintain the following information on all written and oral beneficiary complaints, including telephone complaints, it receives: The name, address, telephone number and health insurance claim number of the complainant, a summary of the complaint and the date it was made; the name of the person taking the complaint, a summary of any actions taken to resolve the complaint; and, if an investigation was not conducted, the name of the person making the decision and the reason for the decision.

[57 FR 27308, June 18, 1992, as amended at 60 FR 63444, Dec. 11, 1995]
§ 424.60 Scope.
(a) This subpart sets forth provisions applicable to payment after the beneficiary's death and payment to entities that provide coverage complementary to Medicare Part B.
(b) The provisions applicable to payment for services excluded as custodial care or services not reasonable and necessary are set forth in §§ 405.332 through 405.336 of this chapter.

§ 424.62 Payment after beneficiary's death: Bill has been paid.
(a) Scope. This section specifies the persons whom Medicare pays, and the conditions for payments, when the beneficiary has died and the bill has been paid.
(b) Situation. (1) The beneficiary has received covered services for which he could receive direct payment under § 424.53.
(2) The beneficiary died without receiving Medicare payment.
(3) The bill has been paid.
(c) Persons whom Medicare pays. In the situation described in paragraph (b) of this section, Medicare pays the following persons in the specified circumstances:
(1) The person or persons who, without a legal obligation to do so, paid for the services with their own funds, before or after the beneficiary's death.
(2) The legal representative of the beneficiary's estate if the services were paid for by the beneficiary before he or she died, or with funds from the estate.
(3) If the deceased beneficiary or his or her estate paid for the services and no legal representative of the estate has been appointed, the survivors, in the following order of priority:
(i) The person found by SSA to be the surviving spouse who was living in the same household with the deceased at the time of death and was not, for the month of death, entitled to monthly social security or railroad retirement benefits on the basis of the same earnings record as the deceased beneficiary;
(ii) The child or children, who were, for the month of death, entitled to monthly social security or railroad retirement benefits on the basis of the same earnings record as the deceased (and, if there is more than one child, in equal parts to each child);
(iii) The parent or parents, who were, for the month of death, entitled to monthly social security or railroad retirement benefits on the basis of the same earnings record as the deceased (and, if there is more than one parent, in equal parts to each parent);
(iv) The person found by SSA to be the surviving spouse who was not living in the same household with the deceased at the time of death and was not, for the month of death, entitled to monthly social security or railroad retirement benefits on the basis of the same earnings record as the deceased beneficiary;
(v) The child or children who were not entitled to monthly social security or railroad retirement benefits on the basis of the same earnings record as the deceased (and, if there is more than one child, in equal parts to each child);
(vi) The parent or parents who were not entitled to monthly social security or railroad retirement benefits on the basis of the same earnings record as the deceased (and, if there is more than one parent, in equal parts to each parent).
(4) If none of the listed relatives survive, no payment is made.
(5) If the services were paid for by a person other than the deceased beneficiary, and that person died before payment was completed, Medicare does not pay that person's estate. Medicare pays a surviving relative of the deceased beneficiary in accordance with the priorities in paragraph (c)(3) of this section. If none of those relatives survive. Medicare pays the legal representative of the deceased beneficiary's estate. If there is no legal representative of the estate, no payment is made.
(d) Amount of payment. The amount of payment is the amount due, including unegotiated checks issued for the purpose of making direct payment to the beneficiary.
(e) Conditions for payment. For payment to be made under this section—
(1) The person who claims payment must meet the following requirements:
   (i) Submit a claim on a HCFA-prescribed form and an itemized bill in accordance with the requirements of this subpart. (See paragraph (g) of this section for an exception.)
   (ii) Provide evidence that the services were furnished if the intermediary or carrier requests it.
   (iii) Provide evidence of payment of the bill and of the identity of the person who paid it.

(2) If a person claims payment as the legal representative of the deceased beneficiary's estate, he or she must also submit a copy of the papers showing appointment as legal representative.

(3) If a person claims payment as a survivor of the beneficiary, he or she must also submit evidence, if the intermediary or carrier requests it, that he or she is highest on the priority list of paragraph (c)(3) of this section.

(f) Evidence of payment. Evidence of payment may be—
   (1) A receipted bill, or a properly completed "Report of Services" section of a claim form, showing who paid the bill;
   (2) A cancelled check;
   (3) A written statement from the provider or supplier or an authorized staff member; or
   (4) Other probative evidence.

(g) Exception: Claim submitted before beneficiary died. If a claim and itemized bill has been submitted by or on behalf of the beneficiary before he or she died, submission of another claim form is not required; any written request by the person seeking payment is sufficient.

§ 424.66 Payment after beneficiary's death: Bill has not been paid.

(a) Scope. This section specifies whom Medicare pays, and the conditions for payment when the beneficiary has died and the bill has not been paid.

(b) Situation. (1) The beneficiary has received covered Part B services furnished by a physician or other supplier.

(2) The beneficiary died without making an assignment to the physician or other supplier or receiving Medicare payment.

(3) The bill has not been paid.

(c) To whom payment is made. In the situation described in paragraph (b) of this section, Medicare pays as follows:
   (1) Payment to the supplier. Medicare pays the physician or other supplier if he or she—
      (i) Files a claim on a HCFA-prescribed form in accordance with the applicable requirements of this subpart;
      (ii) Upon request from the carrier, provides evidence that the services for which it claims payment were, in fact, furnished; and
      (iii) Agrees in writing to accept the reasonable charge as full charge for the services.
   (2) Payment to a person who assumes legal obligation to pay for the services. If the physician or other supplier does not agree to accept the reasonable charge as full charge for the service, Medicare pays any person who submits to the carrier all of the following:
      (i) A statement indicating that he or she has assumed legal obligation to pay for the services.
      (ii) A claim on a HCFA-prescribed form in accordance with the requirements of this subpart. (If a claim had been submitted by or on behalf of the beneficiary before he or she died, submission of another claim form is not required; a written request by the person seeking payment meets the requirement for a claim.)
      (iii) An itemized bill that identifies the claimant as the person to whom the physician or other supplier holds responsible for payment. (If such an itemized bill had been submitted by or on behalf of the beneficiary before he or she died, submission of another itemized bill is not required.)
      (iv) If the intermediary or carrier requests it, evidence that the services were actually furnished.


§ 424.66 Payment to entities that provide coverage complementary to Medicare Part B.

(a) Conditions for payment. Medicare may pay an entity for Part B services furnished by a physician or other supplier if the entity meets all of the following requirements:
   (1) Provides coverage of the service under a complementary health benefit plan that meets the requirements specified in subpart H of this part.
§ 424.70 Basis and scope.

(a) Statutory basis. This subpart implements sections 1815(c) and 1842(b)(6) of the Act, which establish limitations on who may receive payments due a provider or supplier of services or a beneficiary.

(b) Scope. This subpart—

(1) Prohibits the assignment, reassignment, or other transfer of the right to Medicare payments except under specified conditions;

(2) Sets forth the sanctions that HCFA may impose on a provider or supplier that violates this prohibition, or on a supplier that violates the conditions to which it agreed in accepting assignment from the individual; and

(3) Specifies the conditions for payment under court-ordered assignments or reassignments.

§ 424.71 Definitions.

As used in this subpart, unless the context indicates otherwise—

Court of competent jurisdiction means a court that has jurisdiction over the subject matter and the parties before it.

Facility means a hospital or other institution that furnishes health care services to inpatients.

Health care delivery system or system means a public or private organization for delivering health services. The term includes, but is not limited to, clinics and health care prepayment plans.

Power of attorney means any written documents by which a principal authorizes an agent to—

(1) Receive, in the agent's name, any payments due the principal;

(2) Negotiate checks payable to the principal; or

(3) Receive, in any other manner, direct payment of amounts due the principal.

§ 424.73 Prohibition of assignment of claims by providers.

(a) Basic prohibition. Except as specified in paragraph (b) of this section, Medicare does not pay amounts that are due a provider to any other person under assignment, or power of attorney, or any other direct payment arrangement.

(b) Exceptions to the prohibition—(1) Payment to a government agency or entity. Subject to the requirements of the Assignment of Claims Act (31 U.S.C. 3727), Medicare may pay a government agency or entity under an assignment by the provider.

(2) Payment under assignment established by court order. Medicare may pay under an assignment established by, or in accordance with, the order of a court.
of competent jurisdiction if the assignment meets the conditions set forth in §424.90.

(3) Payment to an agent. Medicare may pay an agent who furnishes billing and collection services to the provider if the following conditions are met:

(i) The agent receives the payment under an agency agreement with the provider;
(ii) The agent’s compensation is not related in any way to the dollar amounts billed or collected;
(iii) The agent’s compensation is not dependent upon the actual collection of payment;
(iv) The agent acts under payment disposition instructions that the provider may modify or revoke at any time and
(v) The agent, in receiving the payment, acts only on behalf of the provider.

Payment to an agent will always be made in the name of the provider.

§ 424.74 Termination of provider agreement.

HCFA may terminate a provider agreement, in accordance with §489.53(a)(1) of this chapter, if the provider—
(a) Executes or continues a power of attorney, or enters into or continues any other arrangement, that authorizes or permits payment contrary to the provisions of this subpart; or
(b) Fails to furnish, upon request by HCFA or the intermediary, evidence necessary to establish compliance with the requirements of this subpart.

§ 424.80 Prohibition of reassignment of claims by suppliers.

(a) Basic prohibition. Except as specified in paragraph (b) of this section, Medicare does not pay amounts that are due a supplier under an assignment to any other person under reassignment, power of attorney, or any other direct arrangement.

(b) Exceptions to the basic rule—(1) Payment to employer. Medicare may pay the supplier’s employer if the supplier is required, as a condition of employment, to turn over to the employer the fees for his or her services.

(2) Payment to a facility. Medicare may pay the facility in which the services were furnished if there is a contractual arrangement between the facility and the supplier under which the facility bills for the supplier’s services.

(3) Payment to health care delivery system. Medicare may pay a health care delivery system if there is a contractual arrangement between the system and the supplier under which the system bills for the supplier’s services.

(4) Payment to a government agency or entity. Subject to the requirements of the Assignment of Claims Act (31 U.S.C. 3727), Medicare may pay a government agency or entity under a reassignment by the supplier.

(5) Payment under a reassignment established by court order. Medicare may pay under a reassignment established by, or in accordance with, the order of a court competent jurisdiction, if the reassignment meets the conditions set forth in §424.90.

(6) Payment to an agent. Medicare may pay an agent who furnishes billing and collection services to the supplier, or to the employer, facility, or system specified in paragraphs (b) (1), (2) and (3) of this section, if the conditions of §424.73(b)(3) for payment to a provider’s agent are met by the agent of the supplier or of the employer, facility, or system. Payment to an agent will always be made in the name of the supplier or the employer, facility, or system.

(c) Rules applicable to an employer, facility, or system. An employer, facility, or system that may receive payment under paragraph (b)(1), (b)(2), or (b)(3) of this section will itself be considered the supplier of those services for purposes of the rules of subparts C, D, and E of this part.

[53 FR 6634, Mar. 2, 1988, as amended at 54 FR 4027, Jan. 27, 1989]

§ 424.82 Revocation of right to receive assigned benefits.

(a) Scope. This section sets forth the conditions and procedures for revocation of the right of a supplier or other party to receive Medicare payments.

(b) Definition. As used in this section, other party means an employer, facility, or health care delivery system to which Medicare may make payment under §424.80(b) (1), (2), or (3).
§ 424.83 Hearings on revocation of right to receive assigned benefits.

If the supplier or other party requests a hearing under § 424.82(e)(2)—

(a) The hearing is conducted—
   (1) By a HCFA hearing official who was not involved in the decision to revoke; and
   (2) In accordance with the procedures set forth in §§ 405.824 through 405.833 (but excepting § 405.832(d)) and 405.860 through 405.872 of this chapter. In applying those procedures, “HCFA” is substituted for “carrier”; and “hearing official”, for “hearing officer”.

(b) As soon as practicable after the close of the hearing, the official who conducted it issues a hearing decision that—
   (1) Is based on all the evidence presented at the hearing and included in the hearing record; and
   (2) Contains findings of fact and a statement of reasons.

§ 424.84 Final determination on revocation of right to receive assigned benefits.

(a) Basis of final determination—
   (1) Final determination without a hearing. If the supplier or other party does not request a hearing, HCFA’s revocation determination becomes final at the end of the period specified in the notice of revocation.
   (2) Final determination following a hearing. If there is a hearing, the hearing decision constitutes HCFA’s final determination.

(b) Notice of final determination. HCFA sends the supplier or other party a written notice of the final determination and, if there was a hearing, includes a copy of the hearing decision.
(c) Application of the final determination—(1) A final determination not to revoke is the final administrative decision by HCFA on the matter.

(2) A final determination to revoke remains in effect until HCFA finds that the reason for the revocation has been removed and that there is reasonable assurance that it will not recur.

(d) Effect of revocation when supplier or other party has a financial interest in another entity. Revocation of the party’s right to accept assignment also applies to any corporation, partnership, or other entity in which the party, directly or indirectly, has or acquires all or all but a nominal part of the financial interest.


§ 424.86 Prohibition of assignment of claims by beneficiaries.

(a) Basic prohibition. Except as specified in paragraph (b) of this section, Medicare does not pay amounts that are due a beneficiary under §424.53 to any other person under assignment, power of attorney, or any other direct payment arrangement.

(b) Exceptions—(1) Payment to a government agency or entity. Subject to the requirements of the Assignment of Claims Act (31 U.S.C. 3727), Medicare may pay a government agency or entity under an assignment by a beneficiary (or by the beneficiary’s legal guardian or representative payee).

(2) Payment under an assignment established by court order. Medicare may pay under an assignment established by, or in accordance with, a court order if the assignment meets the conditions set forth in §424.90.

§ 424.90 Court ordered assignments: Conditions and limitations.

(a) Conditions for acceptance. An assignment or reassignment established by or in accordance with a court order is effective for Medicare payments only if—

(1) Someone files a certified copy of the court order and of the executed assignment or reassignment (if it was necessary to execute one) with the intermediary or carrier responsible for processing the claim; and

(2) The assignment or reassignment—

(i) Applies to all Medicare benefits payable to a particular person or entity during a specified or indefinite time period; or

(ii) Specifies a particular amount of money, payable to a particular person or entity by a particular intermediary or carrier.

(b) Retention of authority to reduce interim payments to providers. A court-ordered assignment does not preclude the intermediary or carrier from reducing interim payments, as set forth in §413.64(i) of this chapter, if the provider or assignee is in imminent danger of insolvency or bankruptcy.

(c) Liability of the parties. The party that receives payments under a court-ordered assignment or reassignment that meets the conditions of paragraph (a) of this section and the party that would have received payment if the court order had not been issued are jointly and severally responsible for any Medicare overpayment to the former.

Subpart G—Special Conditions: Emergency Services Furnished by a Nonparticipating Hospital

§ 424.100 Scope.

This subpart sets forth procedures and criteria that are followed in determining whether Medicare will pay for emergency services furnished by a hospital that is located in the United States and does not have in effect a provider agreement, that is, an agreement to participate in Medicare.

§ 424.101 Definitions.

As used in this subpart, unless the context indicates otherwise—

Emergency services means inpatient or outpatient hospital services that are necessary to prevent death or serious impairment of health and, because of the danger to life or health, require use of the most accessible hospital available and equipped to furnish those services.

Hospital means a facility that—

(1) Is primarily engaged in providing, by or under the supervision of doctors of medicine or osteopathy, inpatient services for the diagnosis, treatment,
§ 424.102 Situations that do not constitute an emergency.

Without additional evidence of a threat to life or health, the following situations do not in themselves indicate a need for emergency services:

(a) Lack of care at home.
(b) Lack of transportation to a participating hospital.
(c) Death of the patient in the hospital.

§ 424.103 Conditions for payment for emergency services.

Medicare pays for emergency services furnished to a beneficiary by a non-participating hospital or under arrangements made by such a hospital if the conditions of this section are met.

(a) General requirements. (1) The services are of the type that Medicare would pay for if they were furnished by a participating hospital.
(2) The hospital has in effect an election to claim payment for all emergency services furnished in a calendar year in accordance with § 424.104.
(3) The need for emergency services arose while the beneficiary was not an inpatient in a hospital.
(4) In the case of inpatient hospital services, the services are furnished during a period in which the beneficiary could not be safely discharged or transferred to a participating hospital or other institution.
(5) The determination that the hospital was the most accessible hospital available and equipped to furnish the services is made in accordance with § 424.106.

(b) Medical information requirements. A physician (or, if appropriate, the hospital) submits medical information that—
(1) Describes the nature of the emergency and specifies why it required that the beneficiary be treated in the most accessible hospital;
(2) Establishes that all the conditions in paragraph (a) of this section are met; and
(3) Indicates when the emergency ended, which, for inpatient hospital services, is the earliest date on which the beneficiary could be safely discharged or transferred to a participating hospital or other institution.

§ 424.104 Election to claim payment for emergency services furnished during a calendar year.

(a) Terms of the election. The hospital agrees to the following:
(1) To comply with the provisions of subpart C of part 489 of this chapter relating to charges for items and services the hospital may make to the beneficiary, or any other person on his or her behalf.
(2) To comply with the provisions of subpart D of part 489 of this chapter relating to proper disposition of monies incorrectly collected from, or on behalf of a beneficiary.
(3) To request payment under the Medicare program based on amounts specified in § 413.74 of this chapter.

(b) Filing of election statement. An election statement must be filed on a form designated by HCFA, signed by an authorized official of the hospital, and either received by HCFA, or postmarked, before the close of the calendar year of election.

(c) Acceptance and effective date of election. If HCFA accepts the election statement, the election is effective as of the earliest day of the calendar year of election from which HCFA determines the hospital has been in continuous compliance with the requirements of section 1814(d) of the Act.

(d) Appeal by hospital. Any hospital dissatisfied with a determination that it does not qualify to claim reimbursement shall be entitled to appeal the determination as provided in part 498 of this chapter.

(e) Conditions for reinstatement after notice of failure to continue to qualify. If HCFA has notified a hospital that it no
§ 424.108 Payment to a hospital.

(a) Conditions for payment. Medicare pays the hospital for emergency services if the hospital—

(1) Has in effect a statement of election to claim payment for all covered emergency services furnished during a calendar year, in accordance with §424.104;

(2) Claims payment in accordance with §424.32; and

(3) Submits evidence requested by HCFA to establish that the services meet the requirements of this subpart.

(b) Subsequent claims. If the hospital files subsequent claims because the initial claim did not include all the services furnished, those claims must include physicians' statements that—

(1) Contain sufficient information to clearly establish that, when the additional services were furnished, the emergency still existed; and

(2) Indicate when the emergency ended, which, for inpatient hospital services, is the earliest date on which the beneficiary could be safely discharged or transferred to a participating hospital or other institution.
§ 424.109 Payment to the beneficiary.

Medicare pays the beneficiary for emergency services if the following conditions are met:
(a) The hospital does not have in effect an election to claim payment.
(b) The beneficiary, or someone on his or her behalf, submits—
   (1) A claim that meets the requirements of § 424.32;
   (2) An itemized hospital bill; and
   (3) Evidence requested by HCFA to establish that the services meet the requirements of this subpart.

Subpart H—Special Conditions: Services Furnished in a Foreign Country

§ 424.120 Scope.

This subpart sets forth the conditions for payment for services furnished in a foreign country.

§ 424.121 Scope of payments.

Subject to the conditions set forth in this subpart—
(a) Medicare Part A pays, in the amounts specified in § 413.74 of this chapter, for emergency and non-emergency inpatient hospital services furnished by a foreign hospital.
(b) Medicare Part B pays for certain physicians’ services and ambulance services furnished in connection with covered inpatient care in a foreign hospital, as specified in § 424.124.
(c) All other services furnished outside the United States are excluded from Medicare coverage, as specified in § 405.313 of this chapter.

§ 424.122 Conditions for payment for emergency inpatient hospital services.

Medicare Part A pays for emergency inpatient hospital services furnished by a foreign hospital if the following conditions are met:
(a) At the time of the emergency that required the inpatient hospital services, the beneficiary was—
   (1) In the United States; or
   (2) In Canada traveling between Alaska and another State without unreasonable delay and by the most direct route.
(b) The foreign hospital was closer to, or more accessible from, the site of the emergency than the nearest United States hospital equipped to deal with, and available to treat, the individual’s illness or injury.
(c) The conditions for payment for emergency services set forth in § 424.103 are met.
(d) The hospital is a hospital as defined in § 424.101, and is licensed, or approved as meeting the conditions for licensing, by the appropriate agency of the country in which it is located.
(e) The determination of whether the hospital was more accessible is made in accordance with § 424.106.

§ 424.123 Conditions for payment for nonemergency inpatient services furnished by a hospital closer to the individual’s residence.

Medicare Part A pays for inpatient hospital services furnished by a foreign hospital if the following conditions are met:
(a) The beneficiary is a resident of the United States.
(b) The foreign hospital is closer or more accessible to the beneficiary’s residence than the nearest United States hospital equipped to deal with, and available to treat, the individual’s illness or injury.
(c) The foreign hospital is—
   (1) A hospital as defined in § 424.101 and, it is licensed, or approved as meeting the conditions for licensing, by the appropriate agency of the country in which it is located; and
   (2) Accredited by the Joint Commission on Accreditation of Hospitals (JCAH) or accredited or approved by a program of the country where it is located under standards that HCFA finds to be essentially equivalent to those of the JCAH.
(d) The services are covered services that Medicare would pay for if they were furnished by a participating hospital.

§ 424.124 Conditions for payment for physician services and ambulance services.

(a) Basic rules. Medicare Part B pays for physician and ambulance services if—
   (1) They are furnished—
§ 424.350 Replacement of checks that are lost, stolen, defaced, mutilated, destroyed, or paid on forged endorsements.

(a) U.S. Government checks—(1) Responsibility. The Treasury Department is responsible for the investigation and settlement of claims in connection with Treasury checks issued on behalf of HCFA.

(2) Action by HCFA. HCFA forwards reports of lost, stolen, defaced, mutilated, destroyed, or forged Treasury checks to the Treasury Department disbursing center responsible for issuing checks.

(3) Action by the Treasury Department. The Treasury Department will replace and begin reclamation of Treasury checks in accordance with Treasury Department regulations (31 CFR parts 235, 240, and 245).

(b) Intermediary and carrier benefit checks. Checks issued by intermediaries and carriers are drawn on commercial banks and are not subject to the Federal laws and Treasury Department regulations.
§ 424.352 Intermediary and carrier checks that are lost, stolen, defaced, mutilated, destroyed or paid on forged endorsements.

(a) When an intermediary or carrier is notified by a payee that a check has been lost, stolen, defaced, mutilated, destroyed, or paid on forged endorsement, the intermediary or carrier contacts the commercial bank on whose paper the check was drawn and determines whether the check has been negotiated.

(b) If the check has been negotiated—

(1) The intermediary or carrier provides the payee with a copy of the check and other pertinent information (such as a claim form, affidavit or questionnaire to be completed by the payee) required to pursue his or her claim in accordance with State law and commercial banking regulations.

(2) To pursue the claim, the payee must examine the check and certify (by completing the claim form, questionnaire or affidavit) that the endorsement is not the payee’s.

(3) The claim form and other pertinent information is sent to the intermediary or carrier for review and processing of the claim.

(4) The intermediary or carrier reviews the payee’s claim. If the intermediary or carrier determines that the claim appears to be valid, it forwards the claim and a copy of the check to the issuing bank. The intermediary or carrier takes further action to recover the proceeds of the check in accordance with the State law and regulations.

(5) Once the intermediary or carrier recovers the proceeds of the initial check, the intermediary or carrier issues a replacement check to the payee.

(6) If the bank of first deposit refuses to settle on the check for good cause, the payee must pursue the claim on his or her own and the intermediary or carrier will not reissue the check to the payee.

(c) If the check has not been negotiated—

(1) The intermediary or carrier arranges with the bank to stop payment on the check; and

(2) Except as provided in paragraph (d), the intermediary or carrier reissues the check to the payee.

(d) No check may be reissued under (c)(2) unless the claim for a replacement check is received by the intermediary or carrier no later than 1 year from the date of issuance of the original check, unless State law (including any applicable Federal banking laws or regulations that may affect the relevant State proceeding) provides a longer period which will control.

[58 FR 65130, Dec. 13, 1993]
FINDING AIDS

A list of CFR titles, subtitles, chapters, subchapters and parts and an alphabetical list of agencies publishing in the CFR are included in the CFR Index and Finding Aids volume to the Code of Federal Regulations which is published separately and revised annually.

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(Revised as of October 1, 2000)

The Director of the Federal Register has approved under 5 U.S.C. 552(a) and 1 CFR part 51 the incorporation by reference of the following publications. This list contains only those incorporations by reference effective as of the revision date of this volume. Incorporations by reference found within a regulation are effective upon the effective date of that regulation. For more information on incorporation by reference, see the preliminary pages of this volume.

42 CFR (PARTS 400–429)
HEALTH CARE FINANCING ADMINISTRATION, DEPARTMENT OF HEALTH AND HUMAN SERVICES

American Hospital Association
Available from: Health Care Financing Administration, Office of Management and Budget, Division of Communication Services, Printing and Publishing Branch, Gwynn Oak Bldg., Baltimore, MD 21235
Chart of Accounts for Hospitals (1973 Ed.) .............................................. 405.415(b)(7)(i)

American Lung Association
(Formerly National Tuberculosis Association)
1740 Broadway, New York, NY 10019
Diagnostic Standards and Classification of Tuberculosis (1974 Ed.) 405.1039(b)(2)

American National Standards Institute
11 West 42nd St., New York, NY 10036 Telephone: (212) 642-4900
ANSI A117.1–1971 Specifications for Making Buildings and Facilities Accessible to and Usable by Physically Handicapped People. 405.1134(c)

American Psychiatric Association
Division of Publications and Marketing, 1400 K Street NW., Washington, DC 20005
Diagnostic and Statistical Manual of Mental Disorders, 1980 (3rd Ed.). 405.243(b); 405.1037(a)(12)

Association for the Advancement of Medical Instrumentation
3330 Washington Blvd., Arlington, VA 22201–4598
ASNI/AAMI RD5–1992, Hemodialysis Systems, second edition .......... 405.2150(a);
Recommended Practice for Reuse of Hemodialyzers ......................... 405.2150

National Academy of Sciences
Academy Press, 2101 Constitution Ave. NW., Washington, DC 20418
Recommended Dietary Allowances Current Edition, 1980 (9th Ed.) 405.1025(c)(2)

National Fire Protection Association (NFPA)
1 Batterymarch Park, P.O. Box 9101, Quincy, MA 02269–9101

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XXIX   Federal Communications Commission (Parts 3900—3999)
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XXXI   Farm Credit Administration (Parts 4100—4199)
XXXIII   Overseas Private Investment Corporation (Part 4301)
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IX Federal Housing Finance Board (Parts 900—999)
XI Federal Financial Institutions Examination Council (Parts 1100—1199)
XIV Farm Credit System Insurance Corporation (Parts 1400—1499)
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XVII Office of Federal Housing Enterprise Oversight, Department of Housing and Urban Development (Parts 1700—1799)
XVIII Community Development Financial Institutions Fund, Department of the Treasury (Parts 1800—1899)

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Redesignation Table I

At 51 FR 22040, June 17, 1986, §§ 405.1011—405.1042 (subpart J), formerly appearing in title 42, part 405, were redesignated as part 482 of title 42.

For the convenience of the user, the following table shows the relationship of the redesignated sections.

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Redesignation Table II

At 51 FR 34790, Sept. 30, 1986, certain sections of regulations formerly appearing in Title 42, Part 405, Subpart D, were redesignated as Part 413 of Title 42.

For the convenience of the user, the following table shows the relationship of the redesignated sections.

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Redesignation Table III

At 51 FR 34764, Sept. 30, 1986, a document was published by the Department of Health and Human Services which established in title 42 a new chapter V, Office of Inspector General—Health Care, consisting of parts 1001 through 1004. This document also removed part 101 from title 45. A second document at 51 FR 34786, Sept. 30, 1986 amended title 42, parts 412, 420, 455, 466, 474, and 489. The following derivation table identifies the sections of 42 CFR chapter IV and 45 CFR subtitle A from which the new 42 CFR chapter V derives its content.

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Redesignation Table IV

At 51 FR 41335, Nov. 14, 1986, certain sections of regulations formerly appearing in title 42, part 405, subpart B were redesignated as part 410 of title 42.

For the convenience of the user, the following table shows the relationship of the redesignated sections.

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Redesignation Table V

At 52 FR 22446, June 12, 1987, certain sections of regulations appearing in title 42, part 405, subparts F and O were redesignated as part 498 of title 42.

For the convenience of the user, the following table shows the relationship of the redesignated sections.

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Redesignation Table VI

At 52 FR 36746, Sept. 30, 1987, regulations appearing in title 42, part 110 were redesignated as part 417, subpart A of title 42.

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Redesignation Table VII

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### Title 42—Public Health

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Redesignation Table VIII

At 53 FR 23100, June 17, 1988, §§ 405.1901—405.1913 (subpart S), formerly appearing in title 42, part 405, were redesignated as part 488 of title 42. For the convenience of the user, the following table shows the relationship of the redesignated sections.

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Redesignation Table IX

At 53 FR 47201, November 22, 1988, §§ 405.201—405.226 (subpart B), formerly appearing in title 42, part 405, were redesignated as part 407 of title 42.

For the convenience of the user, the following table shows the relationship of the redesignated sections.

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Redesignation Table X

At 52 FR 48114, Dec. 18, 1988, regulations appearing in Title 42, part 405 were redesignated as part 408 of Title 42. The redesignation table was correctly published at 53 FR 4156, Feb. 12, 1988.

For the convenience of the user, the following table shows the relationship of the redesignated sections.

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Redesignation Table XI

At 54 FR 33355, August 14, 1989, §§405.1201—405.1230 (subpart L), formerly appearing in title 42, part 405, were redesignated as part 484 of title 42.

For the convenience of the user, the following table shows the relationship of the redesignated sections.

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<td>405.1230</td>
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Redesignation Table XII

At 54 FR 41733 and 41734, October 11, 1989, subpart C of part 405 was amended by removing §§ 405.308 through 405.344 and a new part 411 was added, to redesignate, revise, and amplify the content removed from subpart C of part 405.

For the convenience of the user, the following table (appearing at 54 FR 41733, Oct. 11, 1989 and corrected at 55 FR 1820, Jan. 19, 1990) shows the relationship of the old §§ 405.308 through 405.344 and the new part 411.

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Redesignation Table XIII

At 56 FR 51985, October 17, 1991, §§417.110—417.137 (subpart A) were redesignated as subpart V, part 417 of title 42.

For the convenience of the user, the following table shows the relationship of the redesignated sections.

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Redesignation Table XIV

At 60 FR 2326, January 9, 1995, §§ 405.1411—405.1416 (subpart N), formerly appearing in title 42, part 405, were redesignated as subpart C, part 486, title 42. Additionally, §§ 405.1701, 405.1702, and 405.1715—405.1726, formerly appearing in part 405, were redesignated as subpart H, part 485, title 42. Sections 405.1730—405.1737 (subpart Q), formerly appearing in part 405, were redesignated as subpart D, part 486, title 42. And §§ 411.60 through 411.65 (subpart E) and §§ 411.70 through 411.75 (subpart F), appearing in part 411 of title 42, were redesignated as subparts F and G of the same part.

For the convenience of the user, the following table shows the relationship of the redesignated sections.

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### List of CFR Sections Affected

All changes in this volume of the Code of Federal Regulations which were made by documents published in the Federal Register since January 1, 1986, are enumerated in the following list. Entries indicate the nature of the changes effected. Page numbers refer to Federal Register pages. The user should consult the entries for chapters and parts as well as sections for revisions.


#### 42 CFR

<table>
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<th>1986</th>
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<td>Chapter IV—Continued 405.210 (b)(1) (iii) and (iv) amended</td>
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<td>405.240-405.241 Removed (regulations transferred to Part 410)</td>
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<td>15492</td>
<td>405.243-405.246 Removed (regulations transferred to Part 410)</td>
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<td>405.250-2 Removed (regulations transferred to Part 410)</td>
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