PART 482—CONDITIONS OF PARTICIPATION FOR HOSPITALS

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

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Subpart A—General Provisions

§ 482.1 Basis and scope.
(a) Statutory basis. (1) Section 1861(e) of the Act provides that—
(i) Hospitals participating in Medicare must meet certain specified requirements; and
(ii) The Secretary may impose additional requirements if they are found necessary in the interest of the health and safety of the individuals who are furnished services in hospitals.
(2) Section 1861(f) of the Act provides that an institution participating in Medicare as a psychiatric hospital must meet certain specified requirements imposed on hospitals under section 1861(e), must be primarily engaged in providing, by or under the supervision of a physician, psychiatric services for the diagnosis and treatment of mentally ill persons, must maintain clinical records and other records that the Secretary finds necessary, and must meet staffing requirements that the Secretary finds necessary, and must meet staffing requirements that the Secretary finds necessary to carry out an active program of treatment for individuals who are furnished services in the hospital. A distinct part of an institution can participate as a psychiatric hospital if the institution meets the specified 1861(e) requirements and is primarily engaged in providing psychiatric services, and if the
§ 482.2 Provision of emergency services by nonparticipating hospitals.

(a) The services of an institution that does not have an agreement to participate in the Medicare program may, nevertheless, be reimbursed under the program if—

(1) The services are emergency services; and

(2) The institution meets the requirements of section 1861(e) (1) through (5) and (7) of the Act. Rules applicable to emergency services furnished by non-participating hospitals are set forth in subpart G of part 424 of this chapter.

(b) Section 440.170(e) of this chapter defines emergency hospital services for purposes of Medicaid reimbursement.

in the hospital dependent solely upon certification, fellowship, or membership in a specialty body or society.

(b) Standard: Chief executive officer. The governing body must appoint a chief executive officer who is responsible for managing the hospital.

(c) Standard: Care of patients. In accordance with hospital policy, the governing body must ensure that the following requirements are met:

(1) Every Medicare patient is under the care of:
   (i) A doctor of medicine or osteopathy (This provision is not to be construed to limit the authority of a doctor of medicine or osteopathy to delegate tasks to other qualified health care personnel to the extent recognized under State law or a State's regulatory mechanism);
   (ii) A doctor of dental surgery or dental medicine who is legally authorized to practice dentistry by the State and who is acting within the scope of his or her license;
   (iii) A doctor of podiatric medicine, but only with respect to functions which he or she is legally authorized by the State to perform;
   (iv) A doctor of optometry who is legally authorized to practice optometry by the State in which he or she practices;
   (v) A chiropractor who is licensed by the State or legally authorized to perform the services of a chiropractor, but only with respect to treatment by means of manual manipulation of the spine to correct a subluxation demonstrated by x-ray to exist; and
   (vi) A clinical psychologist as defined in §410.71 of this chapter, but only with respect to clinical psychologist services as defined in §410.71 of this chapter and only to the extent permitted by State law.

(2) Patients are admitted to the hospital only on the recommendation of a licensed practitioner permitted by the State to admit patients to a hospital. If a Medicare patient is admitted by a practitioner not specified in paragraph (c)(1) of this section, that patient is under the care of a doctor of medicine or osteopathy.

(3) A doctor of medicine or osteopathy is on duty or on call at all times.

(4) A doctor of medicine or osteopathy is responsible for the care of each Medicare patient with respect to any medical or psychiatric problem that—
   (i) is present on admission or develops during hospitalization; and
   (ii) Is not specifically within the scope of practice of a doctor of dental surgery, dental medicine, podiatric medicine, or optometry; a chiropractor; or clinical psychologist, as that scope is—
      (A) Defined by the medical staff;
      (B) Permitted by State law; and
      (C) Limited, under paragraph (c)(1)(v) of this section, with respect to chiropractors.

(d) Standard: Institutional plan and budget. The institution must have an overall institutional plan that meets the following conditions:

(1) The plan must include an annual operating budget that is prepared according to generally accepted accounting principles.

(2) The budget must include all anticipated income and expenses. This provision does not require that the budget identify item by item the components of each anticipated income or expense.

(3) The plan must provide for capital expenditures for at least a 3-year period, including the year in which the operating budget specified in paragraph (d)(2) of this section is applicable.

(4) The plan must include and identify in detail the objective of, and the anticipated sources of financing for, each anticipated capital expenditure in excess of $600,000 (or a lesser amount that is established, in accordance with section 1122(g)(1) of the Act, by the State in which the hospital is located) that relates to any of the following:
   (i) Acquisition of land;
   (ii) Improvement of land, buildings, and equipment; or
   (iii) The replacement, modernization, and expansion of buildings and equipment.

(5) The plan must be submitted for review to the planning agency designated in accordance with section 1122(b) of the Act, or if an agency is not designated, to the appropriate health planning agency in the State. (See part 100 of this title.) A capital expenditure
is not subject to section 1122 review if 75 percent of the health care facility’s patients who are expected to use the service for which the capital expenditure is made are individuals enrolled in a health maintenance organization (HMO) or competitive medical plan (CMP) that meets the requirements of section 1876(b) of the Act, and if the Department determines that the capital expenditure is for services and facilities that are needed by the HMO or CMP in order to operate efficiently and economically and that are not otherwise readily accessible to the HMO or CMP because—
   (i) The facilities do not provide common services at the same site;
   (ii) The facilities are not available under a contract of reasonable duration;
   (iii) Full and equal medical staff privileges in the facilities are not available;
   (iv) Arrangements with these facilities are not administratively feasible; or
   (v) The purchase of these services is more costly than if the HMO or CMP provided the services directly.

(6) The plan must be reviewed and updated annually.

(7) The plan must be prepared—
   (i) Under the direction of the governing body; and
   (ii) By a committee consisting of representatives of the governing body, the administrative staff, and the medical staff of the institution.

(e) Standard: Contracted services. The governing body must be responsible for services furnished in the hospital whether or not they are furnished under contracts. The governing body must ensure that a contractor of services (including one for shared services and joint ventures) furnishes services that permit the hospital to comply with all applicable conditions of participation and standards for the contracted services.

(1) The governing body must ensure that the services performed under a contract are provided in a safe and effective manner.

(2) The hospital must maintain a list of all contracted services, including the scope and nature of the services provided.

(f) Standard: Emergency services. (1) If emergency services are provided at the hospital, the hospital must comply with the requirements of §482.55.

(2) If emergency services are not provided at the hospital, the governing body must assure that the medical staff has written policies and procedures for appraisal of emergencies, initial treatment, and referral when appropriate.


§ 482.13 Condition of participation: Patients’ rights.

A hospital must protect and promote each patient’s rights.

(a) Standard: Notice of rights. (1) A hospital must inform each patient, or when appropriate, the patient’s representative (as allowed under State law), of the patient’s rights, in advance of furnishing or discontinuing patient care whenever possible.

(2) The hospital must establish a process for prompt resolution of patient grievances and must inform each patient whom to contact to file a grievance. The hospital’s governing body must approve and be responsible for the effective operation of the grievance process and must review and resolve grievances, unless it delegates the responsibility in writing to a grievance committee. The grievance process must include a mechanism for timely referral of patient concerns regarding quality of care or premature discharge to the appropriate Utilization and Quality Control Peer Review Organization. At a minimum:

   (i) The hospital must establish a clearly explained procedure for the submission of a patient’s written or verbal grievance to the hospital.
   (ii) The grievance process must specify time frames for review of the grievance and the provision of a response.
   (iii) In its resolution of the grievance, the hospital must provide the patient with written notice of its decision that contains the name of the hospital contact person, the steps taken on behalf of the patient to investigate the
grievance, the results of the grievance process, and the date of completion.

(b) Standard: Exercise of rights. (1) The patient has the right to participate in the development and implementation of his or her plan of care.

(2) The patient or his or her representative (as allowed under State law) has the right to make informed decisions regarding his or her care. The patient’s rights include being informed of his or her health status, being involved in care planning and treatment, and being able to refuse treatment. This right must not be construed as a mechanism to demand the provision of treatment or services deemed medically unnecessary or inappropriate.

(3) The patient has the right to formulate advance directives and to have hospital staff and practitioners who provide care in the hospital comply with these directives, in accordance with § 489.100 of this part (Definition), § 489.102 of this part (Requirements for providers), and § 489.104 of this part (Effective dates).

(4) The patient has the right to have a family member or representative of his or her choice and his or her own physician notified promptly of his or her admission to the hospital.

(c) Standard: Privacy and safety. (1) The patient has the right to personal privacy.

(2) The patient has the right to receive care in a safe setting.

(3) The patient has the right to be free from all forms of abuse or harassment.

(d) Standard: Confidentiality of patient records. (1) The patient has the right to the confidentiality of his or her clinical records.

(2) The patient has the right to access information contained in his or her clinical records within a reasonable time frame. The hospital must not frustrate the legitimate efforts of individuals to gain access to their own medical records and must actively seek to meet these requests as quickly as its recordkeeping system permits.

(e) Standard: Restraint for acute medical and surgical care. (1) The patient has the right to be free from restraints of any form that are not medically necessary or are used as a means of coercion, discipline, convenience, or retaliation by staff. The term “restraint” includes either a physical restraint or a drug that is being used as a restraint. A physical restraint is any manual method or physical or mechanical device, material, or equipment attached or adjacent to the patient’s body that he or she cannot easily remove that restricts freedom of movement or normal access to one’s body. A drug used as a restraint is a medication used to control behavior or to restrict the patient’s freedom of movement and is not a standard treatment for the patient’s medical or psychiatric condition.

(2) A restraint can only be used if needed to improve the patient’s well-being and less restrictive interventions have been determined to be ineffective.

(3) The use of a restraint must be—

(i) Selected only when other less restrictive measures have been found to be ineffective to protect the patient or others from harm;

(ii) In accordance with the order of a physician or other licensed independent practitioner permitted by the State and hospital to order a restraint. This order must—

(A) Never be written as a standing or on an as needed basis (that is, PRN); and

(B) Be followed by consultation with the patient’s treating physician, as soon as possible, if the restraint is not ordered by the patient’s treating physician;

(iii) In accordance with a written modification to the patient’s plan of care;

(iv) Implemented in the least restrictive manner possible;

(v) In accordance with safe and appropriate restraining techniques; and

(vi) Ended at the earliest possible time.

(4) The condition of the restrained patient must be continually assessed, monitored, and reevaluated.

(5) All staff who have direct patient contact must have ongoing education and training in the proper and safe use of restraints.

(f) Standard: Seclusion and restraint for behavior management. (1) The patient has the right to be free from seclusion and restraints, of any form, imposed as
§ 482.21 Condition of participation: Quality assurance.

(a) Standard: Clinical plan. The organized, hospital-wide quality assurance program must be ongoing and have a written plan of implementation.

A means of coercion, discipline, convenience, or retaliation by staff. The term “restraint” includes either a physical restraint or a drug that is being used as a restraint. A physical restraint is any manual method or physical or mechanical device, material, or equipment attached or adjacent to the patient’s body that he or she cannot easily remove that restricts freedom of movement or normal access to one’s body. A drug used as a restraint is a medication used to control behavior or to restrict the patient’s freedom of movement and is not a standard treatment for the patient’s medical or psychiatric condition. Seclusion is the involuntary confinement of a person in a room or an area where the person is physically prevented from leaving.

(2) Seclusion or a restraint can only be used in emergency situations if needed to ensure the patient’s physical safety and less restrictive interventions have been determined to be ineffective.

(3) The use of a restraint or seclusion must be—

(i) Selected only when less restrictive measures have been found to be ineffective to protect the patient or others from harm;

(ii) In accordance with the order of a physician or other licensed independent practitioner permitted by the State and hospital to order seclusion or restraint. The following requirements will be superseded by existing State laws that are more restrictive:

(A) Orders for the use of seclusion or a restraint must never be written as a standing order or on an as needed basis (that is, PRN).

(B) The treating physician must be consulted as soon as possible, if the restraint or seclusion is not ordered by the patient’s treating physician.

(C) A physician or other licensed independent practitioner must see and evaluate the need for restraint or seclusion within 1 hour after the initiation of this intervention.

(D) Each written order for a physical restraint or seclusion is limited to 4 hours for adults; 2 hours for children and adolescents ages 9 to 17; or 1 hour for patients under 9. The original order may only be renewed in accordance with these limits for up to a total of 24 hours. After the original order expires, a physician or licensed independent practitioner (if allowed under State law) must see and assess the patient before issuing a new order.

(iii) In accordance with a written modification to the patient’s plan of care;

(iv) Implemented in the least restrictive manner possible;

(v) In accordance with safe appropriate restraining techniques; and

(vi) Ended at the earliest possible time.

(4) A restraint and seclusion may not be used simultaneously unless the patient is—

(i) Continually monitored face-to-face by an assigned staff member; or

(ii) Continually monitored by staff using both video and audio equipment. This monitoring must be in close proximity the patient.

(5) The condition of the patient who is in a restraint or in seclusion must continually be assessed, monitored, and reevaluated.

(6) All staff who have direct patient contact must have ongoing education and training in the proper and safe use of seclusion and restraint application and techniques and alternative methods for handling behavior, symptoms, and situations that traditionally have been treated through the use of restraints or seclusion.

(7) The hospital must report to HCFA any death that occurs while a patient is restrained or in seclusion, or where it is reasonable to assume that a patient’s death is a result of restraint or seclusion.

[64 FR 36088, July 2, 1999]
§ 482.22 Condition of participation: Medical staff.

The hospital must have an organized medical staff that operates under bylaws approved by the governing body and is responsible for the quality of medical care provided to patients by the hospital.

(a) Standard: Composition of the medical staff. The medical staff must be composed of doctors of medicine or osteopathy and, in accordance with State law, may also be composed of other practitioners appointed by the governing body.

(1) The medical staff must periodically conduct appraisals of its members.

(2) The medical staff must examine credentials of candidates for medical staff membership and make recommendations to the governing body on the appointment of the candidates.

(b) Standard: Medical staff organization and accountability. The medical staff must be well organized and accountable to the governing body for the quality of the medical care provided to patients.

(1) The medical staff must be organized in a manner approved by the governing body.

(2) If the medical staff has an executive committee, a majority of the members of the committee must be doctors of medicine or osteopathy.

(3) The responsibility for organization and conduct of the medical staff must be assigned only to an individual doctor of medicine or osteopathy or, when permitted by State law of the State in which the hospital is located, a doctor of dental surgery or dental medicine.

(c) Standard: Medical staff bylaws. The medical staff must adopt and enforce bylaws to carry out its responsibilities. The bylaws must:

(1) Be approved by the governing body.

(2) Include a statement of the duties and privileges of each category of medical staff (e.g., active, courtesy, etc.)

(3) Describe the organization of the medical staff.

(d) Standard: Autopsies. The medical staff should attempt to secure autopsies in all cases of unusual deaths and of medical-legal and educational interest. The mechanism for documenting permission to perform an autopsy must be defined. There must be a system for notifying the medical staff, and specifically the attending practitioner, when an autopsy is being performed.

§ 482.23 Condition of participation: Nursing services.

The hospital must have an organized nursing service that provides 24-hour nursing services. The nursing services must be furnished or supervised by a registered nurse.

(a) Standard: Organization. The hospital must have a well-organized service with a plan of administrative authority and delineation of responsibilities for patient care. The director of the nursing service must be a licensed registered nurse. He or she is responsible for the operation of the service, including determining the types and numbers of nursing personnel and staff necessary to provide nursing care for all areas of the hospital.

(b) Standard: Staffing and delivery of care. The nursing service must have adequate numbers of licensed registered nurses, licensed practical (vocational) nurses, and other personnel to provide nursing care to all patients as needed. There must be supervisory and staff personnel for each department or nursing unit to ensure, when needed, the immediate availability of a registered nurse for bedside care of any patient.

(1) The hospital must provide 24-hour nursing services furnished or supervised by a registered nurse, and have a licensed practical nurse or registered nurse on duty at all times, except for rural hospitals that have in effect a 24-hour nursing waiver granted under §405.1910(c) of this chapter.

(2) The nursing service must have a procedure to ensure that hospital nursing personnel for whom licensure is required have valid and current licensure.

(3) A registered nurse must supervise and evaluate the nursing care for each patient.

(4) The hospital must ensure that the nursing staff develops, and keeps current, a nursing care plan for each patient.

(5) A registered nurse must assign the nursing care of each patient to other nursing personnel in accordance with the patient's needs and the specialized qualifications and competence of the nursing staff available.

(6) Non-employee licensed nurses who are working in the hospital must adhere to the policies and procedures of the hospital. The director of nursing service must provide for the adequate supervision and evaluation of the clinical activities of non-employee nursing personnel which occur within the responsibility of the nursing service.

(c) Standard: Preparation and administration of drugs. Drugs and biologicals must be prepared and administered in accordance with Federal and State laws, the orders of the practitioner or practitioners responsible for the patient's care as specified under §482.12(c), and accepted standards of practice.

(1) All drugs and biologicals must be administered by, or under supervision of, nursing or other personnel in accordance with Federal and State laws and regulations, including applicable licensing requirements, and in accordance with the approved medical staff policies and procedures.

(2) All orders for drugs and biologicals must be in writing and signed by the practitioner or practitioners responsible for the care of the patient as specified under §482.12(c). When telephone or oral orders must be used, they must be—

(i) Accepted only by personnel that are authorized to do so by the medical staff policies and procedures, consistent with Federal and State law;

(ii) Signed or initialed by the prescribing practitioner as soon as possible; and

(iii) Used infrequently.

(3) Blood transfusions and intravenous medications must be administered in accordance with State law and approved medical staff policies and procedures. If blood transfusions and intravenous medications are administered by personnel other than doctors of medicine or osteopathy, the personnel must have special training for this duty.

(4) There must be a hospital procedure for reporting transfusion reactions, adverse drug reactions, and errors in administration of drugs.

§ 482.24 Condition of participation: Medical record services.

The hospital must have a medical record service that has administrative responsibility for medical records. A
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The medical record must be maintained for every individual evaluated or treated in the hospital.

(a) Standard: Organization and staffing. The organization of the medical record service must be appropriate to the scope and complexity of the services performed. The hospital must employ adequate personnel to ensure prompt completion, filing, and retrieval of records.

(b) Standard: Form and retention of record. The hospital must maintain a medical record for each inpatient and outpatient. Medical records must be accurately written, promptly completed, properly filed and retained, and accessible. The hospital must use a system of author identification and record maintenance that ensures the integrity of the authentification and protects the security of all record entries.

(1) Medical records must be retained in their original or legally reproduced form for a period of at least 5 years.

(2) The hospital must have a system of coding and indexing medical records. The system must allow for timely retrieval by diagnosis and procedure, in order to support medical care evaluation studies.

(3) The hospital must have a procedure for ensuring the confidentiality of patient records. Information from or copies of records may be released only to authorized individuals, and the hospital must ensure that unauthorized individuals cannot gain access to or alter patient records. Original medical records must be released by the hospital only in accordance with Federal or State laws, court orders, or subpoenas.

(c) Standard: Content of record. The medical record must contain information to justify admission and continued hospitalization, support the diagnosis, and describe the patient’s progress and response to medications and services.

(ii) Authentication may include signatures, written initials or computer entry.

(2) All records must document the following, as appropriate:

(i) Evidence of a physical examination, including a health history, performed no more than 7 days prior to admission or within 48 hours after admission.

(ii) Admitting diagnosis.

(iii) Results of all consultative evaluations of the patient and appropriate findings by clinical and other staff involved in the care of the patient.

(iv) Documentation of complications, hospital acquired infections, and unfavorable reactions to drugs and anesthesia.

(v) Properly executed informed consent forms for procedures and treatments specified by the medical staff, or by Federal or State law if applicable, to require written patient consent.

(vi) All practitioners’ orders, nursing notes, reports of treatment, medication records, radiology, and laboratory reports, and vital signs and other information necessary to monitor the patient’s condition.

(vii) Discharge summary with outcome of hospitalization, disposition of case, and provisions for follow-up care.

(viii) Final diagnosis with completion of medical records within 30 days following discharge.

§ 482.25 Condition of participation: Pharmaceutical services.

The hospital must have pharmaceutical services that meet the needs of the patients. The institution must have a pharmacy directed by a registered pharmacist or a drug storage area under competent supervision. The medical staff is responsible for developing policies and procedures that minimize drug errors. This function may be delegated to the hospital’s organized pharmaceutical service.

(a) Standard: Pharmacy management and administration. The pharmacy or drug storage area must be administered in accordance with accepted professional principles.

(i) A full-time, part-time, or consulting pharmacist must be responsible
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§ 482.26 Condition of participation: Radiologic services.

The hospital must maintain, or have available, diagnostic radiologic services. If therapeutic services are also provided, they, as well as the diagnostic services, must meet professionally approved standards for safety and personnel qualifications.

(a) Standard: Radiologic services. The hospital must maintain, or have available, radiologic services according to needs of the patients.

(b) Standard: Safety for patients and personnel. The radiologic services, particularly ionizing radiology procedures, must be free from hazards for patients and personnel.

(1) Proper safety precautions must be maintained against radiation hazards. This includes adequate shielding for patients, personnel, and facilities, as well as appropriate storage, use, and disposal of radioactive materials.

(2) Periodic inspection of equipment must be made and hazards identified must be promptly corrected.

(3) Radiation workers must be checked periodically, by the use of exposure meters or badge tests, for amount of radiation exposure.

(4) Radiologic services must be provided only on the order of practitioners with clinical privileges or, consistent with State law, of other practitioners authorized by the medical staff and the governing body to order the services.

(c) Standard: Personnel. (1) A qualified full-time, part-time, or consulting radiologist must supervise the ionizing radiology services and must interpret only those radiologic tests that are determined by the medical staff to require a radiologist's specialized knowledge. For purposes of this section, a radiologist is a doctor of medicine or osteopathy who is qualified by education and experience in radiology.

(2) Only personnel designated as qualified by the medical staff may use the radiologic equipment and administer procedures.
(d) Standard: Records. Records of radiologic services must be maintained.

(1) The radiologist or other practitioner who performs radiology services must sign reports of his or her interpretations.

(2) The hospital must maintain the following for at least 5 years:
   (i) Copies of reports and printouts.
   (ii) Films, scans, and other image records, as appropriate.

§ 482.27 Condition of participation: Laboratory services.

(a) The hospital must maintain, or have available, adequate laboratory services to meet the needs of its patients. The hospital must ensure that all laboratory services provided to its patients are performed in a facility certified in accordance with part 493 of this chapter.

(b) Standard: Adequacy of laboratory services. The hospital must have laboratory services available, either directly or through a contractual agreement with a certified laboratory that meets requirements of part 493 of this chapter.

(1) Emergency laboratory services must be available 24 hours a day.

(2) A written description of services provided must be available to the medical staff.

(3) The laboratory must make provision for proper receipt and reporting of tissue specimens.

(4) The medical staff and a pathologist must determine which tissue specimens require a macroscopic (gross) examination and which require both macroscopic and microscopic examinations.

(c) Standard: Potentially infectious blood and blood products—(1) Potentially HIV infectious blood and blood products are prior collections from a donor who tested negative at the time of donation but tests repeatedly reactive for the antibody to the human immunodeficiency virus (HIV) on a later donation, and the FDA-licensed, more specific test or other followup testing recommended or required by FDA completed within 30 calendar days after the donor's repeatedly reactive screening test. (FDA regulations concerning HIV testing and lookback procedures are set forth at 21 CFR 610.45 et seq.)

(2) Services furnished by an outside blood bank. If a hospital regularly uses the services of an outside blood bank, it must have an agreement with the blood bank that governs the procurement, transfer, and availability of blood and blood products. The agreement must require that the blood bank promptly notify the hospital of the following:
   (i) If it supplied blood and blood products collected from a donor who tested negative at the time of donation but tests repeatedly reactive for the antibody to HIV on a later donation; and
   (ii) The results of the FDA-licensed, more specific test or other followup testing recommended or required by FDA completed within 30 calendar days after the donor's repeatedly reactive screening test. (FDA regulations concerning HIV testing and lookback procedures are set forth at 21 CFR 610.45 et seq.)

(3) Quarantine of blood and blood products pending completion of testing. If the blood bank notifies the hospital of the repeatedly reactive HIV screening test results as required by paragraph (c)(2)(i) of this section, the hospital must determine the disposition of the blood or blood product and quarantine all blood and blood products from previous donations in inventory.

   (i) If the blood bank notifies the hospital that the result of the FDA-licensed, more specific test or other followup testing recommended or required by FDA is negative, absent other informative test results, the hospital may release the blood and blood products from quarantine.

   (ii) If the blood bank notifies the hospital that the result of the FDA-licensed, more specific test or other followup testing recommended or required by FDA is positive, the hospital must dispose of the blood and blood products in accordance with 21 CFR 606.40 and notify patients in accordance with paragraph (c)(4) of this section.

(4) Patient notification. If the hospital has administered potentially HIV infectious blood or blood products (either directly through its own blood bank or under an agreement described in paragraph (c)(2) of this section) or released such blood or blood products to another entity or appropriate individual,
the hospital must take the following actions:

(i) Promptly make at least three attempts to notify the patient's attending physician (that is, the physician of record) or the physician who ordered the blood or blood product that potentially HIV infectious blood or blood products were transfused to the patient.

(ii) Ask the physician to immediately notify the patient, or other individual as permitted under paragraph (c)(8) of this section, of the need for HIV testing and counseling.

(iii) If the physician is unavailable, declines to make the notification, or later informs the hospital that he or she was unable to notify the patient, promptly make at least three attempts to notify the patient, or other individual as permitted under paragraph (c)(8) of this section, of the need for HIV testing and counseling.

(iv) Document in the patient's medical record the notification or attempts to give the required notification.

(5) Timeframe for notification. The notification effort begins when the blood bank notifies the hospital that it received potentially HIV infectious blood and blood products and continues for 8 weeks unless—

(i) The patient is located and notified; or

(ii) The hospital is unable to locate the patient and documents in the patient's medical record the extenuating circumstances beyond the hospital's control that caused the notification timeframe to exceed 8 weeks.

(6) Content of notification. The notification given under paragraphs (c)(4) (ii) and (iii) of this section must include the following information:

(i) A basic explanation of the need for HIV testing and counseling.

(ii) Enough oral or written information so that the transfused patient can make an informed decision about whether to obtain HIV testing and counseling.

(iii) A list of programs or places where the patient can obtain HIV testing and counseling, including any requirements or restrictions the program may impose.

(7) Policies and procedures. The hospital must establish policies and procedures for notification and documentation that conform to Federal, State, and local laws, including requirements for confidentiality and medical records.

(8) Notification to legal representative or relative. If the patient has been adjudged incompetent by a State court, the physician or hospital must notify a legal representative designated in accordance with State law. If the patient is competent, but State law permits a legal representative or relative to receive the information on the patient's behalf, the physician or hospital must notify the patient or his or her legal representative or relative. If the patient is deceased, the physician or hospital must continue the notification process and inform the deceased patient's legal representative or relative.

[57 FR 7136, Feb. 28, 1992, as amended at 61 FR 47433, Sept. 9, 1996]

§ 482.28 Condition of participation: Food and dietetic services.

The hospital must have organized dietary services that are directed and staffed by adequate qualified personnel. However, a hospital that has a contract with an outside food management company may be found to meet this Condition of participation if the company has a dietitian who serves the hospital on a full-time, part-time, or consultant basis, and if the company maintains at least the minimum standards specified in this section and provides for constant liaison with the hospital medical staff for recommendations on dietetic policies affecting patient treatment.

(a) Standard: Organization. (1) The hospital must have a full-time employee who—

(i) Serves as director of the food and dietetic service;

(ii) Is responsible for the daily management of the dietary services; and

(iii) Is qualified by experience or training.

(2) There must be a qualified dietitian, full-time, part-time, or on a consultant basis.

(3) There must be administrative and technical personnel competent in their respective duties.

(b) Standard: Diets. Menus must meet the needs of the patients.
(1) Therapeutic diets must be prescribed by the practitioner or practitioners responsible for the care of the patients.

(2) Nutritional needs must be met in accordance with recognized dietary practices and in accordance with orders of the practitioner or practitioners responsible for the care of the patients.

(3) A current therapeutic diet manual approved by the dietitian and medical staff must be readily available to all medical, nursing, and food service personnel.

§ 482.30 Condition of participation: Utilization review.

The hospital must have in effect a utilization review (UR) plan that provides for review of services furnished by the institution and by members of the medical staff to patients entitled to benefits under the Medicare and Medicaid programs.

(a) Applicability. The provisions of this section apply except in either of the following circumstances:

(1) A Utilization and Quality Control Peer Review Organization (PRO) has assumed binding review for the hospital.

(2) HCFA has determined that the UR procedures established by the State under title XIX of the Act are superior to the procedures required in this section, and has required hospitals in that State to meet the UR plan requirements under §§ 456.50 through 456.245 of this chapter.

(b) Standard: Composition of utilization review committee. A UR committee consisting of two or more practitioners must carry out the UR function. At least two of the members of the committee must be doctors of medicine or osteopathy. The other members may be any of the other types of practitioners specified in § 482.12(c)(1).

(B) Established in a manner approved by HCFA.

(2) If, because of the small size of the institution, it is impracticable to have a properly functioning staff committee, the UR committee must be established as specified in paragraph (b)(1)(ii) of this section.

(3) The committee’s or group’s reviews may not be conducted by any individual who—

(i) Has a direct financial interest (for example, an ownership interest) in that hospital; or

(ii) Was professionally involved in the care of the patient whose case is being reviewed.

(c) Standard: Scope and frequency of review. (1) The UR plan must provide for review for Medicare and Medicaid patients with respect to the medical necessity of—

(i) Admissions to the institution; (ii) The duration of stays; and (iii) Professional services furnished, including drugs and biologicals.

(2) Review of admissions may be performed before, at, or after hospital admission.

(3) Except as specified in paragraph (e) of this section, reviews may be conducted on a sample basis.

(4) Hospitals that are paid for inpatient hospital services under the prospective payment system set forth in Part 412 of this chapter must conduct review of duration of stays and review of professional services as follows:

(i) For duration of stays, these hospitals need review only cases that they reasonably assume to be outlier cases based on extended length of stay, as described in § 412.80(a)(1)(i) of this chapter; and

(ii) For professional services, these hospitals need review only cases that they reasonably assume to be outlier cases based on extraordinarily high costs, as described in § 412.80(a)(1)(ii) of this chapter.

(d) Standard: Determination regarding admissions or continued stays. (1) The determination that an admission or continued stay is not medically necessary—

(i) May be made by one member of the UR committee if the practitioner or practitioners responsible for the care of the patient, as specified of
§ 482.12(c), concur with the determination or fail to present their views when afforded the opportunity; and
(ii) Must be made by at least two members of the UR committee in all other cases.
(2) Before making a determination that an admission or continued stay is not medically necessary, the UR committee must consult the practitioner or practitioners responsible for the care of the patient, as specified in §482.12(c), and afford the practitioner or practitioners the opportunity to present their views.
(3) If the committee decides that admission to or continued stay in the hospital is not medically necessary, written notification must be given, no later than 2 days after the determination, to the hospital, the patient, and the practitioner or practitioners responsible for the care of the patient, as specified in §482.12(c);

(e) Standard: Extended stay review. (1) In hospitals that are not paid under the prospective payment system, the UR committee must make a periodic review, as specified in the UR plan, of each current inpatient receiving hospital services during a continuous period of extended duration. The scheduling of the periodic reviews may—
(i) Be the same for all cases; or
(ii) Differ for different classes of cases.
(2) In hospitals paid under the prospective payment system, the UR committee must review all cases reasonably assumed by the hospital to be outlier cases because the extended length of stay exceeds the threshold criteria for the diagnosis, as described in §412.80(a)(1)(i). The hospital is not required to review an extended stay that does not exceed the outlier threshold for the diagnosis.
(3) The UR committee must make the periodic review no later than 7 days after the day required in the UR plan.

(f) Standard: Review of professional services. The committee must review professional services provided, to determine medical necessity and to promote the most efficient use of available health facilities and services.

§ 482.41 Condition of participation: Physical environment.

The hospital must be constructed, arranged, and maintained to ensure the safety of the patient, and to provide facilities for diagnosis and treatment and for special hospital services appropriate to the needs of the community.

(a) Standard: Buildings. The condition of the physical plant and the overall hospital environment must be developed and maintained in such a manner that the safety and well-being of patients are assured.

(1) There must be emergency power and lighting in at least the operating, recovery, intensive care, and emergency rooms, and stairwells. In all other areas not serviced by the emergency supply source, battery lamps and flashlights must be available.

(2) There must be facilities for emergency gas and water supply.

(b) Standard: Life safety from fire. (1) Except as provided in paragraphs (b)(1)(i) through (b)(1)(iii) of this section, the hospital must meet the applicable provisions of the 1985 edition of the Life Safety Code of the National Fire Protection Association (which is incorporated by reference).1

(i) Any hospital that on November 26, 1982, complied, with or without waivers, with the requirements of the 1967 edition of the Life Safety Code, or on May 9, 1988, complied with the 1981 edition of the Life Safety Code, is considered to be in compliance with this standard as long as the facility continues to remain in compliance with that edition of the Code.

(ii) After consideration of State survey agency findings, HCFA may waive specific provisions of the Life Safety Code which, if rigidly applied, would result in unreasonable hardship upon the facility, but only if the waiver does not adversely affect the health and safety of patients.

(iii) The provisions of the Life Safety Code do not apply in a State where HCFA finds that a fire and safety code imposed by State law adequately protects patients in hospitals.

1See footnote to §405.1134(a) of this chapter.
(2) The hospital must have procedures for the proper routine storage and prompt disposal of trash.

(3) The hospital must have written fire control plans that contain provisions for prompt reporting of fires; extinguishing fires; protection of patients, personnel and guests; evacuation; and cooperation with fire fighting authorities.

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

(c) Standard: Facilities. The hospital must maintain adequate facilities for its services.

(1) Diagnostic and therapeutic facilities must be located for the safety of patients.

(2) Facilities, supplies, and equipment must be maintained to ensure an acceptable level of safety and quality.

(3) The extent and complexity of facilities must be determined by the services offered.

(4) There must be proper ventilation, light, and temperature controls in pharmaceutical, food preparation, and other appropriate areas.

§ 482.43 Condition of participation: Discharge planning.

(a) Standard: Identification of patients in need of discharge planning. The hospital must identify at an early stage of hospitalization all patients who are likely to suffer adverse health consequences upon discharge if there is no adequate discharge planning.

(b) Standard: Responsibilities of chief executive officer, medical staff, and director of nursing services. The chief executive officer, the medical staff, and the director of nursing services must—

(1) Ensure that the hospital-wide quality assurance program and training programs address problems identified by the infection control officer or officers; and

(2) Be responsible for the implementation of successful corrective action plans in affected problem areas.

§ 482.42 Condition of participation: Infection control.

The hospital must provide a sanitary environment to avoid sources and transmission of infections and communicable diseases. There must be an active program for the prevention, control, and investigation of infections and communicable diseases.

(a) Standard: Organization and policies. A person or persons must be designated as infection control officer or officers to develop and implement policies governing control of infections and communicable diseases.

(1) The infection control officer or officers must develop a system for identifying, reporting, investigating, and controlling infections and communicable diseases of patients and personnel.

(2) The infection control officer or officers must maintain a log of incidents related to infections and communicable diseases.
§ 482.45 Condition of participation: Organ, tissue, and eye procurement.

(a) Standard: Organ procurement responsibilities. The hospital must have and implement written protocols that:

(1) Incorporate an agreement with an OPO designated under part 486 of this chapter, under which it must notify, in a timely manner, the OPO or a third party designated by the OPO of individuals whose death is imminent or who have died in the hospital. The OPO determines medical suitability for organ donation and, in the absence of alternative arrangements by the hospital, the OPO determines medical suitability for tissue and eye donation, using the definition of potential tissue and eye donor and the notification protocol developed in consultation with the tissue and eye banks identified by the hospital for this purpose;

(2) Incorporate an agreement with at least one tissue bank and at least one eye bank to cooperate in the retrieval, processing, preservation, storage and distribution of tissues and eyes, as may be appropriate to assure that all usable tissues and eyes are obtained from potential donors, insofar as such an agreement does not interfere with organ procurement;

(3) Ensure, in collaboration with the designated OPO, that the family of each potential donor is informed of its options to donate organs, tissues, or eyes or to decline to donate. The individual designated by the hospital to initiate the request to the family must be an organ procurement representative or a designated requestor. A designated requestor is an individual who has completed a course offered or approved by the OPO and designed in conjunction with the tissue and eye bank community in the methodology for approaching potential donor families and requesting organ or tissue donation;

(4) Encourage discretion and sensitivity with respect to the circumstances, views, and beliefs of the families of potential donors;

(5) Ensure that the hospital works cooperatively with the designated OPO, tissue bank and eye bank in educating staff on donation issues, reviewing death records to improve identification of potential donors, and maintaining potential donors while necessary testing and placement of potential donated organs, tissues, and eyes take place.

(b) Standard: Organ transplantation responsibilities. (1) A hospital in which organ transplants are performed must be a member of the Organ Procurement and Transplantation Network (OPTN) established and operated in accordance with section 372 of the Public Health Service (PHS) Act (42 U.S.C. 274) and abide by its rules. The term "rules of the OPTN" means those rules provided
for in regulations issued by the Secretary in accordance with section 372 of the PHS Act which are enforceable under 42 CFR 121.10. No hospital is considered to be out of compliance with section 1138(a)(1)(B) of the Act, unless the Secretary has given the OPTN formal notice that he or she approves the decision to exclude the hospital from the OPTN and has notified the hospital in writing.

(2) For purposes of these standards, the term "organ" means a human kidney, liver, heart, lung, or pancreas.

(3) If a hospital performs any type of transplants, it must provide organ-transplant-related data, as requested by the OPTN, the Scientific Registry, and the OPOs. The hospital must also provide such data directly to the Department when requested by the Secretary.

[63 FR 33875, June 22, 1998]

Subpart D—Optional Hospital Services

§ 482.51 Condition of participation: Surgical services.

If the hospital provides surgical services, the services must be well organized and provided in accordance with acceptable standards of practice. If outpatient surgical services are offered the services must be consistent in quality with inpatient care in accordance with the complexity of services offered.

(a) Standard: Organization and staffing. The organization of the surgical services must be appropriate to the scope of the services offered.

(1) The operating rooms must be supervised by an experienced registered nurse or a doctor of medicine or osteopathy.

(2) Licensed practical nurses (LPNs) and surgical technologists (operating room technicians) may serve as "scrub nurses" under the supervision of a registered nurse.

(3) Qualified registered nurses may perform circulating duties in the operating room. In accordance with applicable State laws and approved medical staff policies and procedures, LPNs and surgical technologists may assist in circulatory duties under the supervision of a qualified registered nurse who is immediately available to respond to emergencies.

(4) Surgical privileges must be delineated for all practitioners performing surgery in accordance with the competencies of each practitioner. The surgical service must maintain a roster of practitioners specifying the surgical privileges of each practitioner.

(b) Standard: Delivery of service. Surgical services must be consistent with needs and resources. Policies governing surgical care must be designed to assure the achievement and maintenance of high standards of medical practice and patient care.

(1) There must be a complete history and physical work-up in the chart of every patient prior to surgery, except in emergencies. If this has been dictated, but not yet recorded in the patient's chart, there must be a statement to that effect and an admission note in the chart by the practitioner who admitted the patient.

(2) A properly executed informed consent form for the operation must be in the patient's chart before surgery, except in emergencies.

(3) The following equipment must be available to the operating room suites: call-in-system, cardiac monitor, resuscitator, defibrillator, aspirator, and tracheotomy set.

(4) There must be adequate provisions for immediate post-operative care.

(5) The operating room register must be complete and up-to-date.

(6) An operative report describing techniques, findings, and tissues removed or altered must be written or dictated immediately following surgery and signed by the surgeon.

§ 482.52 Condition of participation: Anesthesia services.

If the hospital furnishes anesthesia services, they must be provided in a well-organized manner under the direction of a qualified doctor of medicine or osteopathy. The service is responsible for all anesthesia administered in the hospital.

(a) Standard: Organization and staffing. The organization of anesthesia services must be appropriate to the
scope of the services offered. Anesthesia must be administered by only—
(1) A qualified anesthesiologist:
(2) A doctor of medicine or osteopathy (other than an anesthesiologist);
(3) A dentist, oral surgeon, or podiatrist who is qualified to administer anesthesia under State law;
(4) A certified registered nurse anesthetist (CRNA), as defined in §410.69(b) of this chapter, who is under the supervision of the operating practitioner or of an anesthesiologist who is immediately available if needed; or
(5) An anesthesiologist’s assistant, as defined in §410.69(b) of this chapter, who is under the supervision of an anesthesiologist who is immediately available if needed.

(b) Standard: Delivery of services. Anesthesia services must be consistent with needs and resources. Policies on anesthesia procedures must include the delineation of preanesthesia and post anesthesia responsibilities. The policies must ensure that the following are provided for each patient:
(1) A preanesthesia evaluation by an individual qualified to administer anesthesia under paragraph (a) of this section performed within 48 hours prior to surgery.
(2) An intraoperative anesthesia record.
(3) With respect to inpatients, a postanesthesia followup report by the individual who administers the anesthesia that is written within 48 hours after surgery.
(4) With respect to outpatients, a postanesthesia evaluation for proper anesthesia recovery performed in accordance with policies and procedures approved by the medical staff.

§ 482.53 Condition of participation: Nuclear medicine services.

If the hospital provides nuclear medicine services, those services must meet the needs of the patients in accordance with acceptable standards of practice.

(a) Standard: Organization and staffing. The organization of the nuclear medicine service must be appropriate to the scope and complexity of the services offered.

(1) There must be a director who is a doctor of medicine or osteopathy qualified in nuclear medicine.

(2) The qualifications, training, functions, and responsibilities of nuclear medicine personnel must be specified by the service director and approved by the medical staff.

(b) Standard: Delivery of service. Radioactive materials must be prepared, labeled, used, transported, stored, and disposed of in accordance with acceptable standards of practice.

(1) In-house preparation of radiopharmaceuticals is by, or under, the direct supervision of an appropriately trained registered pharmacist or a doctor of medicine or osteopathy.

(2) There is proper storage and disposal of radioactive material.

(3) If laboratory tests are performed in the nuclear medicine service, the service must meet the applicable requirement for laboratory services specified in §482.27.

(c) Standard: Facilities. Equipment and supplies must be appropriate for the types of nuclear medicine services offered and must be maintained for safe and efficient performance. The equipment must be—
(1) Maintained in safe operating condition; and
(2) Inspected, tested, and calibrated at least annually by qualified personnel.

(d) Standard: Records. The hospital must maintain signed and dated reports of nuclear medicine interpretations, consultations, and procedures.

(1) The hospital must maintain copies of nuclear medicine reports for at least 5 years.

(2) The practitioner approved by the medical staff to interpret diagnostic procedures must sign and date the interpretation of these tests.

(3) The hospital must maintain records of the receipt and disposition of radiopharmaceuticals.

(4) Nuclear medicine services must be ordered only by practitioner whose scope of Federal or State licensure and whose defined staff privileges allow such referrals.

[51 FR 22042, June 17, 1986, as amended at 57 FR 7136, Feb. 28, 1992]
§ 482.54 Condition of participation: Outpatient services.

If the hospital provides outpatient services, the services must meet the needs of the patients in accordance with acceptable standards of practice.

(a) Standard: Organization. Outpatient services must be appropriately organized and integrated with inpatient services.

(b) Standard: Personnel. The hospitals must—

1. Assign an individual to be responsible for outpatient services; and
2. Have appropriate professional and nonprofessional personnel available.

§ 482.55 Condition of participation: Emergency services.

The hospital must meet the emergency needs of patients in accordance with acceptable standards of practice.

(a) Standard: Organization and direction. If emergency services are provided at the hospital—

1. The services must be organized under the direction of a qualified member of the medical staff;
2. The services must be integrated with other departments of the hospital;
3. The policies and procedures governing medical care provided in the emergency service or department are established by and are a continuing responsibility of the medical staff.

(b) Standard: Personnel. (1) The emergency services must be supervised by a qualified member of the medical staff.

2. There must be adequate medical and nursing personnel qualified in emergency care to meet the written emergency procedures and needs anticipated by the facility.

§ 482.56 Condition of participation: Rehabilitation services.

If the hospital provides rehabilitation, physical therapy, occupational therapy, audiology, or speech pathology services, the services must be organized and staffed to ensure the health and safety of patients.

(a) Standard: Organization and staffing. The organization of the service must be appropriate to the scope of the services offered.

1. The director of the services must have the necessary knowledge, experience, and capabilities to properly supervise and administer the services.
2. Physical therapy, occupational therapy, or speech therapy, or audiology services, if provided, must be provided by staff who meet the qualifications specified by the medical staff, consistent with State law.

(b) Standard: Delivery of services. Services must be furnished in accordance with a written plan of treatment. Services must be given in accordance with orders of practitioners who are authorized by the medical staff to order the services, and the orders must be incorporated in the patient's record.

§ 482.57 Condition of participation: Respiratory care services.

The hospital must meet the needs of the patients in accordance with acceptable standards of practice. The following requirements apply if the hospital provides respiratory care service.

(a) Standard: Organization and staffing. The organization of the respiratory care services must be appropriate to the scope and complexity of the services offered.

1. There must be a director of respiratory care services who is a doctor of medicine or osteopathy with the knowledge experience, and capabilities to supervise and administer the service properly. The director may serve on either a full-time or part-time basis.
2. There must be adequate numbers of respiratory therapists, respiratory therapy technicians, and other personnel who meet the qualifications specified by the medical staff, consistent with State law.

(b) Standard: Delivery of Services. Services must be delivered in accordance with medical staff directives.

1. Personnel qualified to perform specific procedures and the amount of supervision required for personnel to carry out specific procedures must be designated in writing.
2. If blood gases or other laboratory tests are performed in the respiratory care unit, the unit must meet the applicable requirements for laboratory services specified in §482.27.
§ 482.60 Special provisions applying to psychiatric hospitals.

Psychiatric hospital must—
(a) Be primarily engaged in providing, by or under the supervision of a doctor of medicine or osteopathy, psychiatric services for the diagnosis and treatment of mentally ill persons;
(b) Meet the conditions of participation specified in §§482.1 through 482.23 and §§482.25 through 482.57;
(c) Maintain clinical records on all patients, including records sufficient to permit HCFA to determine the degree and intensity of treatment furnished to Medicare beneficiaries, as specified in §482.61; and
(d) Meet the staffing requirements specified in §482.62.

§ 482.61 Condition of participation: Special medical record requirements for psychiatric hospitals.

The medical records maintained by a psychiatric hospital must permit determination of the degree and intensity of the treatment provided to individuals who are furnished services in the institution.

(a) Standard: 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 patient is hospitalized.

(1) The identification data must include the patient’s legal status.

(2) A provisional or admitting diagnosis must be made on every patient at the time of admission, and must include the diagnoses of intercurrent diseases as well as the psychiatric diagnoses.

(3) The reasons for admission must be clearly documented as stated by the patient and/or others significantly involved.

(4) The social service records, including reports of interviews with patients, family members, and others, must provide an assessment of home plans and family attitudes, and community resource contacts as well as a social history.

(5) When indicated, a complete neurological examination must be recorded at the time of the admission physical examination.

(b) Standard: Psychiatric evaluation. Each patient must receive a psychiatric evaluation that must—

(1) Be completed within 60 hours of admission;

(2) Include a medical history;

(3) Contain a record of mental status;

(4) Note the onset of illness and the circumstances leading to admission;

(5) Describe attitudes and behavior;

(6) Estimate intellectual functioning, memory functioning, and orientation; and

(7) Include an inventory of the patient’s assets in descriptive, not interpretative, fashion.

(c) Standard: Treatment plan. (1) Each patient must have an individual comprehensive treatment plan that must be based on an inventory of the patient’s strengths and disabilities. The written plan must include—

(i) A substantiated diagnosis;

(ii) Short-term and long-range goals;

(iii) The specific treatment modalities utilized;

(iv) The responsibilities of each member of the treatment team; and

(v) Adequate documentation to justify the diagnosis and the treatment and rehabilitation activities carried out.

(2) The treatment received by the patient must be documented in such a way to assure that all active therapeutic efforts are included.

(d) Standard: Recording progress. Progress notes must be recorded by the doctor of medicine or osteopathy responsible for the care of the patient as specified in §482.12(c), 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 patient but must be
recorded at least weekly for the first 2 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 patient’s progress in accordance with the original or revised treatment plan.

(e) Standard: 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 patient’s hospitalization and qualifications of doctors of from appropriate services concerning follow-up or aftercare as well as a brief summary of the patient’s condition on discharge.

[51 FR 22042, June 17, 1986; 51 FR 27848, Aug. 4, 1986]

§ 482.62 Condition of participation: Special staff requirements for psychiatric hospitals.

The hospital must have adequate numbers of qualified professional and supportive staff to evaluate patients, formulate written, individualized comprehensive treatment plans, provide active treatment measures, and engage in discharge planning.

(a) Standard: Personnel. The hospital must employ or undertake to provide adequate numbers of qualified professional, technical, and consultative personnel to:

(1) Evaluate patients;
(2) Formulate written individualized comprehensive treatment plans;
(3) Provide active treatment measures; and
(4) Engage in discharge planning.

(b) Standard: 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.

(1) 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.

(2) The director must monitor and evaluate the quality and appropriateness of services and treatment provided by the medical staff.

(c) Standard: Availability of medical personnel. Doctors of medicine or osteopathy and other appropriate professional personnel must be available to provide necessary medical and surgical diagnostic and treatment services. If medical and surgical diagnostic and treatment services are not available within the institution, the institution must have an agreement with an outside source of these services to ensure that they are immediately available or a satisfactory agreement must be established for transferring patients to a general hospital that participates in the Medicare program.

(d) Standard: Nursing services. The hospital 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 necessary medical and surgical diagnostic and treatment services. If medical and surgical diagnostic and treatment services are not available within the institution, the institution must have an agreement with an outside source of these services to ensure that they are immediately available or a satisfactory agreement must be established for transferring patients to a general hospital that participates in the Medicare program.

(1) 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.

(2) The staffing pattern must insure the availability of a registered professional 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 patient’s active treatment program.

(e) Standard: Psychological services. The hospital must provide or have
available psychological services to meet the needs of the patients.

(f) Standard: 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.

(1) The director of the social work department or service must have a master’s degree from an accredited school of social work or must be qualified by education and experience in the social services needs of the mentally ill. If the director does not hold a masters degree in social work, at least one staff member must have this qualification.

(2) 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.

(g) Standard: Therapeutic activities. The hospital must provide a therapeutic activities program.

(1) The program must be appropriate to the needs and interests of patients and be directed toward restoring and maintaining optimal levels of physical and psychosocial functioning.

(2) The number of qualified therapists, support personnel, and consultants must be adequate to provide comprehensive therapeutic activities consistent with each patient’s active treatment program.

§ 482.66 Special requirements for hospital providers of long-term care services ("swing-beds").

A hospital that has a Medicare provider agreement must meet the following requirements in order to be granted an approval from HCFA to provide post-hospital extended care services, as specified in §409.30 of this chapter, and be reimbursed as a swing-bed hospital, as specified in §413.114 of this chapter:

(a) Eligibility. A hospital must meet the following eligibility requirements:

(1) The facility has fewer than 100 hospital beds, excluding beds for newborns and beds in intensive care type inpatient units (for eligibility of hospitals with distinct parts electing the optional reimbursement method, see §413.24(d)(5) of this chapter).

(2) The hospital is located in a rural area. This includes all areas not delineated as “urbanized” areas by the Census Bureau, based on the most recent census.

(3) The hospital does not have in effect a 24-hour nursing waiver granted under §408.54(c) of this chapter.

(4) The hospital has not had a swing-bed approval terminated within the two years previous to application.

(b) Skilled nursing facility services. The facility is substantially in compliance with the following skilled nursing facility requirements contained in subpart B of part 483 of this chapter.

(1) Resident rights (§483.10 (b)(3), (b)(4), (b)(5), (b)(6), (d), (e), (h), (1), (j)(1)(vii), (j)(1)(viii), (1), and (m)).

(2) Admission, transfer, and discharge rights (§483.12 (a)(1), (a)(2), (a)(3), (a)(4), (a)(5), (a)(6), and (a)(7)).

(3) Resident behavior and facility practices (§483.13).

(4) Patient activities (§483.15(f)).

(5) Social services (§483.15(g)).

(6) Discharge planning (§483.20(e)).

(7) Specialized rehabilitative services (§483.45).

(8) Dental services (§483.55).

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(7) Specialized rehabilitative services (§483.45).

(8) Dental services (§483.55).

[51 FR 22042, June 17, 1986; 51 FR 27848, Aug. 4, 1986]
§ 483.1 Basis and scope.

(a) Statutory basis. (1) Sections 1819 (a), (b), (c), and (d) of the Act provide that—

(i) Skilled nursing facilities participating in Medicare must meet certain specified requirements; and

(2) Sections 1861 (a), (b), (c), (d), (e), and (f) of the Social Security Act provide that—

(i) Inpatient psychiatric facilities participating in Medicare must meet certain specified requirements; and

(ii) Hospices participating in Medicare must meet certain specified requirements.

§ 483.2 Definitions.

§ 483.30 Nursing services.

§ 483.35 Dietary services.

§ 483.40 Physician services.

§ 483.45 Specialized rehabilitative services.

§ 483.50 Dental services.

§ 483.55 Pharmacy services.

§ 483.60 Infection control.

§ 483.65 Physical environment.

§ 483.70 Administration.

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§ 483.100 Basis.

§ 483.102 Applicability and definitions.

§ 483.104 State plan requirement.

§ 483.106 Basic rule.

§ 483.108 Relationship of PASARR to other Medicaid processes.

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§ 483.150 Statutory basis; Deemed meeting or waiver of requirements.

§ 483.151 State review and approval of nurse aide training and competency evaluation programs and competency evaluation programs.

§ 483.152 Requirements for approval of a nurse aide training and competency evaluation program.

§ 483.154 Nurse aide competency evaluation.

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§ 483.315 Specification of resident assessment instrument.

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Subpart I—Conditions of Participation for Intermediate Care Facilities for the Mentally Retarded

§ 483.400 Basis and purpose.

§ 483.405 Relationship to other HHS regulations.

§ 483.410 Condition of participation: Governing body and management.

§ 483.420 Condition of participation: Client protections.

§ 483.430 Condition of participation: Facility staffing.

§ 483.440 Condition of participation: Active treatment services.

§ 483.450 Condition of participation: Client behavior and facility practices.

§ 483.460 Condition of participation: Health care services.

§ 483.470 Condition of participation: Physical environment.

§ 483.480 Condition of participation: Dietetic services.

Authority: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1396hh).

Subpart A [Reserved]

Subpart B—Requirements for Long Term Care Facilities

Source: 54 FR 5359, Feb. 2, 1989, unless otherwise noted.

§ 483.1 Basis and scope.

(a) Statutory basis. (1) Sections 1819 (a), (b), (c), and (d) of the Act provide that—

(i) Skilled nursing facilities participating in Medicare must meet certain specified requirements; and
§ 483.5 Definitions.

Facility means, a skilled nursing facility (SNF) or a nursing facility (NF) which meets the requirements of sections 1819 or 1919 (a), (b), (c), and (d) of the Act. “Facility” may include a distinct part of an institution specified in §440.40 of this chapter, but does not include an institution for the mentally retarded or persons with related conditions described in §440.150 of this chapter. For Medicare and Medicaid purposes (including eligibility, coverage, certification, and payment), the “facility” is always the entity which participates in the program, whether that entity is comprised of all of, or a distinct part of a larger institution. For Medicare, a SNF (see section 1819(a)(1)), and for Medicaid, a NF (see section 1919(a)(1)) may not be an institution for mental diseases as defined in §435.1009.

§ 483.10 Resident rights.

The resident has a right to a dignified existence, self-determination, and communication with and access to persons and services inside and outside the facility. A facility must protect and promote the rights of each resident, including each of the following rights:

(a) Exercise of rights. (1) The resident has the right to exercise his or her rights as a resident of the facility and as a citizen or resident of the United States.

(b) Scope. The provisions of this part contain the requirements that an institution must meet in order to qualify to participate as a SNF in the Medicare program, and as a nursing facility in the Medicaid program. They serve as the basis for survey activities for the purpose of determining whether a facility meets the requirements for participation in Medicare and Medicaid.

§ 483.10 Definition.

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Facility means, a skilled nursing facility (SNF) or a nursing facility (NF) which meets the requirements of sections 1819 or 1919 (a), (b), (c), and (d) of the Act. “Facility” may include a distinct part of an institution specified in §440.40 of this chapter, but does not include an institution for the mentally retarded or persons with related conditions described in §440.150 of this chapter. For Medicare and Medicaid purposes (including eligibility, coverage, certification, and payment), the “facility” is always the entity which participates in the program, whether that entity is comprised of all of, or a distinct part of a larger institution. For Medicare, a SNF (see section 1819(a)(1)), and for Medicaid, a NF (see section 1919(a)(1)) may not be an institution for mental diseases as defined in §435.1009.

(ii) The Secretary may impose additional requirements (see section 1819(d)(4)(B)) if they are necessary for the health and safety of individuals to whom services are furnished in the facilities.

(2) Section 1861(l) of the Act requires the facility to have in effect a transfer agreement with a hospital.

(3) Sections 1919 (a), (b), (c), and (d) of the Act provide that nursing facilities participating in Medicaid must meet certain specific requirements.

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(4) The resident has the right to refuse treatment, to refuse to participate in experimental research, and to formulate an advance directive as specified in paragraph (8) of this section; and

(5) The facility must—

(i) Inform each resident who is entitled to Medicaid benefits, in writing, at the time of admission to the nursing facility or, when the resident becomes eligible for Medicaid of—

(A) The items and services that are included in nursing facility services under the State plan and for which the resident may not be charged;

(B) Those other items and services that the facility offers and for which the resident may be charged, and the amount of charges for those services; and

(ii) Inform each resident when changes are made to the items and services specified in paragraphs (5)(i) (A) and (B) of this section.

(6) The facility must inform each resident before, or at the time of admission, and periodically during the resident's stay, of services available in the facility and of charges for those services, including any charges for services not covered under Medicare or by the facility's per diem rate.

(7) The facility must furnish a written description of legal rights which includes—

(i) A description of the manner of protecting personal funds, under paragraph (c) of this section;

(ii) A description of the requirements and procedures for establishing eligibility for Medicaid, including the right to request an assessment under section 1924(c) which determines the extent of a couple's non-exempt resources at the time of institutionalization and attributes to the community spouse an equitable share of resources which cannot be considered available for payment toward the cost of the institutionalized spouse's medical care in his or her process of spending down to Medicaid eligibility levels;

(iii) A posting of names, addresses, and telephone numbers of all pertinent State client advocacy groups such as the State survey and certification agency, the State licensure office, the State ombudsman program, the protection and advocacy network, and the Medicaid fraud control unit; and

(iv) A statement that the resident may file a complaint with the State survey and certification agency concerning resident abuse, neglect, misappropriation of resident property in the facility, and non-compliance with the advance directives requirements.

(8) The facility must comply with the requirements specified in subpart I of part 489 of this chapter relating to maintaining written policies and procedures regarding advance directives. These requirements include provisions to inform and provide written information to all adult residents concerning the right to accept or refuse medical or surgical treatment and, at the individual's option, formulate an advance directive. This includes a written description of the facility's policies to implement advance directives and applicable State law. Facilities 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. If an adult individual is incapacitated at the time of admission 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 facility may give advance directive information to the individual's family or surrogate in the same manner that it issues other materials about policies and procedures to the family of the incapacitated individual or to a surrogate or other concerned persons in accordance with State law. The facility is not relieved of its obligation to provide this information to the individual once he or she is no longer incapacitated or unable to receive such information. Follow-up procedures must be in place to provide the information to the individual directly at the appropriate time.

(9) The facility must inform each resident of the name, specialty, and way of contacting the physician responsible for his or her care.

(10) The facility must prominently display in the facility written information, and provide to residents and applicants for admission oral and written information about how to apply for and
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use Medicare and Medicaid benefits, and how to receive refunds for previous payments covered by such benefits.

(11) Notification of changes. (i) A facility must immediately inform the resident; consult with the resident’s physician; and if known, notify the resident’s legal representative or an interested family member when there is—

(A) An accident involving the resident which results in injury and has the potential for requiring physician intervention;

(B) A significant change in the resident’s physical, mental, or psychosocial status (i.e., a deterioration in health, mental, or psychosocial status in either life-threatening conditions or clinical complications);

(C) A need to alter treatment significantly (i.e., a need to discontinue an existing form of treatment due to adverse consequences, or to commence a new form of treatment); or

(D) A decision to transfer or discharge the resident from the facility as specified in § 483.12(a).

(ii) The facility must also promptly notify the resident and, if known, the resident’s legal representative or interested family member when there is—

(A) A change in room or roommate assignment as specified in § 483.15(e)(2); or

(B) A change in resident rights under Federal or State law or regulations as specified in paragraph (b)(1) of this section.

(iii) The facility must record and periodically update the address and phone number of the resident’s legal representative or interested family member.

(c) Protection of resident funds. (1) The resident has the right to manage his or her financial affairs, and the facility may not require residents to deposit their personal funds with the facility.

(2) Management of personal funds. Upon written authorization of a resident, the facility must hold, safeguard, manage, and account for the personal funds of the resident deposited with the facility, as specified in paragraphs (c)(3)-(8) of this section.

(3) Deposit of funds. (i) Funds in excess of $50. The facility must deposit any residents’ personal funds in excess of $50 in an interest bearing account (or accounts) that is separate from any of the facility’s operating accounts, and that credits all interest earned on resident’s funds to that account. (In pooled accounts, there must be a separate accounting for each resident’s share.)

(ii) Funds less than $50. The facility must maintain a resident’s personal funds that do not exceed $50 in a non-interest bearing account, interest-bearing account, or petty cash fund.

(4) Accounting and records. The facility must establish and maintain a system that assures a full and complete and separate accounting, according to generally accepted accounting principles, of each resident’s personal funds entrusted to the facility on the resident’s behalf.

(i) The system must preclude any commingling of resident funds with facility funds or with the funds of any person other than another resident.

(ii) The individual financial record must be available through quarterly statements and on request to the resident or his or her legal representative.

(5) Notice of certain balances. The facility must notify each resident that receives Medicaid benefits—

(i) When the amount in the resident’s account reaches $200 less than the SSI resource limit for one person, specified in section 1611(a)(3)(B) of the Act; and

(ii) That, if the amount in the account, in addition to the value of the resident’s other nonexempt resources, reaches the SSI resource limit for one person, the resident may lose eligibility for Medicaid or SSI.

(6) Conveyance upon death. Upon the death of a resident with a personal fund deposited with the facility, the facility must convey within 30 days the resident’s funds, and a final accounting of those funds, to the individual or probate jurisdiction administering the resident’s estate.

(7) Assurance of financial security. The facility must purchase a surety bond, or otherwise provide assurance satisfactory to the Secretary, to assure the security of all personal funds of residents deposited with the facility.

(8) Limitation on charges to personal funds. The facility may not impose a charge against the personal funds of a resident for any item or service for which payment is made under Medicaid.
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or Medicare (except for applicable deductible and coinsurance amounts). The facility may charge the resident for requested services that are more expensive than or in excess of covered services in accordance with §489.32 of this chapter. (This does not affect the prohibition on facility charges for items and services for which Medicaid has paid. See §447.15, which limits participation in the Medicaid program to providers who accept, as payment in full, Medicaid payment plus any deductible, coinsurance, or copayment required by the plan to be paid by the individual.)

(i) Services included in Medicare or Medicaid payment. During the course of a covered Medicare or Medicaid stay, facilities may not charge a resident for the following categories of items and services:
(A) Nursing services as required at §483.30 of this subpart.
(B) Dietary services as required at §483.35 of this subpart.
(C) An activities program as required at §483.15(f) of this subpart.
(D) Room/bed maintenance services.

(ii) Items and services that may be charged to residents’ funds. Listed below are general categories and examples of items and services that the facility may charge to residents’ funds if they are requested by a resident, if the facility informs the resident that there will be a charge, and if payment is not made by Medicare or Medicaid:
(A) Telephone.
(B) Television/radio for personal use.
(C) Personal comfort items, including smoking materials, notions and novelties, and confections.
(D) Cosmetic and grooming items and services in excess of those for which payment is made under Medicaid or Medicare.
(E) Personal clothing.
(F) Personal reading matter.
(G) Gifts purchased on behalf of a resident.
(H) Flowers and plants.
(I) Social events and entertainment offered outside the scope of the activities program, provided under §483.15(f) of this subpart.
(J) Noncovered special care services such as privately hired nurses or aides.
(K) Private room, except when therapeutically required (for example, isolation for infection control).
(L) Specially prepared or alternative food requested instead of the food generally prepared by the facility, as required by §483.35 of this subpart.

(iii) Requests for items and services. (A) The facility must not charge a resident (or his or her representative) for any item or service not requested by the resident.
(B) The facility must not require a resident (or his or her representative) to request any item or service as a condition of admission or continued stay.
(C) The facility must inform the resident (or his or her representative) requesting an item or service for which a charge will be made that there will be a charge for the item or service and what the charge will be.

(d) Free choice. The resident has the right to—
(1) Choose a personal attending physician;
(2) Be fully informed in advance about care and treatment and of any changes in that care or treatment that may affect the resident’s well-being; and
(3) Unless adjudged incompetent or otherwise found to be incapacitated under the laws of the State, participate in planning care and treatment or changes in care and treatment.

(e) Privacy and confidentiality. The resident has the right to personal privacy and confidentiality of his or her personal and clinical records.
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(1) Personal privacy includes accommodations, medical treatment, written and telephone communications, personal care, visits, and meetings of family and resident groups, but this does not require the facility to provide a private room for each resident;

(2) Except as provided in paragraph (e)(3) of this section, the resident may approve or refuse the release of personal and clinical records to any individual outside the facility;

(3) The resident's right to refuse release of personal and clinical records does not apply when—
   (i) The resident is transferred to another health care institution; or
   (ii) Record release is required by law.

(f) Grievances. A resident has the right to—

(1) Voice grievances without discrimination or reprisal. Such grievances include those with respect to treatment which has been furnished as well as that which has not been furnished; and

(2) Prompt efforts by the facility to resolve grievances the resident may have, including those with respect to the behavior of other residents.

(g) Examination of survey results. A resident has the right to—

(1) Examine the results of the most recent survey of the facility conducted by Federal or State surveyors and any plan of correction in effect with respect to the facility. The facility must make the results available for examination in a place readily accessible to residents, and must post a notice of their availability; and

(2) Receive information from agencies acting as client advocates, and be afforded the opportunity to contact these agencies.

(h) Work. The resident has the right to—

(1) Refuse to perform services for the facility;

(2) Perform services for the facility, if he or she chooses, when—
   (i) The facility has documented the need or desire for work in the plan of care;
   (ii) The plan specifies the nature of the services performed and whether the services are voluntary or paid;
   (iii) Compensation for paid services is at or above prevailing rates; and

(iv) The resident agrees to the work arrangement described in the plan of care.

(i) Mail. The resident has the right to privacy in written communications, including the right to—

(1) Send and promptly receive mail that is unopened; and

(2) Have access to stationery, postage, and writing implements at the resident's own expense.

(j) Access and visitation rights. (1) The resident has the right and the facility must provide immediate access to any resident by the following:

   (i) Any representative of the Secretary;
   (ii) Any representative of the State;
   (iii) The resident's individual physician;
   (iv) The State long term care ombudsman (established under section 307(a)(12) of the Older Americans Act of 1965);
   (v) The agency responsible for the protection and advocacy system for developmentally disabled individuals (established under part C of the Developmental Disabilities Assistance and Bill of Rights Act);
   (vi) The agency responsible for the protection and advocacy system for mentally ill individuals (established under the Protection and Advocacy for Mentally Ill Individuals Act);
   (vii) Subject to the resident's right to deny or withdraw consent at any time, immediate family or other relatives of the resident; and
   (viii) Subject to reasonable restrictions and the resident's right to deny or withdraw consent at any time, others who are visiting with the consent of the resident.

(2) The facility must provide reasonable access to any resident by any entity or individual that provides health, social, legal, or other services to the resident, subject to the resident's right to deny or withdraw consent at any time.

(3) The facility must allow representatives of the State Ombudsman, described in paragraph (j)(1)(iv) of this section, to examine a resident's clinical records with the permission of the resident or the resident's legal representative, and consistent with State law.
§ 483.12 Admission, transfer and discharge rights.

(a) Transfer and discharge—

(1) Definition: Transfer and discharge includes movement of a resident to a bed outside of the certified facility whether that bed is in the same physical plant or not. Transfer and discharge does not refer to movement of a resident to a bed within the same certified facility.

(2) Transfer and discharge requirements. The facility must permit each resident to remain in the facility, and not transfer or discharge the resident from the facility unless—

(i) The transfer or discharge is necessary for the resident's welfare and the resident's needs cannot be met in the facility;

(ii) The transfer or discharge is appropriate because the resident's health has improved sufficiently so the resident no longer needs the services provided by the facility;

(iii) The safety of individuals in the facility is endangered;

(iv) The health of individuals in the facility would otherwise be endangered;

(v) The resident has failed, after reasonable and appropriate notice, to pay for (or to have paid under Medicare or Medicaid) a stay at the facility. For a resident who becomes eligible for Medicaid after admission to a facility, the facility may charge a resident only allowable charges under Medicaid; or

(vi) The facility ceases to operate.

(3) Documentation. When the facility transfers or discharges a resident under any of the circumstances specified in paragraphs (a)(2)(i) through (v) of this section, the resident's clinical record must be documented. The documentation must be made by—

(i) The resident's physician when transfer or discharge is necessary under paragraph (a)(2)(i) or paragraph (a)(2)(ii) of this section; and

(ii) A physician when transfer or discharge is necessary under paragraph (a)(2)(iv) of this section.

(4) Notice before transfer. Before a facility transfers or discharges a resident, the facility must—

(i) Notify the resident and, if known, a family member or legal representative of the resident of the transfer or discharge and the reasons for the move in writing and in a language and manner they understand.

(ii) Record the reasons in the resident's clinical record; and

(iii) Include in the notice the items described in paragraph (a)(6) of this section.

(5) Timing of the notice. (i) Except when specified in paragraph (a)(5)(ii) of this section, the notice of transfer or discharge required under paragraph (a)(4) of this section must be made by the facility at least 30 days before the resident is transferred or discharged.
(ii) Notice may be made as soon as practicable before transfer or discharge when—
(A) the safety of individuals in the facility would be endangered under paragraph (a)(2)(iii) of this section;
(B) The health of individuals in the facility would be endangered, under paragraph (a)(2)(iv) of this section;
(C) The resident’s health improves sufficiently to allow a more immediate transfer or discharge, under paragraph (a)(2)(ii) of this section;
(D) An immediate transfer or discharge is required by the resident’s urgent medical needs, under paragraph (a)(2)(i) of this section; or
(E) A resident has not resided in the facility for 30 days.

(6) Contents of the notice. The written notice specified in paragraph (a)(4) of this section must include the following:
(i) The reason for transfer or discharge;
(ii) The effective date of transfer or discharge;
(iii) The location to which the resident is transferred or discharged;
(iv) A statement that the resident has the right to appeal the action to the State;
(v) The name, address and telephone number of the State long term care ombudsman;
(vi) For nursing facility residents with developmental disabilities, the mailing address and telephone number of the agency responsible for the protection and advocacy of developmentally disabled individuals established under Part C of the Developmental Disabilities Assistance and Bill of Rights Act; and
(vii) For nursing facility residents who are mentally ill, the mailing address and telephone number of the agency responsible for the protection and advocacy of mentally ill individuals established under the Protection and Advocacy for Mentally Ill Individuals Act.

(7) Orientation for transfer or discharge. A facility must provide sufficient preparation and orientation to residents to ensure safe and orderly transfer or discharge from the facility.

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(a) Transfer or discharge of a resident.

(b) Notice of bed-hold policy and readmission—

1) Notice before transfer. Before a nursing facility transfers a resident to a hospital or allows a resident to go on therapeutic leave, the nursing facility must provide written information to the resident and a family member or legal representative that specifies—
(i) The duration of the bed-hold policy under the State plan, if any, during which the resident is permitted to return and resume residence in the nursing facility; and
(ii) The nursing facility’s policies regarding bed-hold periods, which must be consistent with paragraph (b)(3) of this section, permitting a resident to return.

2) Bed-hold notice upon transfer. At the time of transfer of a resident for hospitalization or therapeutic leave, a nursing facility must provide to the resident and a family member or legal representative written notice which specifies the duration of the bed-hold policy described in paragraph (b)(1) of this section.

3) Permits resident to return to facility. A nursing facility must establish and follow a written policy under which a resident, whose hospitalization or therapeutic leave exceeds the bed-hold period under the State plan, is readmitted to the facility immediately upon the first availability of a bed in a semi-private room if the resident—
(i) Requires the services provided by the facility; and
(ii) Is eligible for Medicaid nursing facility services.

(c) Equal access to quality care.

1) A facility must establish and maintain identical policies and practices regarding transfer, discharge, and the provision of services under the State plan for all individuals regardless of source of payment; and
2) The facility may charge any amount for services furnished to non-Medicaid residents consistent with the notice requirement in §483.10(b)(5)(i) and (b)(6) describing the charges; and
3) The State is not required to offer additional services on behalf of a resident other than services provided in the State plan.

(d) Admissions policy.

1) The facility must—
(i) Not require residents or potential residents to waive their rights to Medicare or Medicaid; and
(ii) Not require oral or written assurance that residents or potential residents are not eligible for, or will not apply for, Medicare or Medicaid benefits.

(2) The facility must not require a third party guarantee of payment to the facility as a condition of admission or expedited admission, or continued stay in the facility. However, the facility may require an individual who has legal access to a resident’s income or resources available to pay for facility care to sign a contract, without incurring personal financial liability, to provide facility payment from the resident’s income or resources.

(3) In the case of a person eligible for Medicaid, a nursing facility must not charge, solicit, accept, or receive, in addition to any amount otherwise required to be paid under the State plan, any gift, money, donation, or other consideration as a precondition of admission, expedited admission, or continued stay in the facility. However,—

(i) A nursing facility may charge a resident who is eligible for Medicaid for items and services the resident has requested and received, in addition to any amount otherwise required to be paid under the State plan, any gift, money, donation, or other consideration as a precondition of admission, expedited admission, or continued stay in the facility. However,—

(ii) A nursing facility may solicit, accept, or receive a charitable, religious, or philanthropic contribution from an organization or from a person unrelated to a Medicaid eligible resident or potential resident, but only to the extent that the contribution is not a condition of admission, expedited admission, or continued stay in the facility for a Medicaid eligible resident.

(4) States or political subdivisions may apply stricter admissions standards under State or local laws than are specified in this section, to prohibit discrimination against individuals entitled to Medicaid.

§ 483.13 Resident behavior and facility practices.

(a) Restraints. The resident has the right to be free from any physical or chemical restraints imposed for purposes of discipline or convenience, and not required to treat the resident’s medical symptoms.

(b) Abuse. The resident has the right to be free from verbal, sexual, physical, and mental abuse, corporal punishment, and involuntary seclusion.

(c) Staff treatment of residents. The facility must develop and implement written policies and procedures that prohibit mistreatment, neglect, and abuse of residents and misappropriation of resident property.

(1) The facility must—

(i) Not use verbal, mental, sexual, or physical abuse, corporal punishment, or involuntary seclusion;

(ii) Not employ individuals who have been—

(A) Found guilty of abusing, neglecting, or mistreating residents by a court of law; or

(B) Have had a finding entered into the State nurse aide registry concerning abuse, neglect, mistreatment of residents, or misappropriation of their property; and

(iii) Report any knowledge it has of actions by a court of law against an employee, which would indicate unfitness for service as a nurse aide or other facility staff to the State nurse aide registry or licensing authorities.

(2) The facility must ensure that all alleged violations involving mistreatment, neglect, or abuse, including injuries of unknown source, and misappropriation of resident property are reported immediately to the administrator of the facility and to other officials in accordance with State law through established procedures (including to the State survey and certification agency).

(3) The facility must have evidence that all alleged violations are thoroughly investigated, and must prevent further potential abuse while the investigation is in progress.

(4) The results of all investigations must be reported to the administrator or his designated representative and to other officials in accordance with
§ 483.15 Quality of life.

A facility must care for its residents in a manner and in an environment that promotes maintenance or enhancement of each resident's quality of life.

(a) Dignity. The facility must promote care for residents in a manner and in an environment that maintains or enhances each resident's dignity and respect in full recognition of his or her individuality.

(b) Self-determination and participation. The resident has the right to—

(1) Choose activities, schedules, and health care consistent with his or her interests, assessments, and plans of care;

(2) Interact with members of the community both inside and outside the facility; and

(3) Make choices about aspects of his or her life in the facility that are significant to the resident.

(c) Participation in resident and family groups. (1) A resident has the right to organize and participate in resident groups in the facility;

(2) A resident's family has the right to meet in the facility with the families of other residents in the facility;

(3) The facility must provide a resident or family group, if one exists, with private space;

(4) Staff or visitors may attend meetings at the group's invitation;

(5) The facility must provide a designated staff person responsible for providing assistance and responding to written requests that result from group meetings;

(6) When a resident or family group exists, the facility must listen to the views and act upon the grievances and recommendations of residents and families concerning proposed policy and operational decisions affecting resident care and life in the facility.

(d) Participation in other activities. The resident has the right to participate in social, religious, and community activities that do not interfere with the rights of other residents in the facility.

(e) Accommodation of needs. A resident has the right to—

(1) Reside and receive services in the facility with reasonable accommodation of individual needs and preferences, except when the health or safety of the individual or other residents would be endangered; and

(2) Receive notice before the resident's room or roommate in the facility is changed.

(f) Activities. (1) The facility must provide for an ongoing program of activities designed to meet, in accordance with the comprehensive assessment, the interests and the physical, mental, and psychosocial well-being of each resident.

(2) The activities program must be directed by a qualified professional who—

(i) Is a qualified therapeutic recreation specialist or an activities professional who—

(A) Is licensed or registered, if applicable, by the State in which practicing; and

(B) Is eligible for certification as a therapeutic recreation specialist or as an activities professional by a recognized accrediting body on or after October 1, 1990; or

(ii) Has 2 years of experience in a social or recreational program within the last 5 years, 1 of which was full-time in a patient activities program in a health care setting; or

(iii) Is a qualified occupational therapist or occupational therapy assistant; or

(iv) Has completed a training course approved by the State.

(g) Social Services. (1) The facility must provide medically-related social services to attain or maintain the highest practicable physical, mental, and psychosocial well-being of each resident.

(2) A facility with more than 120 beds must employ a qualified social worker on a full-time basis.

(3) Qualifications of social worker. A qualified social worker is an individual who—

(i) Has a bachelor's degree in social work or a bachelor's degree in a human services field including but not limited to
§ 483.20 Resident assessment.

The facility must conduct initially and periodically a comprehensive, accurate, standardized, reproducible assessment of each resident's functional capacity.

(a) Admission orders. At the time each resident is admitted, the facility must have physician orders for the resident's immediate care.

(b) Comprehensive assessments.

(1) Resident assessment instrument. A facility must make a comprehensive assessment of a resident's needs, using the resident assessment instrument (RAI) specified by the State. The assessment must include at least the following:

(i) Identification and demographic information.

(ii) Customary routine.

(iii) Cognitive patterns.

(iv) Communication.

(v) Vision.

(vi) Mood and behavior patterns.

(vii) Psychosocial well-being.

(viii) Physical functioning and structural problems.

(ix) Continence.

(x) Disease diagnoses and health conditions.

(xi) Dental and nutritional status.

(xii) Skin condition.

(xiii) Activity pursuit.

(xiv) Medications.

(xv) Special treatments and procedures.

(xvi) Discharge potential.

(xvii) Documentation of summary information regarding the additional assessment performed through the resident assessment protocols.

(xviii) Documentation of participation in assessment.

The assessment process must include direct observation and communication with the resident, as well as communication with licensed and nonlicensed direct care staff members on all shifts.

(2) When required. Subject to the timeframes prescribed in §413.343(b) of this chapter, a facility must conduct a comprehensive assessment of a resident in accordance with the timeframes specified in paragraphs (b)(2)(i) through (iii) of this section. The timeframes prescribed in §413.343(b) of this chapter do not apply to CAHs.

(i) Within 14 calendar days after admission, excluding readmissions in which there is no significant change in the resident's physical or mental condition. (For purposes of this section, “readmission” means a return to the facility following a temporary absence for hospitalization or for therapeutic leave.)

(ii) Within 14 calendar days after the facility determines, or should have determined, that there has been a significant change in the resident's physical or mental condition. (For purposes of this section, a “significant change” means a major decline or improvement in the resident's status that will not normally resolve itself without further intervention by staff or by implementing standard disease-related clinical interventions, that has an impact on more than one area of the resident's health status, and requires interdisciplinary review or revision of the care plan, or both.)

(iii) Not less often than once every 12 months.

(c) Quarterly review assessment. A facility must assess a resident using the quarterly review instrument specified in §413.343(c) of this chapter.

The quarterly review instrument includes the following:

(i) Current health status.

(ii) Status of any additional assessment performed during the previous 12 months.

(iii) Discharge potential.

(iv) Documentation of summary information regarding the additional assessment performed through the resident assessment protocols, the care plan, or both.

(v) Documentation of participation in assessment.

The quarterly review process must include direct observation and communication with the resident, as well as communication with licensed and nonlicensed direct care staff members on all shifts.

(3) After discharge. A facility must conduct a comprehensive assessment of a resident prior to discharge.

(i) Upon discharge, or upon return from any leave (whether formal or informal).

(ii) As required by the resident.

(iii) Not less often than once every 12 months.

(4) Discharge summary. The facility must prepare a discharge summary that includes the following:

(i) Identification and demographic information.

(ii) Status of any additional assessment performed during the previous 12 months.

(iii) Care plan.

(iv) Discharge potential.

(v) Documentation of summary information regarding the additional assessment performed through the resident assessment protocols, the care plan, or both.

(vi) Documentation of participation in assessment.

(5) Discharge planning. The facility must conduct an initial discharge planning assessment.

(i) At the time each resident is determined to be eligible for discharge.

(ii) At the time the facility determines, or should have determined, that there has been a significant change in the resident's status.

(iii) At least once every 12 months.

(6) Discharge planning reviews. The facility must conduct discharge planning reviews.

(i) Within 14 calendar days after discharge planning.

(ii) At least once every 12 months.

(iii) Not less often than once every 12 months.

(7) Changes in resident status. A facility must conduct a comprehensive assessment when there is a significant change in the resident's status.

(i) Upon discharge.

(ii) Upon return from any leave (whether formal or informal).

(iii) As required by the resident.

(iv) Not less often than once every 12 months.

(8) Best interests. The facility must conduct a comprehensive assessment when the resident is not able to make an informed decision.

(i) Upon admission.

(ii) When the resident is not able to make an informed decision.

(iii) As required by the resident.

(iv) Not less often than once every 12 months.

(9) Interdisciplinary review. The facility must conduct an interdisciplinary review of the resident's care plan when there is a significant change in the resident's status.

(i) Upon discharge.

(ii) Upon return from any leave (whether formal or informal).

(iii) As required by the resident.

(iv) Not less often than once every 12 months.

(10) Family conferences. The facility must conduct family conferences when there is a significant change in the resident's status.

(i) Upon discharge.

(ii) Upon return from any leave (whether formal or informal).

(iii) As required by the resident.

(iv) Not less often than once every 12 months.

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sociology, special education, rehabilitation counseling, and psychology; and

(ii) One year of supervised social work experience in a health care setting working directly with individuals.

(h) Environment. The facility must provide—

(1) A safe, clean, comfortable, and homelike environment, allowing the resident to use his or her personal belongings to the extent possible;

(2) Housekeeping and maintenance services necessary to maintain a sanitary, orderly, and comfortable interior;

(3) Clean bed and bath linens that are in good condition;

(4) Private closet space in each resident room, as specified in §483.70(d)(2)(iv) of this part;

(5) Adequate and comfortable lighting levels in all areas;

(6) Comfortable and safe temperature levels. Facilities initially certified after October 1, 1990 must maintain a temperature range of 71±81° F; and

(7) For the maintenance of comfortable sound levels.

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by the State and approved by HCFA not less frequently than once every 3 months.

(d) Use. A facility must maintain all resident assessments completed within the previous 15 months in the resident’s active record and use the results of the assessments to develop, review, and revise the resident’s comprehensive plan of care.

(e) Coordination. A facility must coordinate assessments with the preadmission screening and resident review program under Medicaid in part 483, subpart C to the maximum extent practicable to avoid duplicative testing and effort.

(f) Automated data processing requirement. (1) Encoding data. Within 7 days after a facility completes a resident’s assessment, a facility must encode the following information for each resident in the facility:

(i) Admission assessment.
(ii) Annual assessment updates.
(iii) Significant change in status assessments.
(iv) Quarterly review assessments.
(v) A subset of items upon a resident’s transfer, reentry, discharge, and death.
(vi) Background (face-sheet) information, if there is no admission assessment.

(2) Transmitting data. Within 7 days after a facility completes a resident’s assessment, a facility must be capable of transmitting to the State information for each resident contained in the MDS in a format that conforms to standard record layouts and data dictionaries, and that passes standardized edits defined by HCFA and the State.

(3) Monthly transmittal requirements. A facility must electronically transmit, at least monthly, encoded, accurate, complete MDS data to the State for all assessments conducted during the previous month, including the following:

(i) Admission assessment.
(ii) Annual assessment.
(iii) Significant change in status assessment.
(iv) Significant correction of prior full assessment.
(v) Significant correction of prior quarterly assessment.
(vi) Quarterly review.

(vii) A subset of items upon a resident’s transfer, reentry, discharge, and death.

(viii) Background (face-sheet) information, for an initial transmission of MDS data on a resident that does not have an admission assessment.

(4) Data format. The facility must transmit data in the format specified by HCFA or, for a State which has an alternate RAI approved by HCFA, in the format specified by the State and approved by HCFA.

(5) Resident-identifiable information. (i) A facility may not release information that is resident-identifiable to the public.

(ii) The facility may release information that is resident-identifiable to an agent only in accordance with a contract under which the agent agrees not to use or disclose the information except to the extent the facility itself is permitted to do so.

(g) Accuracy of assessments. The assessment must accurately reflect the resident’s status.

(h) Coordination. A registered nurse must conduct or coordinate each assessment with the appropriate participation of health professionals.

(i) Certification. (1) A registered nurse must sign and certify that the assessment is completed.

(2) Each individual who completes a portion of the assessment must sign and certify the accuracy of that portion of the assessment.

(j) Penalty for falsification. (1) Under Medicare and Medicaid, an individual who willfully and knowingly—

(i) Certifies a material and false statement in a resident assessment is subject to a civil money penalty of not more than $1,000 for each assessment; or

(ii) Causes another individual to certify a material and false statement in a resident assessment is subject to a civil money penalty of not more than $5,000 for each assessment.

(2) Clinical disagreement does not constitute a material and false statement.

(k) Comprehensive care plans. (1) The facility must develop a comprehensive care plan for each resident that includes measurable objectives and timetables to meet a resident’s medical,
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nursing, and mental and psychosocial needs that are identified in the comprehensive assessment. The care plan must describe the following—

(i) The services that are to be furnished to attain or maintain the resident’s highest practicable physical, mental, and psychosocial well-being as required under § 483.25; and

(ii) Any services that would otherwise be required under § 483.25 but are not provided due to the resident’s exercise of rights under § 483.10, including the right to refuse treatment under § 483.10(b)(4).

(2) A comprehensive care plan must be—

(i) Developed within 7 days after completion of the comprehensive assessment;

(ii) Prepared by an interdisciplinary team, that includes the attending physician, a registered nurse with responsibility for the resident, and other appropriate staff in disciplines as determined by the resident’s needs, and, to the extent practicable, the participation of the resident, the resident’s family or the resident’s legal representative; and

(iii) Periodically reviewed and revised by a team of qualified persons after each assessment.

(3) The services provided or arranged by the facility must—

(i) Meet professional standards of quality; and

(ii) Be provided by qualified persons in accordance with each resident’s written plan of care.

(1) Discharge summary. When the facility anticipates discharge a resident must have a discharge summary that includes—

A recapitulation of the resident’s stay;

A final summary of the resident’s status to include items in paragraph (b)(2) of this section, at the time of the discharge that is available for release to authorized persons and agencies, with the consent of the resident or legal representative; and

A post-discharge plan of care that is developed with the participation of the resident and his or her family, which will assist the resident to adjust to his or her new living environment.

(m) Preadmission screening for mentally ill individuals and individuals with mental retardation. (1) A nursing facility must not admit, on or after January 1, 1989, any new resident with—

(i) Mental illness as defined in paragraph (f)(2)(i) of this section, unless the State mental health authority has determined, based on an independent physical and mental evaluation performed by a person or entity other than the State mental health authority, prior to admission,

(A) That, because of the physical and mental condition of the individual, the individual requires the level of services provided by a nursing facility; and

(B) If the individual requires such level of services, whether the individual requires specialized services; or

(ii) Mental retardation, as defined in paragraph (f)(2)(ii) of this section, unless the State mental retardation or developmental disability authority has determined prior to admission—

(A) That, because of the physical and mental condition of the individual, the individual requires the level of services provided by a nursing facility; and

(B) If the individual requires such level of services, whether the individual requires specialized services for mental retardation.

(2) Definition. For purposes of this section—

(i) An individual is considered to have mental illness if the individual has a serious mental illness as defined in § 483.102(b)(1).

(ii) An individual is considered to be mentally retarded if the individual is mentally retarded as defined in § 483.102(b)(3) or is a person with a related condition as described in 42 CFR 435.1009.

(a) Activities of daily living. Based on the comprehensive assessment of a resident, the facility must ensure that—

(1) A resident's abilities in activities of daily living do not diminish unless circumstances of the individual's clinical condition demonstrate that diminution was unavoidable. This includes the resident's ability to—

(i) Bathe, dress, and groom;
(ii) Transfer and ambulate;
(iii) Toilet;
(iv) Eat; and
(v) Use speech, language, or other functional communication systems.

(2) A resident is given the appropriate treatment and services to maintain or improve his or her abilities specified in paragraph (a)(1) of this section; and

(3) A resident who is unable to carry out activities of daily living receives the necessary services to maintain good nutrition, grooming, and personal and oral hygiene.

(b) Vision and hearing. To ensure that residents receive proper treatment and assistive devices to maintain vision and hearing abilities, the facility must, if necessary, assist the resident—

(1) In making appointments, and

(2) By arranging for transportation to and from the office of a practitioner specializing in the treatment of vision or hearing impairment or the office of a professional specializing in the provision of vision or hearing assistive devices.

(c) Pressure sores. Based on the comprehensive assessment of a resident, the facility must ensure that—

(1) A resident who enters the facility without pressure sores does not develop pressure sores unless the individual's clinical condition demonstrates that they were unavoidable; and

(2) A resident having pressure sores receives necessary treatment and services to promote healing, prevent infection and prevent new sores from developing.

(d) Urinary Incontinence. Based on the resident's comprehensive assessment, the facility must ensure that—

(1) A resident who enters the facility without an indwelling catheter is not catheterized unless the resident's clinical condition demonstrates that catheterization was necessary; and

(2) A resident who is incontinent of bladder receives appropriate treatment and services to prevent urinary tract infections and to restore as much normal bladder function as possible.

(e) Range of motion. Based on the comprehensive assessment of a resident, the facility must ensure that—

(1) A resident who enters the facility without a limited range of motion does not experience reduction in range of motion unless the resident's clinical condition demonstrates that a reduction in range of motion is unavoidable; and

(2) A resident with a limited range of motion receives appropriate treatment and services to increase range of motion and/or to prevent further decrease in range of motion.

(f) Mental and Psychosocial functioning. Based on the comprehensive assessment of a resident, the facility must ensure that—

(1) A resident who displays mental or psychosocial adjustment difficulty, receives appropriate treatment and services to correct the assessed problem, and

(2) A resident whose assessment did not reveal a mental or psychosocial adjustment difficulty does not display a pattern of decreased social interaction and/or increased withdrawn, angry, or depressive behaviors, unless the resident's clinical condition demonstrates that such a pattern was unavoidable.

(g) Naso-gastric tubes. Based on the comprehensive assessment of a resident, the facility must ensure that—

(1) A resident who has been able to eat enough alone or with assistance is not fed by naso-gastric tube unless the resident's clinical condition demonstrates that use of a naso-gastric tube was unavoidable; and

(2) A resident who is fed by a naso-gastric or gastrostomy tube receives the appropriate treatment and services to prevent aspiration pneumonia, diarrhea, vomiting, dehydration, metabolic abnormalities, and nasal-pharyngeal ulcers and to restore, if possible, normal eating skills.

(h) Accidents. The facility must ensure that—
(1) The resident environment remains as free of accident hazards as is possible; and
(2) Each resident receives adequate supervision and assistance devices to prevent accidents.

(i) Nutrition. Based on a resident’s comprehensive assessment, the facility must ensure that a resident—
(1) Maintains acceptable parameters of nutritional status, such as body weight and protein levels, unless the resident’s clinical condition demonstrates that this is not possible; and
(2) Receives a therapeutic diet when there is a nutritional problem.

(j) Hydration. The facility must provide each resident with sufficient fluid intake to maintain proper hydration and health.

(k) Special needs. The facility must ensure that residents receive proper treatment and care for the following special services:
(1) Injections;
(2) Parenteral and enteral fluids;
(3) Colostomy, ureterostomy, or ileostomy care;
(4) Tracheostomy care;
(5) Tracheal suctioning;
(6) Respiratory care;
(7) Foot care; and
(8) Prostheses.

(l) Unnecessary drugs—(1) General. Each resident’s drug regimen must be free from unnecessary drugs. An unnecessary drug is any drug when used:
(i) In excessive dose (including duplicate drug therapy); or
(ii) For excessive duration; or
(iii) Without adequate monitoring; or
(iv) Without adequate indications for its use; or
(v) In the presence of adverse consequences which indicate the dose should be reduced or discontinued; or
(vi) Any combinations of the reasons above.

(2) Antipsychotic Drugs. Based on a comprehensive assessment of a resident, the facility must ensure that—
(i) Residents who have not used antipsychotic drugs are not given these drugs unless antipsychotic drug therapy is necessary to treat a specific condition as diagnosed and documented in the clinical record; and
(ii) Residents who use antipsychotic drugs receive gradual dose reductions, and behavioral interventions, unless clinically contraindicated, in an effort to discontinue these drugs.

(m) Medication Errors. The facility must ensure that—
(1) It is free of medication error rates of five percent or greater; and
(2) Residents are free of any significant medication errors.

§ 483.30 Nursing services.

The facility must have sufficient nursing staff to provide nursing and related services to attain or maintain the highest practicable physical, mental, and psychosocial well-being of each resident, as determined by resident assessments and individual plans of care.

(a) Sufficient staff. (1) The facility must provide services by sufficient numbers of each of the following types of personnel on a 24-hour basis to provide nursing care to all residents in accordance with resident care plans:
(i) Except when waived under paragraph (c) of this section, licensed nurses; and
(ii) Other nursing personnel.

(2) Except when waived under paragraph (c) or (d) of this section, the facility must designate a registered nurse to serve as a charge nurse on each tour of duty.

(b) Registered nurse. (1) Except when waived under paragraph (c) of this section, the facility must designate a licensed nurse to serve as a charge nurse on each tour of duty.

(2) Except when waived under paragraph (c) or (d) of this section, the facility must designate a registered nurse to serve as the director of nursing on a full time basis.

(3) The director of nursing may serve as a charge nurse only when the facility has an average daily occupancy of 60 or fewer residents.

(c) Nursing facilities: Waiver of requirement to provide licensed nurses. To the extent that a facility is unable to meet the requirements of paragraphs (a)(2) and (b)(1) of this section, a State may waive such requirements with respect to the facility if—
§ 483.35 Dietary services.

The facility must provide each resident with a nourishing, palatable, well-balanced diet that meets the daily nutritional and special dietary needs of each resident.

(a) Staffing. The facility must employ a qualified dietitian either full-time, part-time, or on a consultant basis.

(1) If a qualified dietitian is not employed full-time, the facility must designate a person to serve as the director of food service who receives frequently scheduled consultation from a qualified dietitian.

(2) A qualified dietitian is one who is qualified based upon either registration by the Commission on Dietetic Registration of the American Dietetic Association, or on the basis of education, training, or experience in identification of dietary needs, planning, and implementation of dietary programs.

(b) Sufficient staff. The facility must employ sufficient support personnel to—

(A) Has only patients whose physicians have indicated (through physicians’ orders or admission notes) that they do not require the services of a registered nurse or a physician for a 48-hours period, or

(B) Has made arrangements for a registered nurse or a physician to spend time at the facility, as determined necessary by the physician, to provide necessary skilled nursing services on days when the regular full-time registered nurse is not on duty;

(iv) The Secretary provides notice of the waiver to the State long term care ombudsman (established under section 307(a)(12) of the Older Americans Act of 1965) and the protection and advocacy system in the State for the mentally ill and mentally retarded; and

(v) The facility that is granted such a waiver notifies residents of the facility (or, where appropriate, the guardians or legal representatives of such residents) and members of their immediate families of the waiver.

(2) A waiver of the registered nurse requirement under paragraph (d)(1) of this section is subject to annual renewal by the Secretary.
§ 483.40 Physician services.

A physician must personally approve in writing a recommendation that an individual be admitted to a facility. Each resident must remain under the care of a physician.

(a) Physician supervision. The facility must ensure that—

(1) The medical care of each resident is supervised by a physician; and

(2) Another physician supervises the medical care of residents when their attending physician is unavailable.

(b) Physician visits. The physician must—

(1) Review the resident’s total program of care, including medications and treatments, at each visit required by paragraph (c) of this section;

(2) Write, sign, and date progress notes at each visit; and

(3) Sign and date all orders.

(c) Frequency of physician visits.

(1) The resident must be seen by a physician at least once every 30 days for the first 90 days after admission, and at least once every 60 days thereafter.

(2) A physician visit is considered timely if it occurs not later than 10 days after the date the visit was required.

(3) Except as provided in paragraphs (c)(4) and (f) of this section, all required physician visits must be made by the physician personally.

(4) At the option of the physician, required visits in SNFs after the initial visit may alternate between personal visits by the physician and visits by a physician assistant, nurse practitioner, or clinical nurse specialist in accordance with paragraph (e) of this section.

(d) Availability of physicians for emergency care. The facility must provide or arrange for the provision of physician services 24 hours a day, in case of an emergency.

(e) Physician delegation of tasks in SNFs.

(1) Except as specified in paragraph (e)(2) of this section, a physician may delegate tasks to a physician assistant, nurse practitioner, or clinical nurse specialist who—

(i) Meets the applicable definition in §491.2 of this chapter or, in the case of a clinical nurse specialist, is licensed as such by the State;

(2) Has at least one hour of direct supervision by a physician or a provider who is employed by the facility.

(f) Access to medical records. The facility must provide access to a resident’s medical record to the following individuals—

(1) The resident;

(2) Any member of the resident’s family; and

(3) Any health care provider who is involved in the resident’s care.
§ 483.45 Specialized rehabilitative services.

(a) Provision of services. If specialized rehabilitative services such as but not limited to physical therapy, speech-language pathology, occupational therapy, and mental health rehabilitative services for mental illness and mental retardation, are required in the resident’s comprehensive plan of care, the facility must—

(1) Provide the required services; or
(2) Obtain the required services from an outside resource (in accordance with §483.75(h) of this part) from a provider of specialized rehabilitative services.

(b) Qualifications. Specialized rehabilitative services must be provided under the written order of a physician by qualified personnel.


§ 483.55 Dental services.

The facility must assist residents in obtaining routine and 24-hour emergency dental care.

(a) Skilled nursing facilities. A facility (1) Must provide or obtain from an outside resource, in accordance with §483.75(h) of this part, routine and emergency dental services to meet the needs of each resident;

(ii) May charge a Medicare resident an additional amount for routine and emergency dental services;

(iii) Must if necessary, assist the resident—

(i) In making appointments; and

(ii) By arranging for transportation to and from the dentist’s office; and

(4) Promptly refer residents with lost or damaged dentures to a dentist.

(b) Nursing facilities. The facility (1) Must provide or obtain from an outside resource, in accordance with §483.75(h) of this part, the following dental services to meet the needs of each resident:

(i) Routine dental services (to the extent covered under the State plan); and
(ii) Emergency dental services;

(2) Must, if necessary, assist the resident—

(i) In making appointments; and

(ii) By arranging for transportation to and from the dentist’s office; and

(3) Must promptly refer residents with lost or damaged dentures to a dentist.

[56 FR 48875, Sept. 26, 1991]

§ 483.60 Pharmacy services.

The facility must provide routine and emergency drugs and biologicals to its residents, or obtain them under an agreement described in §483.75(h) of this part. The facility may permit unlicensed personnel to administer drugs if State law permits, but only under the general supervision of a licensed nurse.

(a) Procedures. A facility must provide pharmaceutical services (including procedures that assure the accurate acquiring, receiving, dispensing, and administering of all drugs and biologicals) to meet the needs of each resident.

(b) Service consultation. The facility must employ or obtain the services of a licensed pharmacist who—

(1) Provides consultation on all aspects of the provision of pharmacy services in the facility;

(2) Establishes a system of records of receipt and disposition of all controlled drugs in sufficient detail to enable an accurate reconciliation; and

(3) Determines that drug records are in order and that an account of all controlled drugs is maintained and periodically reconciled.
(c) Drug regimen review. (1) The drug regimen of each resident must be reviewed at least once a month by a licensed pharmacist.

(2) The pharmacist must report any irregularities to the attending physician and the director of nursing, and these reports must be acted upon.

(d) Labeling of drugs and biologicals. Drugs and biologicals used in the facility must be labeled in accordance with currently accepted professional principles, and include the appropriate accessory and cautionary instructions, and the expiration date when applicable.

(e) Storage of drugs and biologicals. (1) In accordance with State and Federal laws, the facility must store all drugs and biologicals in locked compartments under proper temperature controls, and permit only authorized personnel to have access to the keys.

(2) The facility must provide separately locked, permanently affixed compartments for storage of controlled drugs listed in Schedule II of the Comprehensive Drug Abuse Prevention and Control Act of 1976 and other drugs subject to abuse, except when the facility uses single unit package drug distribution systems in which the quantity stored is minimal and a missing dose can be readily detected.

§ 483.70 Physical environment.

The facility must be designed, constructed, equipped, and maintained to protect the health and safety of residents, personnel and the public.

(a) Life safety from fire. Except as provided in paragraph (a)(1) or (a)(3) of this section, the facility must meet the applicable provisions of the 1985 edition of the Life Safety Code of the National Fire Protection Association (which is incorporated by reference). Incorporation of the 1985 edition of the National Fire Protection Association’s Life Safety Code (published February 7, 1985; ANSI/NFPA) was approved by the Director of the Federal Register in accordance with 5 U.S.C. 552(a) and 1 CFR part 51 that govern the use of incorporations by reference.1

(1) A facility is considered to be in compliance with this requirement as long as the facility—

(i) On November 26, 1982, complied with or without waivers, with the requirements of the 1967 or 1973 editions of the Life Safety Code and continues to remain in compliance with those editions of the Code; or

(ii) On May 9, 1988, complied, with or without waivers, with the 1981 edition

1The Code is available for inspection at the Office of the Federal Register Information Center, room 8301, 1110 L Street NW., Washington, DC. Copies may be obtained from the National Fire Protection Association, Batterymarch Park, Quincy, MA 02269. If any changes in this code are also to be incorporated by reference, a notice to that effect will be published in the FEDERAL REGISTER.
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(2) After consideration of State survey agency findings, HCFA may waive specific provisions of the Life Safety Code which, if rigidly applied would result in unreasonable hardship upon the facility, but only if the waiver does not adversely affect the health and safety of residents or personnel.

(3) The provisions of the Life Safety Code do not apply in a State where HCFA finds, in accordance with applicable provisions of sections 1819(d)(2)(B)(iii) and 1919(d)(2)(B)(ii) of the Act, that a fire and safety code imposed by State law adequately protects patients, residents and personnel in long-term care facilities.

(b) Emergency power. (1) An emergency electrical power system must supply power adequate at least for lighting all entrances and exits; equipment to maintain the fire detection, alarm, and extinguishing systems; and life support systems in the event the normal electrical supply is interrupted.

(2) When life support systems are used, the facility must provide emergency electrical power with an emergency generator (as defined in NFPA 99, Health Care Facilities) that is located on the premises.

(c) Space and equipment. The facility must—

(1) Provide sufficient space and equipment in dining, health services, recreation, and program areas to enable staff to provide residents with needed services as required by these standards and as identified in each resident's plan of care; and

(2) Maintain all essential mechanical, electrical, and patient care equipment in safe operating condition.

(d) Resident rooms. Resident rooms must be designed and equipped for adequate nursing care, comfort, and privacy of residents.

(1) Bedrooms must—

(i) Accommodate no more than four residents;

(ii) Measure at least 80 square feet per resident in multiple resident bedrooms, and at least 100 square feet in single resident rooms;

(iii) Have direct access to an exit corridor;

(iv) Be designed or equipped to assure full visual privacy for each resident;

(v) In facilities initially certified after March 31, 1992, except in private rooms, each bed must have ceiling suspended curtains, which extend around the bed to provide total visual privacy in combination with adjacent walls and curtains;

(vi) Have at least one window to the outside; and

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

(2) The facility must provide each resident with—

(i) A separate bed of proper size and height for the convenience of the resident;

(ii) A clean, comfortable mattress;

(iii) Bedding appropriate to the weather and climate; and

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

(3) HCFA, or in the case of a nursing facility the survey agency, may permit variations in requirements specified in paragraphs (d)(1)(i) and (ii) of this section relating to rooms in individual cases when the facility demonstrates in writing that the variations—

(i) Are in accordance with the special needs of the residents; and

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

(e) Toilet facilities. Each resident room must be equipped with or located near toilet and bathing facilities.

(f) Resident call system. The nurse's station must be equipped to receive resident calls through a communication system from—

(1) Resident rooms; and

(2) Toilet and bathing facilities.

(g) Dining and resident activities. The facility must provide one or more rooms designated for resident dining and activities. These rooms must—

(1) Be well lighted;

(2) Be well ventilated, with non-smoking areas identified;

(3) Be adequately furnished; and

(4) Have sufficient space to accommodate all activities.
(h) Other environmental conditions. The facility must provide a safe, functional, sanitary, and comfortable environment for the residents, staff and the public. The facility must—
(1) Establish procedures to ensure that water is available to essential areas when there is a loss of normal water supply;
(2) Have adequate outside ventilation by means of windows, or mechanical ventilation, or a combination of the two;
(3) Equip corridors with firmly secured handrails on each side; and
(4) Maintain an effective pest control program so that the facility is free of pests and rodents.

§ 483.75 Administration.
A facility must be administered in a manner that enables it to use its resources effectively and efficiently to attain or maintain the highest practicable physical, mental, and psychosocial well-being of each resident.

(a) Licensure. A facility must be licensed under applicable State and local law.

(b) Compliance with Federal, State, and local laws and professional standards. The facility must operate and provide services in compliance with all applicable Federal, State, and local laws, regulations, and codes, and with accepted professional standards and principles that apply to professionals providing services in such a facility.

(c) Relationship to other HHS regulations. In addition to compliance with the regulations set forth in this subpart, facilities are obliged to meet the applicable provisions of other HHS regulations, including but not limited to those pertaining to nondiscrimination on the basis of race, color, or national origin (45 CFR part 80); nondiscrimination on the basis of handicap (45 CFR part 84); nondiscrimination on the basis of age (45 CFR part 91); protection of human subjects of research (45 CFR part 46); and fraud and abuse (42 CFR part 455). Although these regulations are not in themselves considered requirements under this part, their violation may result in the termination or suspension of, or the refusal to grant or continue payment with Federal funds.

(d) Governing body. (1) The facility must have a governing body, or designated persons functioning as a governing body, that is legally responsible for establishing and implementing policies regarding the management and operation of the facility; and
(2) The governing body appoints the administrator who is—
(i) Licensed by the State where licensing is required; and
(ii) Responsible for management of the facility.

(e) Required training of nursing aides—
(1) Definitions. Licensed health professional means a physician; physician assistant; nurse practitioner; physical, speech, or occupational therapist; physical or occupational therapy assistant; registered professional nurse; licensed practical nurse; or licensed or certified social worker.

Nurse aide means any individual providing nursing or nursing-related services to residents in a facility who is not a licensed health professional, a registered dietitian, or someone who volunteers to provide such services without pay.

(2) General rule. A facility must not use any individual working in the facility as a nurse aide for more than 4 months, on a full-time basis, unless:
(i) That individual is competent to provide nursing and nursing-related services; and
(ii)(A) That individual has completed a training and competency evaluation program, or a competency evaluation program approved by the State as meeting the requirements of §§ 483.151–483.154 of this part; or
(B) That individual has been deemed or determined competent as provided in § 483.150 (a) and (b).

(3) Non-permanent employees. A facility must not use on a temporary, per diem, leased, or any basis other than a permanent employee any individual who does not meet the requirements in paragraphs (e)(2) (i) and (ii) of this section.

(4) Competency. A facility must not use any individual who has worked less than 4 months as a nurse aide in that facility unless the individual—
§ 483.75  
(i) Is a full-time employee in a State-approved training and competency evaluation program;  
(ii) Has demonstrated competence through satisfactory participation in a State-approved nurse aide training and competency evaluation program or competency evaluation program; or  
(iii) Has been deemed or determined competent as provided in §483.150 (a) and (b).

(5) Registry verification. Before allowing an individual to serve as a nurse aide, a facility must receive registry verification that the individual has met competency evaluation requirements unless—  
(i) The individual is a full-time employee in a training and competency evaluation program approved by the State; or  
(ii) The individual can prove that he or she has recently successfully completed a training and competency evaluation program or competency evaluation program approved by the State and has not yet been included in the registry. Facilities must follow up to ensure that such an individual actually becomes registered.

(6) Multi-State registry verification. Before allowing an individual to serve as a nurse aide, a facility must receive registry verification from every State registry established under sections 1819(e)(2)(A) or 1919(e)(2)(A) of the Act the facility believes will include information on the individual.

(7) Required retraining. If, since an individual’s most recent completion of a training and competency evaluation program, there has been a continuous period of 24 consecutive months during none of which the individual provided nursing or nursing-related services for monetary compensation, the individual must complete a new training and competency evaluation program or a new competency evaluation program.

(8) Regular in-service education. The facility must complete a performance review of every nurse aide at least once every 12 months, and must provide regular in-service education based on the outcome of these reviews. The in-service training must—  
(i) Be sufficient to ensure the continuing competence of nurse aides, but must be no less than 12 hours per year; and  
(ii) Address areas of weakness as determined in nurse aides’ performance reviews and may address the special needs of residents as determined by the facility staff; and  
(iii) For nurse aides providing services to individuals with cognitive impairments, also address the care of the cognitively impaired.

(f) Proficiency of Nurse aides. The facility must ensure that nurse aides are able to demonstrate competency in skills and techniques necessary to care for residents’ needs, as identified through resident assessments, and described in the plan of care.

(g) Staff qualifications. (1) The facility must employ on a full-time, part-time or consultant basis those professionals necessary to carry out the provisions of these requirements.

(2) Professional staff must be licensed, certified, or registered in accordance with applicable State laws.

(h) Use of outside resources. (1) If the facility does not employ a qualified professional person to furnish a specific service to be provided by the facility, the facility must have that service furnished to residents by a person or agency outside the facility under an arrangement described in section 1861(w) of the Act or (with respect to services furnished to NF residents and dental services furnished to SNF residents) an agreement described in paragraph (h)(2) of this section.

(2) Arrangements as described in section 1861(w) of the Act or agreements pertaining to services furnished by outside resources must specify in writing that the facility assumes responsibility for—  
(i) Obtaining services that meet professional standards and principles that apply to professionals providing services in such a facility; and  
(ii) The timeliness of the services.

(i) Medical director. (1) The facility must designate a physician to serve as medical director.

(2) The medical director is responsible for—  
(i) Implementation of resident care policies; and  
(ii) The coordination of medical care in the facility.

(j) Level B requirement: Laboratory services. (1) The facility must provide
or obtain laboratory services to meet the needs of its residents. The facility is responsible for the quality and timeliness of the services.

(i) If the facility provides its own laboratory services, the services must meet the applicable requirements for laboratories specified in part 493 of this chapter.

(ii) If the facility provides blood bank and transfusion services, it must meet the applicable requirements for laboratories specified in part 493 of this chapter.

(iii) If the laboratory chooses to refer specimens for testing to another laboratory, the referral laboratory must be certified in the appropriate specialties and subspecialties of services in accordance with the requirements of part 493 of this chapter.

(iv) If the facility does not provide laboratory services on site, it must have an agreement to obtain these services from a laboratory that meets the applicable requirements of part 493 of this chapter.

(2) The facility must—

(i) Provide or obtain laboratory services only when ordered by the attending physician;

(ii) Promptly notify the attending physician of the findings;

(iii) Assist the resident in making transportation arrangements to and from the source of service, if the resident needs assistance; and

(iv) File in the resident’s clinical record reports of laboratory results and any other diagnostic services.

(k) Radiology and other diagnostic services. (1) The facility must provide or obtain radiology and other diagnostic services to meet the needs of its residents. The facility is responsible for the quality and timeliness of the services.

(i) If the facility provides its own diagnostic services, the services must meet the applicable conditions of participation for hospitals contained in § 482.26 of this chapter.

(ii) If the facility does not provide its own diagnostic services, it must have an agreement to obtain these services from a provider or supplier that is approved to provide these services under Medicare.

(2) The facility must—

(i) Provide or obtain radiology and other diagnostic services only when ordered by the attending physician;

(ii) Promptly notify the attending physician of the findings;

(iii) Assist the resident in making transportation arrangements to and from the source of service, if the resident needs assistance; and

(iv) File in the resident’s clinical record signed and dated reports of x-ray and other diagnostic services.

(l) Clinical records. (1) The facility must maintain clinical records on each resident in accordance with accepted professional standards and practices that are—

(i) Complete;

(ii) Accurately documented;

(iii) Readily accessible; and

(iv) Systematically organized.

(2) Clinical records must be retained for—

(i) The period of time required by State law; or

(ii) Five years from the date of discharge when there is no requirement in State law; or

(iii) For a minor, three years after a resident reaches legal age under State law.

(3) The facility must safeguard clinical record information against loss, destruction, or unauthorized use.

(4) The facility must keep confidential all information contained in the resident’s records, regardless of the form or storage method of the records, except when release is required by—

(i) Transfer to another health care institution;

(ii) Law;

(iii) Third party payment contract; or

(iv) The resident.

(5) The clinical record must contain—

(i) Sufficient information to identify the resident;

(ii) A record of the resident’s assessments;

(iii) The plan of care and services provided;

(iv) The results of any preadmission screening conducted by the State; and

(v) Progress notes.

(m) Disaster and emergency preparedness. (1) The facility must have detailed written plans and procedures to meet
§ 483.100 Basis.

The requirements of §§483.100 through 483.138 governing the State’s responsibility for preadmission screening and annual resident review (PASARR) of individuals with mental illness and mental retardation are based on section 1919(e)(7) of the Act.

(3) A State or the Secretary may not require disclosure of the records of such committee except in so far as such disclosure is related to the compliance of such committee with the requirements of this section.

(4) Good faith attempts by the committee to identify and correct quality deficiencies will not be used as a basis for sanctions.

(p) Disclosure of ownership. (1) The facility must comply with the disclosure requirements of §§420.206 and 455.104 of this chapter.

(2) The facility must provide written notice to the State agency responsible for licensing the facility at the time of change, if a change occurs in—

(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 corporation, association, or other company responsible for the management of the facility; or

(iv) The facility’s administrator or director of nursing.

(3) The notice specified in paragraph (p)(2) of this section must include the identity of each new individual or company.


Subpart C—Preadmission Screening and Annual Review of Mentally Ill and Mentally Retarded Individuals

SOURCE: 57 FR 56506, Nov. 30, 1992, unless otherwise noted.

§ 483.100 Basis.

The requirements of §§483.100 through 483.138 governing the State’s responsibility for preadmission screening and annual resident review (PASARR) of individuals with mental illness and mental retardation are based on section 1919(e)(7) of the Act.
§ 483.102 Applicability and definitions.

(a) This subpart applies to the screening or reviewing of all individuals with mental illness or mental retardation who apply to or reside in Medicaid certified NFs regardless of the source of payment for the NF services, and regardless of the individual's or resident's known diagnoses.

(b) Definitions. As used in this subpart—

(1) An individual is considered to have a serious mental illness (MI) if the individual meets the following requirements on diagnosis, level of impairment and duration of illness:

(i) Diagnosis. The individual has a major mental disorder diagnosable under the Diagnostic and Statistical Manual of Mental Disorders, 3rd edition, revised in 1987.

Incorporation of the 1987 edition of the Diagnostic and Statistical Manual of Mental Disorders, 3rd edition, was approved by the Director of the Federal Register in accordance with 5 U.S.C. 552(a) and 1 CFR part 51 that govern the use of incorporation by reference.¹

This mental disorder is—

(A) A schizophrenic, mood, paranoid, panic or other severe anxiety disorder; somatoform disorder; personality disorder; other psychotic disorder; or another mental disorder that may lead to a chronic disability; but

(B) Not a primary diagnosis of dementia, including Alzheimer's disease or a related disorder, or a non-primary diagnosis of dementia unless the primary diagnosis is a major mental disorder as defined in paragraph (b)(1)(i)(A) of this section.

(ii) Level of impairment. The disorder results in functional limitations in major life activities within the past 3 to 6 months that would be appropriate for the individual's developmental stage. An individual typically has at least one of the following characteristics on a continuing or intermittent basis:

(A) Interpersonal functioning. The individual has serious difficulty interacting appropriately and communicating effectively with other persons, has a possible history of altercations, evictions, firing, fear of strangers, avoidance of interpersonal relationships and social isolation;

(B) Concentration, persistence, and pace. The individual has serious difficulty in sustaining focused attention for a long enough period to permit the completion of tasks commonly found in work settings or in work-like structured activities occurring in school or home settings, manifests difficulties in concentration, inability to complete simple tasks within an established time period, makes frequent errors, or requires assistance in the completion of these tasks; and

(C) Adaptation to change. The individual has serious difficulty in adapting to typical changes in circumstances associated with work, school, family, or social interaction, manifests agitation, exacerbated signs and symptoms associated with the illness, or withdrawal from the situation, or requires intervention by the mental health or judicial system.

(iii) Recent treatment. The treatment history indicates that the individual has experienced at least one of the following:

(A) Psychiatric treatment more intensive than outpatient care more than once in the past 2 years (e.g., partial hospitalization or inpatient hospitalization); or

(B) Within the last 2 years, due to the mental disorder, experienced an episode of significant disruption to the normal living situation, for which supportive services were required to maintain functioning at home, or in a residential treatment environment, or which resulted in intervention by housing or law enforcement officials.

(2) An individual is considered to have dementia if he or she has a primary diagnosis of dementia, as described in the Diagnostic and Statistical Manual of Mental Disorders, 3rd edition, revised in 1987, or a non-primary diagnosis of dementia unless the

¹The Diagnostic and Statistical Manual of Mental Disorders is available for inspection at the Health Care Financing Administration, room 132, East High Rise Building, 6325 Security Boulevard, Baltimore, Maryland, or at the Office of the Federal Register, suite 700, 800 North Capitol St. NW., Washington, DC. Copies may be obtained from the American Psychiatric Association, Division of Publications and Marketing, 1400 K Street, NW., Washington, DC 20005.
§ 483.104 State plan requirement.

As a condition of approval of the State plan, the State must operate a preadmission screening and annual resident review program that meets the requirements of §§483.100 through 438.138.

§ 483.106 Basic rule.

(a) Requirement. The State PASARR program must require—

(1) Preadmission screening of all individuals with mental illness or mental retardation who apply as new admissions to Medicaid NFs on or after January 1, 1989;

(2) Initial review, by April 1, 1990, of all current residents with mental retardation or mental illness who entered Medicaid NFs prior to January 1, 1989; and

(3) At least annual review, as of April 1, 1990, of all residents with mental illness or mental retardation, regardless of whether they were first screened under the preadmission screening or annual resident review requirements.

(b) Admissions, readmissions and interfacility transfers—

(1) New admission. An individual is a new admission if he or she is admitted to any NF for the first time or does not qualify as a readmission.

(2) Exempted hospital discharge. (i) An exempted hospital discharge means an individual—

(A) Who is admitted to any NF directly from a hospital after receiving acute inpatient care at the hospital;

(B) Who requires NF services for the condition for which he or she received care in the hospital; and

(C) Whose attending physician has certified before admission to the facility that the individual is likely to require less than 30 days nursing facility services.

(ii) If an individual who enters a NF as an exempted hospital discharge is later found to require more than 30 days of NF care, the State mental health or mental retardation authority must conduct an annual resident review within 40 calendar days of admission.

(3) Readmissions. An individual is a readmission if he or she was readmitted to a facility from a hospital to which he or she was transferred for the purpose of receiving care. Readmissions are subject to annual resident review rather than preadmission screening.

(4) Interfacility transfers—

(i) An interfacility transfer occurs when an individual is transferred from one NF to another NF, with or without an intervening hospital stay. Interfacility transfers are subject to annual resident review rather than preadmission screening.

(ii) In cases of transfer of a resident with MI or MR from a NF to a hospital or to another NF, the transferring NF is responsible for ensuring that copies of the resident’s most recent PASARR and resident assessment reports accompany the transferring resident.

2The American Association on Mental Retardation’s Manual on Classification in Mental Retardation is available for inspection at the Health Care Financing Administration, Room 132, East High Rise Building, 620 Security Boulevard, Baltimore, Maryland, or at the Office of the Federal Register Information Center, Suite 700, 800 North Capitol St. NW., Washington, DC. Copies may be obtained from the American Association on Mental Retardation, 1719 Kalorama Rd., NW., Washington, DC 20009.
(c) Purpose. The preadmission screening and annual resident review process must result in determinations based on a physical and mental evaluation of each individual with mental illness or mental retardation, that are described in §§483.112 and 483.114.

(d) Responsibility for evaluations and determinations. The PASARR determinations of whether an individual requires the level of services provided by a NF and whether specialized services are needed—

(1) For individuals with mental illness, must be made by the State mental health authority and be based on an independent physical and mental evaluation performed by a person or entity other than the State mental health authority; and

(2) For individuals with mental retardation, must be made by the State mental retardation or developmental disabilities authority.

(e) Delegation of responsibility—(1) The State mental health and mental retardation authorities may delegate by subcontract or otherwise the evaluation and determination functions for which they are responsible to another entity only if—

(i) The State mental health and mental retardation authorities retain ultimate control and responsibility for the performance of their statutory obligations;

(ii) The two determinations as to the need for NF services and for specialized services are made, based on a consistent analysis of the data; and

(iii) The entity to which the delegation is made is not a NF or an entity that has a direct or indirect affiliation or relationship with a NF.

(2) The State mental health authority has responsibility for both the evaluation and determination functions for individuals with MR whereas the State mental health authority has responsibility only for the determination function.

(3) The evaluation of individuals with MI cannot be delegated by the State mental health authority because it does not have responsibility for this function. The evaluation function must be performed by a person or entity other than the State mental health authority. In designating an independent person or entity to perform MI evaluations, the State must not use a NF or an entity that has a direct or indirect affiliation or relationship with a NF.

[57 FR 56506, Nov. 30, 1992; 58 FR 25784, Apr. 28, 1993]

§ 483.108 Relationship of PASARR to other Medicaid processes.

(a) PASARR determinations made by the State mental health or mental retardation authorities cannot be countermanded by the State Medicaid agency, either in the claims process or through other utilization control/review processes or by the State survey and certification agency. Only appeals determinations made through the system specified in subpart E of this part may overturn a PASARR determination made by the State mental health or mental retardation authorities.

(b) In making their determinations, however, the State mental health and mental retardation authorities must not use criteria relating to the need for NF care or specialized services that are inconsistent with this regulation and any supplementary criteria adopted by the State Medicaid agency under its approved State plan.

(c) To the maximum extent practicable, in order to avoid duplicative testing and effort, the PASARR must be coordinated with the routine resident assessments required by §483.20(b).

§ 483.110 Out-of-State arrangements.

(a) Basic rule. The State in which the individual is a State resident (or would be a State resident at the time he or she becomes eligible for Medicaid), as defined in §435.403 of this chapter, must pay for the PASARR and make the required determinations, in accordance with §431.52(b).

(b) Agreements. A State may include arrangements for PASARR in its provider agreements with out-of-State facilities or reciprocal interstate agreements.

[57 FR 56506, Nov. 30, 1992; 58 FR 25784, Apr. 28, 1993]

§ 483.112 Preadmission screening of applicants for admission to NFs.

(a) Determination of need for NF services. For each NF applicant with MI or
MR, the State mental health or mental retardation authority (as appropriate) must determine, in accordance with §483.130, whether, because of the resident’s physical and mental condition, the individual requires the level of services provided by a NF.

(b) Determination of need for specialized services. If the individual with mental illness or mental retardation is determined to require a NF level of care, the State mental health or mental retardation authority (as appropriate) must also determine, in accordance with §483.130, whether the individual requires specialized services for the mental illness or mental retardation, as defined in §483.120.

(c) Timeliness—(1) Except as specified in paragraph (c)(4) of this section, a preadmission screening determination must be made in writing within an annual average of 7 to 9 working days of referral of the individual with MI or MR by whatever agent performs the Level I identification, under §483.128(a) of this part, to the State mental health or mental retardation authority for screening. (See §483.128(a) for discussion of Level I evaluation.)

(2) The State may convey determinations verbally to nursing facilities and the individual and confirm them in writing.

(3) The State may compute separate annual averages for the mentally ill and the mentally retarded/developmentally disabled populations.

(4) The Secretary may grant an exception to the timeliness standard in paragraph (c)(1) of this section when the State—

(i) Exceeds the annual average; and

(ii) Provides justification satisfactory to the Secretary that a longer time period was necessary.

§483.114 Annual review of NF residents.

(a) Individuals with mental illness. For each resident of a NF who has mental illness, the State mental health authority must determine in accordance with §483.130 whether, because of the resident’s physical and mental condition, the resident requires—

(1) The level of services provided by—

(i) A NF;

(ii) An inpatient psychiatric hospital for individuals under age 21, as described in section 1905(h) of the Act; or

(iii) An institution for mental diseases providing medical assistance to individuals age 65 or older; and

(2) Specialized services for mental illness, as defined in §483.120.

(b) Individuals with mental retardation. For each resident of a NF who has mental retardation, the State mental retardation or developmental disability authority must determine in accordance with §483.130 whether, because of his or her physical or mental condition, the resident requires—

(1) The level of services provided by a NF or an intermediate care facility for the mentally retarded; and

(2) Specialized services for mental retardation as defined in §483.120.

(c) Frequency of review—(1) A review and determination must be conducted for each resident of a Medicaid NF who has mental illness or mental retardation not less often than annually.

(2) “Annually” is defined as occurring within every fourth quarter after the previous preadmission screen or annual resident review.

(d) April 1, 1990 deadline for initial reviews. The first set of annual reviews on residents who entered the NF prior to January 1, 1989, must be completed by April 1, 1990.

§483.116 Residents and applicants determined to require NF level of services.

(a) Individuals needing NF services. If the State mental health or mental retardation authority determines that a resident or applicant for admission to a NF requires a NF level of services, the NF may admit or retain the individual.

(b) Individuals needing NF services and specialized services. If the State mental health or mental retardation authority determines that a resident or applicant for admission requires both a NF level of services and specialized services for the mental illness or mental retardation—

(1) The NF may admit or retain the individual; and

(2) The State must provide or arrange for the provision of the specialized services needed by the individual while he or she resides in the NF.
§ 483.118 Residents and applicants determined not to require NF level of services.

(a) Applicants who do not require NF services. If the State mental health or mental retardation authority determines that an applicant for admission to a NF does not require NF services, the applicant cannot be admitted. NF services are not a covered Medicaid service for that individual, and further screening is not required.

(b) Residents who require neither NF services nor specialized services for MI or MR. If the State mental health or mental retardation authority determines that a resident requires neither the level of services provided by a NF nor specialized services for MI or MR, regardless of the length of stay in the facility, the State must—

(1) Arrange for the safe and orderly discharge of the resident from the facility in accordance with §483.12(a); and

(2) Prepare and orient the resident for discharge.

(c) Residents who do not require NF services but require specialized services for MI or MR—(1) Long term residents. Except as otherwise may be provided in an alternative disposition plan adopted under section 1919(e)(7)(E) of the Act, for any resident who has continuously resided in a NF for at least 30 months before the date of the determination, and who requires only specialized services, as defined in §483.120, and who has not continuously resided in a NF for at least 30 months before the date of the determination, the State must, in consultation with the resident's family or legal representative and caregivers—

(i) Offer the resident the choice of remaining in the facility or of receiving services in an alternative appropriate setting;

(ii) Inform the resident of the institutional and noninstitutional alternatives covered under the State Medicaid plan for the resident;

(iii) Clarify the effect on eligibility for Medicaid services under the State plan if the resident chooses to leave the facility, including its effect on re-admission to the facility; and

(iv) Regardless of the resident's choice, provide for, or arrange for the provision of specialized services for the mental illness or mental retardation.

(2) Short term residents. Except as otherwise may be provided in an alternative disposition plan adopted under section 1919(e)(7)(E) of the Act, for any resident who requires only specialized services, as defined in §483.120, and who has not continuously resided in a NF for at least 30 months before the date of the determination, the State must, in consultation with the resident's family or legal representative and caregivers—

(i) Arrange for the safe and orderly discharge of the resident from the facility in accordance with §483.12(a);

(ii) Prepare and orient the resident for discharge; and

(iii) Provide for, or arrange for the provision of, specialized services for the mental illness or mental retardation.

(3) For the purpose of establishing length of stay in a NF, the 30 months of continuous residence in a NF or longer—

(i) Is calculated back from the date of the first annual resident review determination which finds that the individual is not in need of NF level of services;

(ii) May include temporary absences for hospitalization or therapeutic leave; and

(iii) May consist of consecutive residences in more than one NF.

§ 483.120 Specialized services.

(a) Definition—(1) For mental illness, specialized services means the services specified by the State which, combined with services provided by the NF, results in the continuous and aggressive implementation of an individualized plan of care that—

(i) Is developed and supervised by an interdisciplinary team, which includes a physician, qualified mental health professionals and, as appropriate, other professionals.

(ii) Prescribes specific therapies and activities for the treatment of persons experiencing an acute episode of serious mental illness, which necessitates supervision by trained mental health personnel; and

(iii) Is directed toward diagnosing and reducing the resident's behavioral symptoms that necessitated institutionalization, improving his or her level of independent functioning, and

§ 483.120 Specialized services.

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(iii) Is directed toward diagnosing and reducing the resident's behavioral symptoms that necessitated institutionalization, improving his or her level of independent functioning, and
achieving a functioning level that permits reduction in the intensity of mental health services to below the level of specialized services at the earliest possible time.

(2) For mental retardation, specialized services means the services specified by the State which, combined with services provided by the NF or other service providers, results in treatment which meets the requirements of §483.440(a)(1).

(b) Who must receive specialized services. The State must provide or arrange for the provision of specialized services, in accordance with this subpart, to all NF residents with MI or MR whose needs are such that continuous supervision, treatment and training by qualified mental health or mental retardation personnel is necessary, as identified by the screening provided in §483.130 or §§483.134 and 483.136.

(c) Services of lesser intensity than specialized services. The NF must provide mental health or mental retardation services which are of a lesser intensity than specialized services to all residents who need such services.

§483.122 FFP for NF services.

(a) Basic rule. Except as otherwise may be provided in an alternative disposition plan adopted under section 1919(e)(7)(E) of the Act, FFP is available in State expenditures for NF services provided to a Medicaid eligible individual subject to the requirements of this part only if the individual has been determined—

(1) To need NF care under §483.116(a) or

(2) Not to need NF services but to need specialized services, meets the requirements of §483.118(c)(1), and elects to stay in the NF.

(b) FFP for late reviews. When a predischARGE screening has not been performed prior to admission or an annual review is not performed timely, in accordance with §483.114(c), but either is performed at a later date, FFP is available only for services furnished after the screening or review has been performed, subject to the provisions of paragraph (a) of this section.

§483.124 FFP for specialized services.

FFP is not available for specialized services furnished to NF residents as NF services.

§483.126 Appropriate placement.

Placement of an individual with MI or MR in a NF may be considered appropriate only when the individual's needs are such that he or she meets the minimum standards for admission and the individual's needs for treatment do not exceed the level of services which can be delivered in the NF to which the individual is admitted either through NF services alone or, where necessary, through NF services supplemented by services provided by or arranged for by the State.

§483.128 PASARR evaluation criteria.

(a) Level I: Identification of individuals with MI or MR. The State's PASARR program must identify all individuals who are suspected of having MI or MR as defined in §483.102. This identification function is termed Level I. Level II is the function of evaluating and determining whether NF services and specialized services are needed. The State's performance of the Level I identification function must provide at least, in the case of first time identifications, for the issuance of written noticce to the individual or resident and his or her legal representative that the individual or resident is suspected of having MI or MR and is being referred to the State mental health or mental retardation authority for Level II screening.

(b) Adaptation to culture, language, ethnic origin. Evaluations performed under PASARR and PASARR notices must be adapted to the cultural background, language, ethnic origin and means of communication used by the individual being evaluated.

(c) Participation by individual and family. PASARR evaluations must involve—

(1) The individual being evaluated;

(2) The individual's legal representative, if one has been designated under State law; and

(3) The individual's family if—

(i) Available; and
Health Care Financing Administration, HHS § 483.128

(ii) The individual or the legal representative agrees to family participation.

(d) Interdisciplinary coordination. When parts of a PASARR evaluation are performed by more than one evaluator, the State must ensure that there is interdisciplinary coordination among the evaluators.

(e) The State's PASARR program must use at least the evaluative criteria of § 483.130 (if one or both determinations can easily be made categorically as described in § 483.130) or of §§ 483.132 and 483.134 or § 483.136 (or, in the case of individuals with both MI and MR, §§ 483.132, 483.134 and 483.136 if a more extensive individualized evaluation is required).

(f) Data. In the case of individualized evaluations, information that is necessary for determining whether it is appropriate for the individual with MI or MR to be placed in an NF or in another appropriate setting should be gathered throughout all applicable portions of the PASARR evaluation (§§ 483.132 and 483.134 and/or § 483.136). The two determinations relating to the need for NF level of care and specialized services are interrelated and must be based upon a comprehensive analysis of all data concerning the individual.

(g) Preexisting data. Evaluators may use relevant evaluative data, obtained prior to initiation of preadmission screening or annual resident review, if the data are considered valid and accurate and reflect the current functional status of the individual. However, in the case of individualized evaluations, to supplement and verify the currency and accuracy of existing data, the State's PASARR program may need to gather additional information necessary to assess proper placement and treatment.

(h) Findings. For both categorical and individualized determinations, findings of the evaluation must correspond to the person's current functional status as documented in medical and social history records.

(i) Evaluation report: Individualized determinations. For individualized PASARR determinations, findings must be issued in the form of a written evaluative report which—

(1) Identifies the name and professional title of person(s) who performed the evaluation(s) and the date on which each portion of the evaluation was administered;

(2) Provides a summary of the medical and social history, including the positive traits or developmental strengths and weaknesses or developmental needs of the evaluated individual;

(3) If NF services are recommended, identifies the specific services which are required to meet the evaluated individual's needs, including services required in paragraph (i)(5) of this section;

(4) If specialized services are not recommended, identifies any specific mental retardation or mental health services which are of a lesser intensity than specialized services that are required to meet the evaluated individual's needs;

(5) If specialized services are recommended, identifies the specific mental retardation or mental health services required to meet the evaluated individual's needs; and

(6) Includes the bases for the report's conclusions.

(j) Evaluation report: Categorical determinations. For categorical PASARR determinations, findings must be issued in the form of an abbreviated written evaluative report which—

(1) Identifies the name and professional title of the person applying the categorical determination and the data on which the application was made;

(2) Explains the categorical determination(s) that has (have) been made and, if only one of the two required determinations can be made categorically, describes the nature of any further screening which is required;

(3) Identifies, to the extent possible, based on the available data, NF services, including any mental health or specialized psychiatric rehabilitative services, that may be needed; and

(4) Includes the bases for the report's conclusions.

(k) Interpretation of findings to individual. For both categorical and individualized determinations, findings of the evaluation must be interpreted and explained to the individual and, where
applicable, to a legal representative designated under State law.

(1) Evaluation report. The evaluator must send a copy of the evaluation report to the—

(1) Individual or resident and his or her legal representative;

(2) Appropriate State authority in sufficient time for the State authorities to meet the times identified in §483.112(c) for PASs and §483.114(c) for ARRs;

(3) Admitting or retaining NF;

(4) Individual’s attending physician; and

(5) The discharging hospital if the individual is seeking NF admission from a hospital.

(m) The evaluation may be terminated if the evaluator finds at any time during the evaluation that the individual being evaluated—

(1) Does not have MI or MR; or

(2) Has—

(i) A primary diagnosis of dementia (including Alzheimer’s Disease or a related disorder); or

(ii) A non-primary diagnosis of dementia without a primary diagnosis that is a serious mental illness, and does not have a diagnosis of MR or a related condition.

[57 FR 56506, Nov. 30, 1992; 58 FR 25784, Apr. 28, 1993]

§ 483.130 PASARR determination criteria.

(a) Basis for determinations. Determinations made by the State mental health or mental retardation authority as to whether NF level of services and specialized services are needed must be based on an evaluation of data concerning the individual, as specified in paragraph (b) of this section.

(b) Types of determinations. Determinations may be—

(1) Advance group determinations, in accordance with this section, by category that take into account that certain diagnoses, levels of severity of illness, or need for a particular service clearly indicate that admission to or residence in a NF is normally needed, or that the provision of specialized services is not normally needed; or

(2) Individualized determinations based on more extensive individualized evaluations as required in §483.132.

(c) Group determinations by category. Advance group determinations by category developed by the State mental health or mental retardation authorities may be made applicable to individuals by the NF or other evaluator following Level I review only if existing data on the individual appear to be current and accurate and are sufficient to allow the evaluator readily to determine that the individual fits into the category established by the State authorities (see §483.132(c)). Sources of existing data on the individual that could form the basis for applying a categorical determination by the State authorities would be hospital records, physician’s evaluations, election of hospice status, records of community mental health centers or community mental retardation or developmental disability providers.

(d) Examples of categories. Examples of categories for which the State mental health or mental retardation authority may make an advance group determination that NF services are needed are—

(1) Convalescent care from an acute physical illness which—

(i) Required hospitalization; and

(ii) Does not meet all the criteria for an exempt hospital discharge, which is not subject to preadmission screening, as specified in §483.106(b)(2).

(2) Terminal illness, as defined for hospice purposes in §418.3 of this chapter;

(3) Severe physical illnesses such as coma, ventilator dependence, functioning at a brain stem level, or diagnoses such as chronic obstructive pulmonary disease, Parkinson’s disease, Huntington’s disease, amyotrophic lateral sclerosis, and congestive heart failure which result in a level of impairment so severe that the individual could not be expected to benefit from specialized services;

(4) Provisional admissions pending further assessment in cases of delirium where an accurate diagnosis cannot be made until the delirium clears;

(5) Provisional admissions pending further assessment in emergency situations requiring protective services,
(6) Very brief and finite stays of up to a fixed number of days to provide respite to in-home caregivers to whom the individual with MI or MR is expected to return following the brief NF stay.

e) Time limits. The State may specify time limits for categorical determinations that NF services are needed and in the case of paragraphs (d)(4), (5) and (6) of this section, must specify a time limit which is appropriate for provisional admissions pending further assessment and for emergency situations and respite care. If an individual is later determined to need a longer stay than the State’s limit allows, the individual must be subjected to an annual resident review before continuation of the stay may be permitted and payment made for days of NF care beyond the State’s time limit.

(f) The State mental health and mental retardation authorities may make categorical determinations that specialized services are not needed in the provisional, emergency and respite admission situations identified in §483.130(d)(4)–(6). In all other cases, except for §483.130(h), a determination that specialized services are not needed must be based on a more extensive individualized evaluation under §483.134 or §483.136.

(g) Categorical determinations: No positive specialized treatment determinations. The State mental health and mental retardation authorities must not make categorical determinations that specialized services are needed. Such a determination must be based on a more extensive individualized evaluation under §483.134 or §483.136 to determine the exact nature of the specialized services that are needed.

(h) Categorical determinations: Dementia and MR. The State mental retardation authority may make categorical determinations that individuals with dementia, which exists in combination with mental retardation or a related condition, do not need specialized services.

(i) If a State mental health or mental retardation authority determines NF needs by category, it may not waive the specialized services determination. The appropriate State authority must also determine whether specialized services are needed either by category (if permitted) or by individualized evaluations, as specified in §483.134 or §483.136.

(j) Recording determinations. All determinations made by the State mental health and mental retardation authority, regardless of how they are arrived at, must be recorded in the individual’s record.

(k) Notice of determination. The State mental health or mental retardation authority must notify in writing the following entities of a determination made under this subpart:

(1) The evaluated individual and his or her legal representative;
(2) The admitting or retaining NF;
(3) The individual or resident’s attending physician; and
(4) The discharging hospital, unless the individual is exempt from preadmission screening as provided for at §483.106(b)(2). (l) Contents of notice. Each notice of the determination made by the State mental health or mental retardation authority must include—

(1) Whether a NF level of services is needed;  
(2) Whether specialized services are needed;  
(3) The placement options that are available to the individual consistent with these determinations; and

(4) The rights of the individual to appeal the determination under subpart E of this part.

(m) Placement options. Except as otherwise may be provided in an alternative disposition plan adopted under section 1919(e)(7)(E) of the Act, the placement options and the required State actions are as follows:

(1) Can be admitted to a NF. Any applicant for admission to a NF who has MI or MR and who requires the level of services provided by a NF, regardless of whether specialized services are also needed, may be admitted to a NF, if the placement is appropriate, as determined in §483.126. If specialized services are also needed, the State is responsible for providing or arranging for the provision of the specialized services.

(2) Cannot be admitted to a NF. Any applicant for admission to a NF who...
§483.132 Evaluating the need for NF services and NF level of care (PASARR/NF).

(a) Basic rule. For each applicant for admission to a NF and each NF resident who has MI or MR, the evaluator must assess whether—

1. The individual's total needs are such that his or her needs can be met in an appropriate community setting;

2. The individual's total needs are such that they can be met only on an inpatient basis, which may include the option of placement in a home and community-based services waiver program, but for which the inpatient care would be required;

3. If inpatient care is appropriate and desired, the NF is an appropriate institutional setting for meeting those needs in accordance with §483.126; or

(b) Specialized services needed in a NF.

If a determination is made to admit or allow to remain in a NF any individual who requires specialized services, the determination must be supported by assurances that the specialized services that are needed can and will be provided or arranged for by the State while the individual resides in the NF.

(c) Record retention. The State PASARR system must maintain records of evaluations and determinations, regardless of whether they are performed categorically or individually, in order to support its determinations and actions and to protect the appeal rights of individuals subject to PASARR; and

(d) Tracking system. The State PASARR system must establish and maintain a tracking system for all individuals with MI or MR in NFs to ensure that appeals and future reviews are performed in accordance with this subpart and subpart E.

[57 FR 56506, Nov. 30, 1992; 58 FR 25784, Apr. 28, 1993]
(4) If the inpatient care is appropriate and desired but the NF is not the appropriate setting for meeting the individual's needs in accordance with §483.126, another setting such as an ICF/MR (including small, community-based facilities), an IMD providing services to individuals aged 65 or older, or a psychiatric hospital is an appropriate institutional setting for meeting those needs.

(b) Determining appropriate placement. In determining appropriate placement, the evaluator must prioritize the physical and mental needs of the individual being evaluated, taking into account the severity of each condition.

(c) Data. At a minimum, the data relied on to make a determination must include:

(1) Evaluation of physical status (for example, diagnoses, date of onset, medical history, and prognosis);
(2) Evaluation of mental status (for example, diagnoses, date of onset, medical history, likelihood that the individual may be a danger to himself/herself or others); and
(3) Functional assessment (activities of daily living).

(d) Based on the data compiled in §483.132 and, as appropriate, in §§483.134 and 483.136, the State mental health or mental retardation authority must determine whether an NF level of services is needed.

§ 483.134 Evaluating whether an individual with mental illness requires specialized services (PASARR/MI).

(a) Purpose. The purpose of this section is to identify the minimum data needs and process requirements for the State mental health authority, which is responsible for determining whether or not the applicant or resident with MI, as defined in §483.102(b)(1) of this part, needs a specialized services program for mental illness as defined in §483.120.

(b) Data. Minimum data collected must include—

(1) A comprehensive history and physical examination of the person. The following areas must be included (if not previously addressed):

(i) Complete medical history;
(ii) Review of all body systems;
(iii) Specific evaluation of the person's neurological system in the areas of motor functioning, sensory functioning, gait, deep tendon reflexes, cranial nerves, and abnormal reflexes; and
(iv) In case of abnormal findings which are the basis for an NF placement, additional evaluations conducted by appropriate specialists.

(2) A comprehensive drug history including current or immediate past use of medications that could mask symptoms or mimic mental illness.

(3) A psychosocial evaluation of the person, including current living arrangements and medical and support systems.

(4) A comprehensive psychiatric evaluation including a complete psychiatric history, evaluation of intellectual functioning, memory functioning, and orientation, description of current attitudes and overt behaviors, affect, suicidal or homicidal ideation, paranoia, and degree of reality testing (presence and content of delusions) and hallucinations.

(5) A functional assessment of the individual's ability to engage in activities of daily living and the level of support that would be needed to assist the individual to perform these activities while living in the community. The assessment must determine whether this level of support can be provided to the individual in an alternative community setting or whether the level of support needed is such that NF placement is required.

(6) The functional assessment must address the following areas: Self-monitoring of health status, self-administering and scheduling of medical treatment, including medication compliance, or both, self-monitoring of nutritional status, handling money, dressing appropriately, and grooming.

(c) Personnel requirements. (1) If the history and physical examination are not performed by a physician, then a physician must review and concur with the conclusions.

(2) The State may designate the mental health professionals who are qualified—

(i) To perform the evaluations required under paragraph (b) (2)–(6) of this section including the—

(A) Comprehensive drug history;
(B) Psychosocial evaluation;
(C) Comprehensive psychiatric evaluation;
(D) Functional assessment; and
(ii) To make the determination required in paragraph (d) of this section.

(d) Data interpretation. Based on the data compiled, a qualified mental health professional, as designated by the State, must validate the diagnosis of mental illness and determine whether a program of psychiatric specialized services is needed.

§ 483.136 Evaluating whether an individual with mental retardation requires specialized services (PASARR/MR).

(a) Purpose. The purpose of this section is to identify the minimum data needs and process requirements for the State mental retardation authority to determine whether or not the applicant or resident with mental retardation, as defined in § 483.102(b)(3) of this part, needs a continuous specialized services program, which is analogous to active treatment, as defined in §§ 435.1009 and 483.440 of this chapter.

(b) Data. Minimum data collected must include the individual's comprehensive history and physical examination results to identify the following information or, in the absence of data, must include information that permits a reviewer specifically to assess:

(1) The individual's medical problems;
(2) The level of impact these problems have on the individual's independent functioning;
(3) All current medications used by the individual and the current response of the individual to any prescribed medications in the following drug groups:
   (i) Hypnotics,
   (ii) Antipsychotics (neuroleptics),
   (iii) Mood stabilizers and antidepressants,
   (iv) Antianxiety-sedative agents, and
   (v) Anti-Parkinson agents,
(4) Self-monitoring of health status;
(5) Self-administering and scheduling of medical treatments;
(6) Self-monitoring of nutritional status;
(7) Self-help development such as toileting, dressing, grooming, and eating;
(8) Sensorimotor development, such as ambulation, positioning, transfer skills, gross motor dexterity, visual motor perception, fine motor dexterity, eye-hand coordination, and extent to which prosthetic, orthotic, corrective or mechanical supportive devices can improve the individual's functional capacity;
(9) Speech and language (communication) development, such as expressive language (verbal and nonverbal), receptive language (verbal and nonverbal), extent to which non-oral communication systems can improve the individual's function capacity, auditory functioning, and extent to which amplification devices (for example, hearing aid) or a program of amplification can improve the individual's functional capacity;
(10) Social development, such as interpersonal skills, recreation-leisure skills, and relationships with others;
(11) Academic/educational development, including functional learning skills;
(12) Independent living development such as meal preparation, budgeting and personal finances, survival skills, mobility skills (orientation to the neighborhood, town, city), laundry, housekeeping, shopping, bedmaking, care of clothing, and orientation skills (for individuals with visual impairments);
(13) Vocational development, including present vocational skills;
(14) Affective development such as interests, and skills involved with expressing emotions, making judgments, and making independent decisions; and
(15) The presence of identifiable maladaptive or inappropriate behaviors of the individual based on systematic observation (including, but not limited to, the frequency and intensity of identified maladaptive or inappropriate behaviors).

(c) Data interpretation—(1) The State must ensure that a licensed psychologist identifies the intellectual functioning measurement of individuals with MR or a related condition.

(2) Based on the data compiled in paragraph (b) of this section, the State mental retardation authority, using appropriate personnel, as designated by...
the State, must validate that the individual has MR or is a person with a related condition and must determine whether specialized services for mental retardation are needed. In making this determination, the State mental retardation authority must make a qualitative judgment on the extent to which the person’s status reflects, singly and collectively, the characteristics commonly associated with the need for specialized services, including—

(i) Inability to—

(A) Take care of the most personal care needs;
(B) Understand simple commands;
(C) Communicate basic needs and wants;
(D) Be employed at a productive wage level without systematic long term supervision or support;
(E) Learn new skills without aggressive and consistent training;
(F) Apply skills learned in a training situation to other environments or settings without aggressive and consistent training;
(G) Demonstrate behavior appropriate to the time, situation or place without direct supervision; and
(H) Make decisions requiring informed consent without extreme difficulty;

(ii) Demonstration of severe maladaptive behavior(s) that place the person or others in jeopardy to health and safety; and

(iii) Presence of other skill deficits or specialized training needs that necessitate the availability of trained MR personnel, 24 hours per day, to teach the person functional skills.

[57 FR 56506, Nov. 30, 1992; 58 FR 25784, Apr. 28, 1993]

§ 483.150 Statutory basis; Deemed meeting or waiver of requirements.

(a) Statutory basis. This subpart is based on sections 1819(b)(5) and 1919(b)(5) of the Act, which establish standards for training nurse-aides and for evaluating their competency.

(b) Deemed meeting of requirements. A nurse aide is deemed to satisfy the requirement of completing a training and competency evaluation program before July 1, 1989 if—

(1) The aide would have satisfied this requirement if—

(i) At least 60 hours were substituted for 75 hours in sections 1819(f)(2) and 1919(f)(2) of the Act, and
(ii) The individual has made up at least the difference in the number of hours in the program he or she completed and 75 hours in supervised practical nurse aide training or in regular in-service nurse aide education;

or

(2) The individual was found to be competent (whether or not by the State) after the completion of nurse aide training of at least 100 hours duration.

(c) Waiver of requirements. A State may—

(1) Waive the requirement for an individual to complete a competency evaluation program approved by the State for any individual who can demonstrate to the satisfaction of the State that he or she has served as a nurse aide at one or more facilities of the same employer in the state for at
least 24 consecutive months before December 19, 1989; or
(2) Deem an individual to have completed a nurse aide training and competency evaluation program approved by the State if the individual completed, before July 1, 1989, such a program that the State determines would have met the requirements for approval at the time it was offered.

§ 483.151 State review and approval of nurse aide training and competency evaluation programs and competency evaluation programs.

(a) State review and administration. (1) The State—
(i) Must specify any nurse aide training and competency evaluation programs that the State approves; and
(ii) May choose to offer a nurse aide training and competency evaluation program.
(2) If the State does not choose to offer a nurse aide training and competency evaluation program offered by or in a facility which, in the previous two years—
(i) In the case of a skilled nursing facility, has operated under a waiver under section 1819(b)(4)(C)(i) of the Act for a period in excess of 48 hours per week;
(ii) In the case of a nursing facility, has operated under a waiver under section 1919(b)(4)(C)(ii) of the Act that was granted on the basis of a demonstration that the facility is unable to provide nursing care required under section 1919(b)(4)(C)(i) of the Act for a period in excess of 48 hours per week;
(iii) Has been subject to an extended (or partial extended) survey under sections 1819(g)(2)(B)(i) or 1919(g)(2)(B)(i) of the Act;
(iv) Has been assessed a civil money penalty described in section 1819(h)(2)(B)(i) or 1919(h)(2)(A)(ii) of the Act of not less than $5,000; or
(v) Has been subject to a remedy described in sections 1819(h)(2)(B)(i) or 1919(h)(2)(A)(ii) of the Act,
(3) A State may not, until two years since the assessment of the penalty (or penalties) has elapsed, approve a nurse aide training and competency evaluation program offered by or in a facility that, within the two-year period beginning October 1, 1988—
(i) Had its participation terminated under title XVIII of the Act or under the State plan under title XIX of the Act;
(ii) Was subject to a denial of payment under title XVIII or title XIX;
(iii) Was assessed a civil money penalty of not less than $5,000 for deficiencies in nursing facility standards;
(iv) Operated under temporary management appointed to oversee the operation of the facility and to ensure the health and safety of its residents; or
(v) Pursuant to State action, was closed or had its residents transferred.
(b) Requirements for approval of programs. (1) Before the State approves a nurse aide training and competency evaluation program or competency evaluation program, the State must—
(i) Determine whether the nurse aide training and competency evaluation program meets the course requirements of §§ 483.152;
(ii) Determine whether the nurse aide training and competency evaluation program meets the course requirements of §§ 483.152; and
(iii) In all reviews other than the initial review, visit the entity providing the program.
(2) The State may not approve a nurse aide training and competency evaluation program or competency evaluation program offered by or in a facility which, in the previous two years—
(i) In the case of a skilled nursing facility, has operated under a waiver under section 1819(b)(4)(C)(i) of the Act for a period in excess of 48 hours per week;
(ii) In the case of a nursing facility, has operated under a waiver under section 1919(b)(4)(C)(ii) of the Act that was granted on the basis of a demonstration that the facility is unable to provide nursing care required under section 1919(b)(4)(C)(i) of the Act for a period in excess of 48 hours per week;
(iii) Has been subject to an extended (or partial extended) survey under sections 1819(g)(2)(B)(i) or 1919(g)(2)(B)(i) of the Act;
(iv) Has been assessed a civil money penalty described in section 1819(h)(2)(B)(i) or 1919(h)(2)(A)(ii) of the Act of not less than $5,000; or
(v) Has been subject to a remedy described in sections 1819(h)(2)(B)(i) or 1919(h)(2)(A)(ii) of the Act,
(c) Time frame for acting on a request for approval. The State must, within 90 days of the date of a request under
paragraph (a)(3) of this section or receipt of additional information from the requester—

(1) Advise the requester whether or not the program has been approved; or

(2) Request additional information from the requesting entity.

(d) Duration of approval. The State may not grant approval of a nurse aide training and competency evaluation program for a period longer than 2 years. A program must notify the State and the State must review that program when there are substantive changes made to that program within the 2-year period.

(e) Withdrawal of approval. (1) The State must withdraw approval of a nurse aide training and competency evaluation program or nurse aide competency evaluation program offered by or in a facility described in paragraph (b)(2) of this section.

(2) The State may withdraw approval of a nurse aide training and competency evaluation program or nurse aide competency evaluation program if the State determines that any of the applicable requirements of §§ 483.152 or 483.154 are not met by the program.

(3) The State must withdraw approval of a nurse aide training and competency evaluation program or a nurse aide competency evaluation program if the entity providing the program refuses to permit unannounced visits by the State.

(4) If a State withdraws approval of a nurse aide training and competency evaluation program or competency evaluation program—

(i) The State must notify the program in writing, indicating the reason(s) for withdrawal of approval of the program.

(ii) Students who have started a training and competency evaluation program from which approval has been withdrawn must be allowed to complete the course.

§ 483.152 Requirements for approval of a nurse aide training and competency evaluation program.

(a) For a nurse aide training and competency evaluation program to be approved by the State, it must, at a minimum—

(1) Consist of no less than 75 clock hours of training;

(2) Include at least the subjects specified in paragraph (b) of this section;

(3) Include at least 16 hours of supervised practical training. Supervised practical training means training in a laboratory or other setting in which the trainee demonstrates knowledge while performing tasks on an individual under the direct supervision of a registered nurse or a licensed practical nurse;

(4) Ensure that—

(i) Students do not perform any services for which they have not trained and been found proficient by the instructor; and

(ii) Students who are providing services to residents are under the general supervision of a licensed nurse or a registered nurse;

(5) Meet the following requirements for instructors who train nurse aides;

(i) The training of nurse aides must be performed by or under the general supervision of a registered nurse who possesses a minimum of 2 years of nursing experience, at least 1 year of which must be in the provision of long term care facility services;

(ii) Instructors must have completed a course in teaching adults or have experience in teaching adults or supervising nurse aides;

(iii) In a facility-based program, the training of nurse aides may be performed under the general supervision of the director of nursing for the facility who is prohibited from performing the actual training; and

(iv) Other personnel from the health professions may supplement the instructor, including, but not limited to, registered nurses, licensed practical/vocational nurses, pharmacists, dietitians, social workers, sanitariums, fire safety experts, nursing home administrators, gerontologists, psychologists, physical and occupational therapists, activities specialists, speech/language/hearing therapists, and resident rights experts. Supplemental personnel must have at least 1 year of experience in their fields;

(b) The curriculum of the nurse aide training program must include—
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(1) At least a total of 16 hours of training in the following areas prior to any direct contact with a resident:
   (i) Communication and interpersonal skills;
   (ii) Infection control;
   (iii) Safety/emergency procedures, including the Heimlich maneuver;
   (iv) Promoting residents’ independence; and
   (v) Respecting residents’ rights.

(2) Basic nursing skills:
   (i) Taking and recording vital signs;
   (ii) Measuring and recording height and weight;
   (iii) Caring for the residents’ environment;
   (iv) Recognizing abnormal changes in body functioning and the importance of reporting such changes to a supervisor; and
   (v) Caring for residents when death is imminent.

(3) Personal care skills, including, but not limited to—
   (i) Bathing;
   (ii) Grooming, including mouth care;
   (iii) Dressing;
   (iv) Toileting;
   (v) Assisting with eating and hydration;
   (vi) Proper feeding techniques;
   (vii) Skin care; and
   (viii) Transfers, positioning, and turning.

(4) Mental health and social service needs:
   (i) Modifying aide’s behavior in response to residents’ behavior;
   (ii) Awareness of developmental tasks associated with the aging process;
   (iii) How to respond to resident behavior;
   (iv) Allowing the resident to make personal choices, providing and reinforcing other behavior consistent with the resident’s dignity; and
   (v) Using the resident’s family as a source of emotional support.

(5) Care of cognitively impaired residents:
   (i) Techniques for addressing the unique needs and behaviors of individual with dementia (Alzheimer’s and others);
   (ii) Communicating with cognitively impaired residents;
   (iii) Understanding the behavior of cognitively impaired residents;
   (iv) Appropriate responses to the behavior of cognitively impaired residents; and
   (v) Methods of reducing the effects of cognitive impairments.

(6) Basic restorative services:
   (i) Training the resident in self care according to the resident’s abilities;
   (ii) Use of assistive devices in transferring, ambulation, eating, and dressing;
   (iii) Maintenance of range of motion;
   (iv) Proper turning and positioning in bed and chair;
   (v) Bowel and bladder training; and
   (vi) Care and use of prosthetic and orthotic devices.

(7) Residents’ Rights.
   (i) Providing privacy and maintenance of confidentiality;
   (ii) Promoting the residents’ right to make personal choices to accommodate their needs;
   (iii) Giving assistance in resolving grievances and disputes;
   (iv) Providing needed assistance in getting to and participating in resident and family groups and other activities;
   (v) Maintaining care and security of residents’ personal possessions;
   (vi) Promoting the resident’s right to be free from abuse, mistreatment, and neglect and the need to report any instances of such treatment to appropriate facility staff;
   (vii) Avoiding the need for restraints in accordance with current professional standards.

(c) Prohibition of charges. (1) No nurse aide who is employed by, or who has received an offer of employment from, a facility on the date on which the aide begins a nurse aide training and competency evaluation program may be charged for any portion of the program (including any fees for textbooks or other required course materials).

   (2) If an individual who is not employed, or does not have an offer to be employed, as a nurse aide becomes employed by, or receives an offer of employment from, a facility not later than 12 months after completing a nurse aide training and competency evaluation program, the State must provide for the reimbursement of costs.
§ 483.154 Nurse aide competency evaluation.

(a) Notification to Individual. The State must advise in advance any individual who takes the competency evaluation that a record of the successful completion of the evaluation will be included in the State’s nurse aide registry.

(b) Content of the competency evaluation program—(1) Written or oral examinations. The competency evaluation must—
   (i) Allow an aide to choose between a written and an oral examination;
   (ii) Address each course requirement specified in § 483.152(b);
   (iii) Be developed from a pool of test questions, only a portion of which is used in any one examination;
   (iv) Use a system that prevents disclosure of both the pool of questions and the individual competency evaluations; and
   (v) If oral, must be read from a prepared text in a neutral manner.

(2) Demonstration of skills. The skills demonstration must consist of a demonstration of randomly selected items drawn from a pool consisting of the tasks generally performed by nurse aides. This pool of skills must include all of the personal care skills listed in § 483.152(b)(3).

(c) Administration of the competency evaluation. (1) The competency examination must be administered and evaluated only by—
   (i) The State directly; or
   (ii) A State approved entity which is neither a skilled nursing facility that participates in Medicare nor a nursing facility that participates in Medicaid.

(2) No nurse aide who is employed by, or who has received an offer of employment from, a facility on the date on which the aide begins a nurse aide competency evaluation program may be charged for any portion of the program.

(d) Administration of the skills demonstration part of the evaluation must be—
   (i) Performed in a facility or laboratory setting comparable to the setting in which the individual will function as a nurse aide; and
   (ii) Administered and evaluated by a registered nurse with at least one year’s experience in providing care for the elderly or the chronically ill of any age.

(e) Facility proctoring of the competency evaluation.

(1) The competency evaluation may, at the nurse aide’s option, be conducted at the facility in which the nurse aide is or will be employed unless the facility is described in § 483.151(b)(2).

(2) The State may permit the competency evaluation to be proctored by facility personnel if the State finds that the procedure adopted by the facility assures that the competency evaluation program—
   (i) Is secure from tampering;
   (ii) Is standardized and scored by a testing, educational, or other organization approved by the State; and
   (iii) Requires no scoring by facility personnel.

(3) The State must retract the right to proctor nurse aide competency evaluations from facilities in which the State finds any evidence of impropriety, including evidence of tampering by facility staff.

(f) Successful completion of the competency evaluation program. (1) The State must establish a standard for satisfactory completion of the competency evaluation. To complete the competency evaluation successfully an individual must pass both the written or oral examination and the skills demonstration.

(2) A record of successful completion of the competency evaluation must be included in the nurse aide registry provided in § 483.156 within 30 days of the date if the individual is found to be competent.
§ 483.156  Registry of nurse aides.

(a) Establishment of registry. The State must establish and maintain a registry of nurse aides that meets the requirement of this section. The registry—

1. Must include as a minimum the information contained in paragraph (c) of this section:

2. Must be sufficiently accessible to meet the needs of the public and health care providers promptly;

3. May include home health aides who have successfully completed a home health aide competency evaluation program approved by the State if home health aides are differentiated from nurse aides;

4. Must provide that any response to an inquiry that includes a finding of abuse, neglect, or misappropriation of property includes the following information on any finding by the State survey agency of abuse, neglect, or misappropriation of property by the individual:
   (i) The individual’s full name.
   (ii) Information necessary to identify each individual;
   (iii) The date the individual became eligible for placement in the registry through successfully completing a nurse aide training and competency evaluation program or competency evaluation program or by meeting the requirements of § 483.150; and
   (iv) The following information on any finding by the State survey agency of abuse, neglect, or misappropriation of property by the individual:
      (A) Documentation of the State’s investigation, including the nature of the allegation and the evidence that led the State to conclude that the allegation was valid;
      (B) The date of the hearing, if the individual chose to have one, and its outcome; and
      (C) A statement by the individual disputing the allegation, if he or she chooses to make one; and
      (D) This information must be included in the registry within 10 working days of the finding and must remain in the registry permanently, unless the finding was made in error, the individual was found not guilty in a court of law, or the State is notified of the individual’s death.

(b) Registry operation. (1) The State may contract the daily operation and maintenance of the registry to a non-State entity. However, the State must maintain accountability for overall operation of the registry and compliance with these regulations.

(2) Only the State survey and certification agency may place on the registry findings of abuse, neglect, or misappropriation of property.

(3) The State must determine which individuals who (i) have successfully completed a nurse aide training and competency evaluation program or nurse aide competency evaluation program; (ii) have been deemed as meeting these requirements; or (iii) have had these requirements waived by the State do not qualify to remain on the registry because they have performed no nursing or nursing-related services for a period of 24 consecutive months.

(4) The State may not impose any charges related to registration on individuals listed in the registry.

(5) The State must provide information on the registry promptly.

(c) Registry Content. (1) The registry must contain at least the following information on each individual who has successfully completed a nurse aide training and competency evaluation program which meets the requirements of § 483.152 or a competency evaluation program which meets the requirements of § 483.154 and has been found by the State to be competent to function as a nurse aide or who may function as a nurse aide because of meeting criteria in § 483.150:
   (i) The individual’s full name.
   (ii) Information necessary to identify each individual;
   (iii) The date the individual became eligible for placement in the registry through successfully completing a nurse aide training and competency evaluation program or competency evaluation program or by meeting the requirements of § 483.150; and
   (iv) The following information on any finding by the State survey agency of abuse, neglect, or misappropriation of property by the individual:
      (A) Documentation of the State’s investigation, including the nature of the allegation and the evidence that led the State to conclude that the allegation was valid;
      (B) The date of the hearing, if the individual chose to have one, and its outcome; and
      (C) A statement by the individual disputing the allegation, if he or she chooses to make one; and
      (D) This information must be included in the registry within 10 working days of the finding and must remain in the registry permanently, unless the finding was made in error, the individual was found not guilty in a court of law, or the State is notified of the individual’s death.

(2) The registry must remove entries for individuals who have performed no nursing or nursing-related services for
a period of 24 consecutive months, unless the individual's registry entry includes documented findings of abuse, neglect, or misappropriation of property.

(d) Disclosure of information. The State must—
(1) Disclose all of the information in §483.156(c)(1)(iii) and (iv) to all requesters and may disclose additional information it deems necessary; and
(2) Promptly provide individuals with all information contained in the registry on them when adverse findings are placed on the registry and upon request. Individuals on the registry must have sufficient opportunity to correct any misstatements or inaccuracies contained in the registry.


§483.158 FFP for nurse aide training and competency evaluation.

(a) State expenditures for nurse aide training and competency evaluation programs and competency evaluation programs are administrative costs. They are matched as indicated in §483.15(b)(8) of this chapter.

(b) FFP is available for State expenditures associated with nurse aide training and competency evaluation programs and competency evaluation programs only for—
(1) Nurse aides employed by a facility;
(2) Nurse aides who have an offer of employment from a facility;
(3) Nurse aides who become employed by a facility not later than 12 months after completing a nurse aide training and competency evaluation program or competency evaluation program; or
(4) Nurse aides who receive an offer of employment from a facility not later than 12 months after completing a nurse aide training and competency evaluation program or competency evaluation program.

Subpart E—Appeals of Discharges, Transfers, and Preadmission Screening and Annual Resident Review (PASARR) Determinations

SOURCE: 57 FR 56534, Nov. 30, 1992, unless otherwise noted.

§483.200 Statutory basis.
This subpart is based on sections 1819(e)(3) and (f)(3) and 1919(e)(3) and (f)(3) of the Act, which require States to make available, to individuals who are discharged or transferred from SNFs or NFs, an appeals process that complies with guidelines issued by the Secretary.
[60 FR 50443, Sept. 29, 1995]

§483.202 Definitions.
For purposes of this subpart and subparts B and C—
Discharge means movement from an entity that participates in Medicare as a skilled nursing facility, a Medicare certified distinct part, an entity that participates in Medicaid as a nursing facility, or a Medicaid certified distinct part to a noninstitutional setting when the discharging facility ceases to be legally responsible for the care of the resident.
Individual means an individual or any legal representative of the individual.
Resident means a resident of a SNF or NF or any legal representative of the resident.
Transfer means movement from an entity that participates in Medicare as a skilled nursing facility, a Medicare certified distinct part, an entity that participates in Medicaid as a nursing facility or a Medicaid certified distinct part to another institutional setting when the legal responsibility for the care of the resident changes from the transferring facility to the receiving facility.
§ 483.204 Provision of a hearing and appeal system.

(a) Each State must provide a system for:
(1) A resident of a SNF or a NF to appeal a notice from the SNF or NF of intent to discharge or transfer the resident; and
(2) An individual who has been adversely affected by any PASARR determination made by the State in the context of either a preadmission screening or an annual resident review under subpart C of part 483 to appeal that determination.

(b) The State must provide an appeals system that meets the requirements of this subpart, § 483.12 of this part, and part 431 subpart E of this chapter.

[57 FR 56506, Nov. 30, 1992; 58 FR 25784, Apr. 28, 1993]

§ 483.206 Transfers, discharges and relocations subject to appeal.

(a) “Facility” means a certified entity, either a Medicare SNF or a Medicaid NF (see §§ 483.5 and 483.12(a)(1)).

(b) A resident has appeal rights when he or she is transferred from—
(1) A certified bed into a noncertified bed; and
(2) A bed in a certified entity to a bed in an entity which is certified as a different provider.

(c) A resident has no appeal rights when he or she is moved from one bed in the certified entity to another bed in the same certified entity.

Subpart F—Requirements That Must be Met by States and State Agencies, Resident Assessment

§ 483.315 Specification of resident assessment instrument.

(a) Statutory basis. Sections 1919(e)(5) and 1919(e)(5) of the Act require that a State specify the resident assessment instrument (RAI) to be used by long term care facilities in the State when conducting initial and periodic assessments of each resident’s functional capacity, in accordance with § 483.20.

(b) State options in specifying an RAI. The RAI that the State specifies must be one of the following:
(1) The instrument designated by HCFA.
(2) An alternate instrument specified by the State and approved by HCFA, using the criteria specified in the State Operations Manual issued by HCFA (HCFA Pub. 7), which is available for purchase through the National Technical Information Service, 5285 Port Royal Rd., Springfield, VA 22151.
(c) State requirements in specifying an RAI.
(1) Within 30 days after HCFA notifies the State of the HCFA-designated RAI or changes to it, the State must do one of the following:
(i) Specify the HCFA-designated RAI.
(ii) Notify HCFA of its intent to specify an alternate instrument.
(2) Within 60 days after receiving HCFA approval of an alternate RAI, the State must specify the RAI for use by all long term care facilities participating in the Medicare and Medicaid programs.
(3) After specifying an instrument, the State must provide periodic educational programs for facility staff to assist with implementation of the RAI.
(4) A State must audit implementation of the RAI through the survey process.
(5) A State must obtain approval from HCFA before making any modifications to its RAI.
(6) A State must adopt revisions to the RAI that are specified by HCFA.
(d) HCFA-designated RAI. The HCFA-designated RAI is published in the State Operations Manual issued by HCFA (HCFA Pub. 7), as updated periodically, and consists of the following:
(1) The minimum data set (MDS) and common definitions.
(2) The resident assessment protocols (RAPs) and triggers that are necessary to accurately assess residents, established by HCFA.
(3) The quarterly review, based on a subset of the MDS specified by HCFA.
(4) The requirements for use of the RAI that appear at § 483.20.
(e) Minimum data set (MDS). The MDS includes assessment in the following areas:
(1) Identification and demographic information, which includes information to identify the resident and facility, the resident’s residential history,
education, the reason for the assessment, guardianship status and information regarding advance directives, and information regarding mental health history.

(2) Customary routine, which includes the resident’s lifestyle prior to admission to the facility.

(3) Cognitive patterns, which include memory, decision making, consciousness, behavioral measures of delirium, and stability of condition.

(4) Communication, which includes scales for measuring hearing and communication skills, information on how the resident expresses himself or herself, and stability of communicative ability.

(5) Vision pattern, which includes a scale for measuring vision and vision problems.

(6) Mood and behavior patterns, which include scales for measuring behavioral indicators and symptoms, and stability of condition.

(7) Psychosocial well-being, which includes the resident’s interpersonal relationships and adjustment factors.

(8) Physical functioning and structural problems, which contains scales for measuring activities of daily living, mobility, potential for improvement, and stability of functioning.

(9) Continence, which includes assessment scales for bowel and bladder incontinence, continence patterns, interventions, and stability of continence status.

(10) Disease diagnoses and health conditions, which includes active medical diagnoses, physical problems, pain assessment, and stability of condition.

(11) Dental and nutritional status, which includes information on height and weight, nutritional problems and accommodations, oral care and problems, and measure of nutritional intake.

(12) Skin condition, which includes current and historical assessment of skin problems, treatments, and information regarding foot care.

(13) Activity pursuit, which gathers information on the resident’s activity preferences and the amount of time spent participating in activities.

(14) Medications, which contains information on the types and numbers of medications the resident receives.

(15) Special treatments and procedures, which includes measurements of therapies, assessment of rehabilitation/restorative care, special programs and interventions, and information on hospital visits and physician involvement.

(16) Discharge potential, which assesses the possibility of discharging the resident and discharge status.

(17) Documentation of summary information regarding the additional assessment performed through the resident assessment protocols.

(18) Documentation of participation in assessment.

(f) Resident assessment protocols (RAPs). At a minimum, the RAPs address the following domains:

(1) Delirium.
(2) Cognitive loss.
(3) Visual function.
(4) Communication.
(5) ADL functional/rehabilitation potential.
(6) Urinary incontinence and indwelling catheter.
(7) Psychosocial well-being.
(8) Mood state.
(9) Behavioral symptoms.
(10) Activities.
(11) Falls.
(12) Nutritional status.
(13) Feeding tubes.
(14) Dehydration/fluid maintenance.
(15) Dental care.
(16) Pressure ulcers.
(17) Psychotropic drug use.
(18) Physical restraints.

(g) Criteria for HCFA approval of alternate instrument. To receive HCFA approval, a State’s alternate instrument must use the standardized format, organization, item labels and definitions, and instructions specified by HCFA in the latest issuance of the State Operations Manual issued by HCFA (HCFA Pub. 7).

(h) State MDS collection and data base requirements. (1) As part of facility survey responsibilities, the State must establish and maintain an MDS Database, and must do the following:

(i) Use a system to collect, store, and analyze data that is developed or approved by HCFA.

(ii) Obtain HCFA approval before modifying any parts of the HCFA.
§ 483.400 Basis and purpose.

This subpart implements section 1905 (c) and (d) of the Act which gives the Secretary authority to prescribe regulations for intermediate care facility services in facilities for the mentally retarded or persons with related conditions.

§ 483.405 Relationship to other HHS regulations.

In addition to compliance with the regulations set forth in this subpart, facilities are obliged to meet the applicable provisions of other HHS regulations, including but not limited to those pertaining to nondiscrimination on the basis of race, color, or national origin (45 CFR Part 80), nondiscrimination on the basis of handicap (45 CFR Part 84), nondiscrimination on the basis of age (45 CFR Part 91), protection of human subjects of research (45 CFR Part 46), and fraud and abuse (42 CFR Part 455). Although those regulations are not in themselves considered conditions of participation under this Part, their violation may result in the termination or suspension of, or the refusal to grant or continue, Federal financial assistance.

§ 483.410 Condition of participation: Governing body and management.

(a) Standard: Governing body. The facility must identify an individual or

(j) Resident-identifiable data. (1) The State may not release information that is resident-identifiable to the public.

(2) The State may not release RAI data that is resident-identifiable except in accordance with a written agreement under which the recipient agrees to be bound by the restrictions described in paragraph (i) of this section.


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individuals to constitute the governing body of the facility. The governing body must—

(1) Exercise general policy, budget, and operating direction over the facility;

(2) Set the qualifications (in addition to those already set by State law, if any) for the administrator of the facility; and

(3) Appoint the administrator of the facility.

(b) Standard: Compliance with Federal, State, and local laws. The facility must be in compliance with all applicable provisions of Federal, State and local laws, regulations and codes pertaining to health, safety, and sanitation.

(c) Standard: Client records. (1) The facility must develop and maintain a recordkeeping system that includes a separate record for each client and that documents the client’s health care, active treatment, social information, and protection of the client’s rights.

(2) The facility must keep confidential all information contained in the clients’ records, regardless of the form or storage method of the records.

(3) The facility must develop and implement policies and procedures governing the release of any client information, including consents necessary from the client, or parents (if the client is a minor) or legal guardian.

(4) Any individual who makes an entry in a client’s record must make it legibly, date it, and sign it.

(5) The facility must provide a legend to explain any symbol or abbreviation used in a client’s record.

(6) The facility must provide each identified residential living unit with appropriate aspects of each client’s record.

(d) Standard: Services provided under agreements with outside sources. (1) If a service required under this subpart is not provided directly, the facility must have a written agreement with an outside program, resource, or service to furnish the necessary service, including emergency and other health care.

(2) The agreement must—

(i) Contain the responsibilities, functions, objectives, and other terms agreed to by both parties; and

(ii) Provide that the facility is responsible for assuring that the outside services meet the standards for quality of services contained in this subpart.

(3) The facility must assure that outside services meet the needs of each client.

(4) If living quarters are not provided in a facility owned by the ICF/MR, the ICF/MR remains directly responsible for the standards relating to physical environment that are specified in §483.470 (a) through (g), (j) and (k).

(e) Standard: Licensure. The facility must be licensed under applicable State and local law.


§ 483.420 Condition of participation: Client protections.

(a) Standard: Protection of clients’ rights. The facility must ensure the rights of all clients. Therefore, the facility must—

(1) Inform each client, parent (if the client is a minor), or legal guardian, of the client’s rights and the rules of the facility;

(2) Inform each client, parent (if the client is a minor), or legal guardian, of the client’s medical condition, developmental and behavioral status, attendant risks of treatment, and of the right to refuse treatment;

(3) Allow and encourage individual clients to exercise their rights as clients of the facility, and as citizens of the United States, including the right to file complaints, and the right to due process;

(4) Allow individual clients to manage their financial affairs and teach them to do so to the extent of their capabilities;

(5) Ensure that clients are not subjected to physical, verbal, sexual or psychological abuse or punishment;

(6) Ensure that clients are free from unnecessary drugs and physical restraints and are provided active treatment to reduce dependency on drugs and physical restraints;

(7) Provide each client with the opportunity for personal privacy and ensure privacy during treatment and care of personal needs;

(8) Ensure that clients are not compelled to perform services for the facility and ensure that clients who do
work for the facility are compensated for their efforts at prevailing wages and commensurate with their abilities;

(9) Ensure clients the opportunity to communicate, associate and meet privately with individuals of their choice, and to send and receive unopened mail;

(10) Ensure that clients have access to telephones with privacy for incoming and outgoing local and long distance calls except as contraindicated by factors identified within their individual program plans;

(11) Ensure clients the opportunity to participate in social, religious, and community group activities;

(12) Ensure that clients have the right to retain and use appropriate personal possessions and clothing, and ensure that each client is dressed in his or her own clothing each day; and

(13) Permit a husband and wife who both reside in the facility to share a room.

(b) Standard: Client finances.

(1) The facility must establish and maintain a system that—

(i) Assures a full and complete accounting of clients' personal funds entrusted to the facility on behalf of clients; and

(ii) Precludes any commingling of client funds with facility funds or with the funds of any person other than another client.

(2) The client's financial record must be available on request to the client, parents (if the client is a minor) or legal guardian.

(c) Standard: Communication with clients, parents, and guardians.

The facility must—

(1) Promote participation of parents (if the client is a minor) and legal guardians in the process of providing active treatment to a client unless their participation is unobtainable or inappropriate;

(2) Answer communications from clients' families and friends promptly and appropriately;

(3) Promote visits by individuals with a relationship to the client (such as family, close friends, legal guardians and advocates) at any reasonable hour, without prior notice, consistent with the right of that client's and other clients' privacy, unless the interdisciplinary team determines that the visit would not be appropriate;

(4) Promote visits by parents or guardians to any area of the facility that provides direct client care services to the client, consistent with the right of that client's and other clients' privacy;

(5) Promote frequent and informal leaves from the facility for visits, trips, or vacations; and

(6) Notify promptly the client's parents or guardian of any significant incidents, or changes in the client's condition including, but not limited to, serious illness, accident, death, abuse, or unauthorized absence.

(d) Standard: Staff treatment of clients.

(1) The facility must develop and implement written policies and procedures that prohibit mistreatment, neglect or abuse of the client.

(i) Staff of the facility must not use physical, verbal, sexual or psychological abuse or punishment.

(ii) Staff must not punish a client by withholding food or hydration that contributes to a nutritionally adequate diet.

(iii) The facility must prohibit the employment of individuals with a conviction or prior employment history of child or client abuse, neglect or mistreatment.

(2) The facility must ensure that all allegations of mistreatment, neglect or abuse, as well as injuries of unknown source, are reported immediately to the administrator or to other officials in accordance with State law through established procedures.

(3) The facility must have evidence that all alleged violations are thoroughly investigated and must prevent further potential abuse while the investigation is in progress.

(4) The results of all investigations must be reported to the administrator or designated representative or to other officials in accordance with State law within five working days of the incident and, if the alleged violation is verified, appropriate corrective action must be taken.

§ 483.430 Condition of participation: Facility staffing.

(a) Standard: Qualified mental retardation professional. Each client's active
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A treatment program must be integrated, coordinated and monitored by a qualified mental retardation professional who—

(1) Has at least one year of experience working directly with persons with mental retardation or other developmental disabilities; and

(2) Is one of the following:

(i) A doctor of medicine or osteopathy.

(ii) A registered nurse.

(iii) A social worker who holds at least a bachelor’s degree in a professional category specified in paragraph (b)(5) of this section.

(b) Standard: Professional program services.

(1) Each client must receive the professional program services needed to implement the active treatment program defined by each client’s individual program plan. Professional program staff must work directly with clients and with paraprofessional, nonprofessional and other professional program staff who work with clients.

(2) The facility must have available enough qualified professional staff to carry out and monitor the various professional interventions in accordance with the stated goals and objectives of every individual program plan.

(3) Professional program staff must participate as members of the interdisciplinary team in relevant aspects of the active treatment process.

(4) Professional program staff must participate in ongoing staff development and training in both formal and informal settings with other professional, paraprofessional, and nonprofessional staff members.

(5) Professional program staff must be licensed, certified, or registered, as applicable, to provide professional services by the State in which he or she practices. Those professional program staff who do not fall under the jurisdiction of State licensure, certification, or registration requirements, specified in §483.410(b), must meet the following qualifications:

(i) To be designated as an occupational therapist, an individual must be eligible for certification as an occupational therapist by the American Occupational Therapy Association or another comparable body.

(ii) To be designated as a physical therapist, an individual must be eligible for certification as a physical therapist by the American Physical Therapy Association or another comparable body.

(iii) To be designated as a social worker, an individual must—

(A) Hold a graduate degree from a school of social work accredited or approved by the Council on Social Work Education or another comparable body; or

(B) Hold a Bachelor of Social Work degree from a college or university accredited or approved by the Council on Social Work Education or another comparable body.

(iv) To be designated as a speech-language pathologist or audiologist, an individual must—

(A) Be eligible for a Certificate of Clinical Competence in Speech-Language Pathology or Audiology granted by the American Speech-Language-Hearing Association or another comparable body; or

(B) Meet the educational requirements for certification and be in the process of accumulating the supervised experience required for certification.

(v) To be designated as a professional recreation staff member, an individual must have a bachelor’s degree in recreation or in a specialty area such as art, dance, music or physical education.

(ix) To be designated as a professional dietitian, an individual must be
eligible for registration by the American Dietetics Association.

(x) To be designated as a human services professional an individual must have at least a bachelor’s degree in a human services field (including, but not limited to: sociology, special education, rehabilitation counseling, and psychology).

(xi) If the client’s individual program plan is being successfully implemented by facility staff, professional program staff meeting the qualifications of paragraph (b)(5) (i) through (x) of this section are not required—

(A) Except for qualified mental retardation professionals;

(B) Except for the requirements of paragraph (b)(2) of this section concerning the facility’s provision of enough qualified professional program staff; and

(C) Unless otherwise specified by State licensure and certification requirements.

(c) Standard: Facility staffing. (1) The facility must not depend upon clients or volunteers to perform direct care services for the facility.

(2) There must be responsible direct care staff on duty and awake on a 24-hour basis, when clients are present, to take prompt, appropriate action in case of injury, illness, fire or other emergency, in each defined residential living unit housing—

(i) Clients for whom a physician has ordered a medical care plan;

(ii) Clients who are aggressive, assaultive or security risks;

(iii) More than 16 clients; or

(iv) Fewer than 16 clients within a multi-unit building.

(3) There must be a responsible direct care staff person on duty on a 24 hour basis (when clients are present) to respond to injuries and symptoms of illness, and to handle emergencies, in each defined residential living unit housing—

(i) Clients for whom a physician has not ordered a medical care plan;

(ii) Clients who are not aggressive, assaultive or security risks; and

(iii) Sixteen or fewer clients.

(4) The facility must provide sufficient support staff so that direct care staff are not required to perform support services to the extent that these duties interfere with the exercise of their primary direct client care duties.

(d) Standard: Direct care (residential living unit) staff. (1) The facility must provide sufficient direct care staff to manage and supervise clients in accordance with their individual program plans.

(2) Direct care staff are defined as the present on-duty staff calculated over all shifts in a 24-hour period for each defined residential living unit.

(3) Direct care staff must be provided by the facility in the following minimum ratios of direct care staff to clients:

(i) For each defined residential living unit serving children under the age of 12, severely and profoundly retarded clients, clients with severe physical disabilities, or clients who are aggressive, assaultive, or security risks, or who manifest severely hyperactive or psychotic-like behavior, the staff to client ratio is 1 to 3.2.

(ii) For each defined residential living unit serving moderately retarded clients, the staff to client ratio is 1 to 4.

(iii) For each defined residential living unit serving moderately retarded clients, the staff to client ratio is 1 to 6.4.

(4) When there are no clients present in the living unit, a responsible staff member must be available by telephone.

(e) Standard: Staff training program. (1) The facility must provide each employee with initial and continuing training that enables the employee to perform his or her duties effectively, efficiently, and competently.

(2) For employees who work with clients, training must focus on skills and competencies directed toward clients’ developmental, behavioral, and health needs.

(3) Staff must be able to demonstrate the skills and techniques necessary to administer interventions to manage the inappropriate behavior of clients.

(4) Staff must be able to demonstrate the skills and techniques necessary to implement the individual program plans for each client for whom they are responsible.
§ 483.440 Condition of participation: Active treatment services.

(a) Standard: Active treatment. (1) Each client must receive a continuous active treatment program, which includes aggressive, consistent implementation of a program of specialized and generic training, treatment, health services and related services described in this subpart, that is directed toward—

(i) The acquisition of the behaviors necessary for the client to function with as much self determination and independence as possible; and

(ii) The prevention or deceleration of regression or loss of current optimal functional status.

(2) Active treatment does not include services to maintain generally independent clients who are able to function with little supervision or in the absence of a continuous active treatment program.

(b) Standard: Admissions, transfers, and discharge. (1) Clients who are admitted by the facility must be in need of and receiving active treatment services.

(2) Admission decisions must be based on a preliminary evaluation of the client that is conducted or updated by the facility or by outside sources.

(3) A preliminary evaluation must contain background information as well as currently valid assessments of functional developmental, behavioral, social, health and nutritional status to determine if the facility can provide for the client’s needs and if the client is likely to benefit from placement in the facility.

(4) If a client is to be either transferred or discharged, the facility must—

(i) Have documentation in the client’s record that the client was transferred or discharged for good cause; and

(ii) Provide a reasonable time to prepare the client and his or her parents or guardian for the transfer or discharge (except in emergencies).

(5) At the time of the discharge, the facility must—

(i) Develop a final summary of the client’s developmental, behavioral, social, health and nutritional status and, with the consent of the client, parents (if the client is a minor) or legal guardian, provide a copy to authorized persons and agencies; and

(ii) Provide a post-discharge plan of care that will assist the client to adjust to the new living environment.

(c) Standard: Individual program plan. (1) Each client must have an individual program plan developed by an interdisciplinary team that represents the professions, disciplines or service areas that are relevant to—

(i) Identifying the client’s needs, as described by the comprehensive functional assessments required in paragraph (c)(3) of this section; and

(ii) Designing programs that meet the client’s needs.

(2) Appropriate facility staff must participate in interdisciplinary team meetings. Participation by other agencies serving the client is encouraged. Participation by the client, his or her parent (if the client is a minor), or the client’s legal guardian is required unless that participation is unobtainable or inappropriate.

(3) Within 30 days after admission, the interdisciplinary team must perform accurate assessments or reassessments as needed to supplement the preliminary evaluation conducted prior to admission. The comprehensive functional assessment must take into consideration the client’s age (for example, child, young adult, elderly person) and the implications for active treatment at each stage, as applicable, and must—

(i) Identify the presenting problems and disabilities and where possible, their causes;

(ii) Identify the client’s specific developmental strengths;

(iii) Identify the client’s specific developmental and behavioral management needs;

(iv) Identify the client’s need for services without regard to the actual availability of the services needed; and

(v) Include physical development and health, sensorimotor development, affective development, speech and language development and auditory functioning, cognitive development, social development, adaptive behaviors or independent living skills necessary for the client to be able to function in the...
community, and as applicable, vocational skills.

(4) Within 30 days after admission, the interdisciplinary team must prepare for each client an individual program plan that states the specific objectives necessary to meet the client’s needs, as identified by the comprehensive assessment required by paragraph (c)(3) of this section, and the planned sequence for dealing with those objectives. These objectives must—

(i) Be stated separately, in terms of a single behavioral outcome;
(ii) Be assigned projected completion dates;
(iii) Be expressed in behavioral terms that provide measurable indices of performance;
(iv) Be organized to reflect a developmental progression appropriate to the individual; and
(v) Be assigned priorities.

(5) Each written training program designed to implement the objectives in the individual program plan must specify:

(i) The methods to be used;
(ii) The schedule for use of the method;
(iii) The person responsible for the program;
(iv) The type of data and frequency of data collection necessary to be able to assess progress toward the desired objectives;
(v) The inappropriate client behavior(s), if applicable; and
(vi) Provision for the appropriate expression of behavior and the replacement of inappropriate behavior, if applicable, with behavior that is adaptive or appropriate.

(6) The individual program plan must also:

(i) Describe relevant interventions to support the individual toward independence.
(ii) Identify the location where program strategy information (which must be accessible to any person responsible for implementation) can be found.
(iii) Include, for those clients who lack them, training in personal skills essential for privacy and independence (including, but not limited to, toilet training, personal hygiene, dental hygiene, self-feeding, bathing, dressing, grooming, and communication of basic needs), until it has been demonstrated that the client is developmentally incapable of acquiring them.
(iv) Identify mechanical supports, if needed, to achieve proper body position, balance, or alignment. The plan must specify the reasons for each support, the situations in which each is to be applied, and a schedule for the use of each support.
(v) Provide that clients who have multiple disabling conditions spend a major portion of each waking day out of bed and outside the bedroom area, moving about by various methods and devices whenever possible.
(vi) Include opportunities for client choice and self-management.

(7) A copy of each client’s individual program plan must be made available to all relevant staff, including staff of other agencies who work with the client, and to the client, parents (if the client is a minor) or legal guardian.

(d) Standard: Program implementation.

(1) As soon as the interdisciplinary team has formulated a client’s individual program plan, each client must receive a continuous active treatment program consisting of needed interventions and services in sufficient number and frequency to support the achievement of the objectives identified in the individual program plan.

(2) The facility must develop an active treatment schedule that outlines the current active treatment program and that is readily available for review by relevant staff.

(3) Except for those facets of the individual program plan that must be implemented only by licensed personnel, each client’s individual program plan must be implemented by all staff who work with the client, including professional, paraprofessional and nonprofessional staff.

(e) Standard: Program documentation.

(1) Data relative to accomplishment of the criteria specified in client individual program plan objectives must be documented in measurable terms.

(2) The facility must document significant events that are related to the client’s individual program plan and assessments and that contribute to an overall understanding of the client’s
ongoing level and quality of functioning.

(f) Standard: Program monitoring and change. (1) The individual program plan must be reviewed at least by the qualified mental retardation professional and revised as necessary, including, but not limited to situations in which the client—

(i) Has successfully completed an objective or objectives identified in the individual program plan;
(ii) Is regressing or losing skills already gained;
(iii) Is failing to progress toward identified objectives after reasonable efforts have been made; or
(iv) Is being considered for training towards new objectives.

(2) At least annually, the comprehensive functional assessment of each client must be reviewed by the interdisciplinary team for relevancy and updated as needed, and the individual program plan must be revised, as appropriate, repeating the process set forth in paragraph (c) of this section.

(3) The facility must designate and use a specially constituted committee or committees consisting of members of facility staff, parents, legal guardians, clients (as appropriate), qualified persons who have either experience or training in contemporary practices to change inappropriate client behavior, and persons with no ownership or controlling interest in the facility to—

(i) Review, approve, and monitor individual programs designed to manage inappropriate behavior and other programs that, in the opinion of the committee, involve risks to client protection and rights;

(ii) Insure that these programs are conducted only with the written informed consent of the client, parent (if the client is a minor), or legal guardian; and

(iii) Review, monitor and make suggestions to the facility about its practices and programs as they relate to drug usage, physical restraints, timeout rooms, application of painful or noxious stimuli, control of inappropriate behavior, protection of client rights and funds, and any other area that the committee believes need to be addressed.

(4) The provisions of paragraph (f)(3) of this section may be modified only if, in the judgment of the State survey agency, Court decrees, State law or regulations provide for equivalent client protection and consultation.

§ 483.450 Condition of participation: Client behavior and facility practices.

(a) Standard: Facility practices—Conduct toward clients. (1) The facility must develop and implement written policies and procedures for the management of conduct between staff and clients. These policies and procedures must—

(i) Promote the growth, development and independence of the client;
(ii) Address the extent to which client choice will be accommodated in daily decision-making, emphasizing self-determination and self-management, to the extent possible;
(iii) Specify client conduct to be allowed or not allowed; and
(iv) Be available to all staff, clients, parents of minor children, and legal guardians.

(2) To the extent possible, clients must participate in the formulation of these policies and procedures.

(3) Clients must not discipline other clients, except as part of an organized system of self-government, as set forth in facility policy.

(b) Standard: Management of inappropriate client behavior. (1) The facility must develop and implement written policies and procedures that govern the management of inappropriate client behavior. These policies and procedures must be consistent with the provisions of paragraph (a) of this section. These procedures must—

(i) Specify all facility approved interventions to manage inappropriate client behavior;
(ii) Designate these interventions on a hierarchy to be implemented, ranging from most positive or least intrusive, to least positive or most intrusive;
(iii) Insure, prior to the use of more restrictive techniques, that the client's record documents that programs incorporating the use of less intrusive or more positive techniques have been tried systematically and demonstrated to be ineffective; and
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(iv) Address the following:
(A) The use of time-out rooms.
(B) The use of physical restraints.
(C) The use of drugs to manage inappropriate behavior.
(D) The application of painful or noxious stimuli.
(E) The staff members who may authorize the use of specified interventions.
(F) A mechanism for monitoring and controlling the use of such interventions.
(2) Interventions to manage inappropriate client behavior must be employed with sufficient safeguards and supervision to ensure that the safety, welfare and civil and human rights of clients are adequately protected.
(3) Techniques to manage inappropriate client behavior must never be used for disciplinary purposes, for the convenience of staff or as a substitute for an active treatment program.
(4) The use of systematic interventions to manage inappropriate client behavior must be incorporated into the client’s individual program plan, in accordance with 483.440(c) (4) and (5) of this subpart.
(5) Standing or as needed programs to control inappropriate behavior are not permitted.
(c) Standard: Time-out rooms. (1) A client may be placed in a room from which egress is prevented only if the following conditions are met:
   (i) The placement is a part of an approved systematic time-out program as required by paragraph (b) of this section. (Thus, emergency placement of a client into a time-out room is not allowed.)
   (ii) The client is under the direct constant visual supervision of designated staff.
   (iii) The door to the room is held shut by staff or by a mechanism requiring constant physical pressure from a staff member to keep the mechanism engaged.
(2) Placement of a client in a time-out room must not exceed one hour.
(3) Clients placed in time-out rooms must be protected from hazardous conditions including, but not limited to, presence of sharp corners and objects, uncovered light fixtures, unprotected electrical outlets.
(4) A record of time-out activities must be kept.
(d) Standard: Physical restraints. (1) The facility may employ physical restraint only—
   (i) As an integral part of an individual program plan that is intended to lead to less restrictive means of managing and eliminating the behavior for which the restraint is applied;
   (ii) As an emergency measure, but only if absolutely necessary to protect the client or others from injury; or
   (iii) As a health-related protection prescribed by a physician, but only if absolutely necessary during the conduct of a specific medical or surgical procedure, or only if absolutely necessary for client protection during the time that a medical condition exists.
(2) Authorizations to use or extend restraints as an emergency must be:
   (i) In effect no longer than 12 consecutive hours; and
   (ii) Obtained as soon as the client is restrained or stable.
(3) The facility must not issue orders for restraint on a standing or as needed basis.
(4) A client placed in restraint must be checked at least every 30 minutes by staff trained in the use of restraints, released from the restraint as quickly as possible, and a record of these checks and usage must be kept.
(5) Restraints must be designed and used so as not to cause physical injury to the client and so as to cause the least possible discomfort.
(6) Opportunity for motion and exercise must be provided for a period of not less than 10 minutes during each two hour period in which restraint is employed, and a record of such activity must be kept.
(7) Barred enclosures must not be more than three feet in height and must not have tops.
(e) Standard: Drug usage. (1) The facility must not use drugs in doses that interfere with the individual client’s daily living activities.
(2) Drugs used for control of inappropriate behavior must be approved by the interdisciplinary team and be used only as an integral part of the client’s individual program plan that is directed specifically towards the reduction of and eventual elimination of the

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behaviors for which the drugs are employed.

(3) Drugs used for control of inappropriate behavior must not be used until it can be justified that the harmful effects of the behavior clearly outweigh the potentially harmful effects of the drugs.

(4) Drugs used for control of inappropriate behavior must be—

(i) Monitored closely, in conjunction with the physician and the drug regimen review requirement at §483.460(j), for desired responses and adverse consequences by facility staff; and

(ii) Gradually withdrawn at least annually in a carefully monitored program conducted in conjunction with the interdisciplinary team, unless clinical evidence justifies that this is contraindicated.

§ 483.460 Condition of participation: Health care services.

(a) Standard: Physician services.

(1) The facility must ensure the availability of physician services 24 hours a day.

(2) The physician must develop, in coordination with licensed nursing personnel, a medical care plan of treatment for a client if the physician determines that an individual client requires 24-hour licensed nursing care. This plan must be integrated in the individual program plan.

(3) The facility must provide or obtain preventive and general medical care as well as annual physical examinations of each client that at a minimum include the following:

(i) Evaluation of vision and hearing.

(ii) Immunizations, using as a guide the recommendations of the Public Health Service Advisory Committee on Immunization Practices or of the Committee on the Control of Infectious Diseases of the American Academy of Pediatrics.

(iii) Routine screening laboratory examinations as determined necessary by the physician, and special studies when needed.

(iv) Tuberculosis control, appropriate to the facility’s population, and in accordance with the recommendations of the American College of Chest Physicians or the section of diseases of the chest of the American Academy of Pediatrics, or both.

(4) To the extent permitted by State law, the facility may utilize physician assistants and nurse practitioners to provide physician services as described in this section.

(b) Standard: Physician participation in the individual program plan. A physician must participate in—

(1) The establishment of each newly admitted client’s initial individual program plan as required by §456.380 of this chapter that specified plan of care requirements for ICFs; and

(2) If appropriate, physicians must participate in the review and update of an individual program plan as part of the interdisciplinary team process either in person or through written report to the interdisciplinary team.

(c) Standard: Nursing services. The facility must provide clients with nursing services in accordance with their needs. These services must include—

(1) Participation as appropriate in the development, review, and update of an individual program plan as part of the interdisciplinary team process;

(2) The development, with a physician, of a medical care plan of treatment for a client when the physician has determined that an individual client requires such a plan;

(3) For those clients certified as not needing a medical care plan, a review of their health status which must—

(i) Be by a direct physical examination;

(ii) Be by a licensed nurse;

(iii) Be on a quarterly or more frequent basis depending on client need;

(iv) Be recorded in the client’s record; and

(v) Result in any necessary action (including referral to a physician to address client health problems).

(4) Other nursing care as prescribed by the physician or as identified by client needs; and

(5) Implementing, with other members of the interdisciplinary team, appropriate protective and preventive health measures that include, but are not limited to—

(i) Training clients and staff as needed in appropriate health and hygiene methods;
(ii) Control of communicable diseases and infections, including the instruction of other personnel in methods of infection control; and

(iii) Training direct care staff in detecting signs and symptoms of illness or dysfunction, first aid for accidents or illness, and basic skills required to meet the health needs of the clients.

(d) Standard: Nursing staff. (1) Nurses providing services in the facility must have a current license to practice in the State.

(2) The facility must employ or arrange for licensed nursing services sufficient to care for clients health needs including those clients with medical care plans.

(3) The facility must utilize registered nurses as appropriate and required by State law to perform the health services specified in this section.

(4) If the facility utilizes only licensed practical or vocational nurses to provide health services, it must have a formal arrangement with a registered nurse to be available for verbal or on-site consultation to the licensed practical or vocational nurse.

(5) Non-licensed nursing personnel who work with clients under a medical care plan must do so under the supervision of licensed persons.

(e) Standard: Dental services. (1) The facility must provide or make arrangements for comprehensive diagnostic and treatment services for each client from qualified personnel, including licensed dentists and dental hygienists either through organized dental services in-house or through arrangement.

(2) If appropriate, dental professionals must participate, in the development, review and update of an individual program plan as part of the interdisciplinary process either in person or through written report to the interdisciplinary team.

(3) The facility must provide education and training in the maintenance of oral health.

(f) Standard: Comprehensive dental diagnostic services. Comprehensive dental diagnostic services include—

(1) A complete extraoral and intraoral examination, using all diagnostic aids necessary to properly evaluate the client's oral condition, not later than one month after admission to the facility (unless the examination was completed within twelve months before admission);

(2) Periodic examination and diagnosis performed at least annually, including radiographs when indicated and detection of manifestations of systemic disease; and

(3) A review of the results of examination and entry of the results in the client's dental record.

(g) Standard: Comprehensive dental treatment. The facility must ensure comprehensive dental treatment services that include—

(1) The availability for emergency dental treatment on a 24-hour-a-day basis by a licensed dentist; and

(2) Dental care needed for relief of pain and infections, restoration of teeth, and maintenance of dental health.

(h) Standard: Documentation of dental services. (1) If the facility maintains an in-house dental service, the facility must keep a permanent dental record for each client, with a dental summary maintained in the client's living unit.

(2) If the facility does not maintain an in-house dental service, the facility must obtain a dental summary of the results of dental visits and maintain the summary in the client's living unit.

(i) Standard: Pharmacy services. The facility must provide or make arrangements for the provision of routine and emergency drugs and biologicals to its clients. Drugs and biologicals may be obtained from community or contract pharmacists or the facility may maintain a licensed pharmacy.

(j) Standard: Drug regimen review. (1) A pharmacist with input from the interdisciplinary team must review the drug regimen of each client at least quarterly.

(2) The pharmacist must report any irregularities in clients' drug regimens to the prescribing physician and interdisciplinary team.

(3) The pharmacist must prepare a record of each client's drug regimen reviews and the facility must maintain that record.

(4) An individual medication administration record must be maintained for each client.
(5) As appropriate the pharmacist must participate in the development, implementation, and review of each client's individual program plan either in person or through written report to the interdisciplinary team.

(k) Standard: Drug administration. The facility must have an organized system for drug administration that identifies each drug up to the point of administration. The system must assure that—
(1) All drugs are administered in compliance with the physician's orders;
(2) All drugs, including those that are self-administered, are administered without error;
(3) Unlicensed personnel are allowed to administer drugs only if State law permits;
(4) Clients are taught how to administer their own medications if the interdisciplinary team determines that self administration of medications is an appropriate objective, and if the physician does not specify otherwise;
(5) The client's physician is informed of the interdisciplinary team's decision that self-administration of medications is an objective for the client;
(6) No client self-administers medications until he or she demonstrates the competency to do so;
(7) Drugs used by clients while not under the direct care of the facility are packaged and labeled in accordance with State law; and
(8) Drug administration errors and adverse drug reactions are recorded and reported immediately to a physician.

(l) Standard: Drug storage and record-keeping. (1) The facility must store drugs under proper conditions of sanitation, temperature, light, humidity, and security.
(2) The facility must keep all drugs and biologicals locked except when being prepared for administration.
(3) The facility must maintain records of the receipt and disposition of all controlled drugs in schedules II through IV (drugs subject to the Comprehensive Drug Abuse Prevention and Control Act of 1970, 21 U.S.C. 801 et seq., as implemented by 21 CFR part 308).
(5) If the facility maintains a licensed pharmacy, the facility must comply with the regulations for controlled drugs.

(m) Standard: Drug labeling. (1) Labeling of drugs and biologicals must—
(i) Be based on currently accepted professional principles and practices; and
(ii) Include the appropriate accessory and cautionary instructions, as well as the expiration date, if applicable.
(2) The facility must remove from use—
(i) Outdated drugs; and
(ii) Drug containers with worn, illegible, or missing labels.
(3) Drugs and biologicals packaged in containers designated for a particular client must be immediately removed from the client's current medication supply if discontinued by the physician.

(n) Standard: Laboratory services. (1) If a facility chooses to provide laboratory services, the laboratory must meet the requirements specified in part 493 of this chapter.
(2) If the laboratory chooses to refer specimens for testing to another laboratory, the referral laboratory must be certified in the appropriate specialties and subspecialities of service in accordance with the requirements of part 493 of this chapter.

§ 483.470 Condition of participation: Physical environment.

(a) Standard: Client living environment.
(1) The facility must not house clients of grossly different ages, developmental levels, and social needs in close physical or social proximity unless the housing is planned to promote the growth and development of all those housed together.
(2) The facility must not segregate clients solely on the basis of their physical disabilities. It must integrate clients who have ambulation deficits or
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who are deaf, blind, or have seizure disorders, etc., with others of comparable social and intellectual development.

(b) Standard: Client bedrooms. (1) Bedrooms must—
   (i) Be rooms that have at least one outside wall;
   (ii) Be equipped with or located near toilet and bathing facilities;
   (iii) Accommodate no more than four clients unless granted a variance under paragraph (b)(3) of this section;
   (iv) Measure at least 60 square feet per client in multiple client bedrooms and at least 80 square feet in single client bedrooms; and
   (v) In all facilities initially certified, or in buildings constructed or with major renovations or conversions on or after October 3, 1988, have walls that extend from floor to ceiling.

(2) If a bedroom is below grade level, it must have a window that—
   (i) Is usable as a second means of escape by the client(s) occupying the room; and
   (ii) Is no more than 44 inches (measured to the window sill) above the floor unless the facility is surveyed under the Health Care Occupancy Chapter of the Life Safety Code, in which case the window must be no more than 36 inches (measured to the window sill) above the floor.

(3) The survey agency may grant a variance from the limit of four clients per room only if a physician who is a member of the interdisciplinary team and who is a qualified mental retardation professional—
   (i) Certifies that each client to be placed in a bedroom housing more than four persons is so severely medically impaired as to require direct and continuous monitoring during sleeping hours; and
   (ii) Documents the reasons why housing in a room of only four or fewer persons would not be medically feasible.

(4) The facility must provide each client with—
   (i) A separate bed of proper size and height for the convenience of the client;
   (ii) A clean, comfortable, mattress;
   (iii) Bedding appropriate to the weather and climate; and
   (iv) Functional furniture appropriate to the client's needs, and individual closet space in the client's bedroom with clothes racks and shelves accessible to the client.

(c) Standard: Storage space in bedroom. The facility must provide—
   (1) Space and equipment for daily out-of-bed activity for all clients who are not yet mobile, except those who have a short-term illness or those few clients for whom out-of-bed activity is a threat to health and safety; and
   (2) Suitable storage space, accessible to clients, for personal possessions, such as TVs, radios, prosthetic equipment and clothing.

(d) Standard: Client bathrooms. The facility must—
   (1) Provide toilet and bathing facilities appropriate in number, size, and design to meet the needs of the clients;
   (2) Provide for individual privacy in toilets, bathtubs, and showers; and
   (3) In areas of the facility where clients who have not been trained to regulate water temperature are exposed to hot water, ensure that the temperature of the water does not exceed 110 °Fahrenheit.

(e) Standard: Heating and ventilation. (1) Each client bedroom in the facility must have—
   (i) At least one window to the outside; and
   (ii) Direct outside ventilation by means of windows, air conditioning, or mechanical ventilation.

(2) The facility must—
   (i) Maintain the temperature and humidity within a normal comfort range by heating, air conditioning or other means; and
   (ii) Ensure that the heating apparatus does not constitute a burn or smoke hazard to clients.

(f) Standard: Floors. The facility must have—
   (1) Floors that have a resilient, non-abrasive, and slip-resistant surface;
   (2) Nonabrasive carpeting, if the area used by clients is carpeted and serves clients who lie on the floor or ambulate with parts of their bodies, other than feet, touching the floor; and
   (3) Exposed floor surfaces and floor coverings that promote mobility in areas used by clients, and promote maintenance of sanitary conditions.

(g) Standard: Space and equipment. The facility must—
(1) Provide sufficient space and equipment in dining, living, health services, recreation, and program areas (including adequately equipped and sound treated areas for hearing and other evaluations if they are conducted in the facility) to enable staff to provide clients with needed services as required by this subpart and as identified in each client's individual program plan.

(2) Furnish, maintain in good repair, and teach clients to use and to make informed choices about the use of dentures, eyeglasses, hearing and other communications aids, braces, and other devices identified by the interdisciplinary team as needed by the client.

(3) Provide adequate clean linen and dirty linen storage areas.

(h) Standard: Emergency plan and procedures. (1) The facility must develop and implement detailed written plans and procedures to meet all potential emergencies and disasters such as fire, severe weather, and missing clients.

(2) The facility must communicate, periodically review, make the plan available, and provide training to the staff.

(i) Standard: Evacuation drills. (1) The facility must hold evacuation drills at least quarterly for each shift of personnel and under varied conditions to—

(i) Ensure that all personnel on all shifts are trained to perform assigned tasks;

(ii) Ensure that all personnel on all shifts are familiar with the use of the facility's fire protection features; and

(iii) Evaluate the effectiveness of emergency and disaster plans and procedures.

(2) The facility must—

(i) Actually evacuate clients during at least one drill each year on each shift;

(ii) Make special provisions for the evacuation of clients with physical disabilities;

(iii) File a report and evaluation on each evacuation drill;

(iv) Investigate all problems with evacuation drills, including accidents, and take corrective action; and

(v) During fire drills, clients may be evacuated to a safe area in facilities certified under the Health Care Occupancies Chapter of the Life Safety Code.

(3) Facilities must meet the requirements of paragraphs (i)(1) and (2) of this section for any live-in and relief staff they utilize.

(j) Standard: Fire protection—(1) General. (i) Except as specified in paragraph (j)(2) of this section, the facility must meet the applicable provisions of either the Health Care Occupancies Chapters or the Residential Board and Care Occupancies Chapter of the Life Safety Code (LSC) of the National Fire Protection Association, 1985 edition, which is incorporated by reference.

(ii) The State survey agency may apply a single chapter of the LSC to the entire facility or may apply different chapters to different buildings or parts of buildings as permitted by the LSC.

(iii) A facility that meets the LSC definition of a residential board and care occupancy and that has 16 or fewer beds, must have its evacuation capability evaluated in accordance with the Evacuation Difficulty Index of the LSC (appendix F).

(2) Exceptions. (i) For facilities that meet the LSC definition of a health care occupancy:

(A) The State survey agency may waive, for a period it considers appropriate, specific provisions of the LSC if—

(1) The waiver would not adversely affect the health and safety of the clients; and

(2) Rigid application of specific provisions would result in an unreasonable hardship for the facility.
(B) The State survey agency may apply the State's fire and safety code instead of the LSC if the Secretary finds that the State has a code imposed by State law that adequately protects a facility's clients.

(C) Compliance on November 26, 1982 with the 1967 edition of the LSC or compliance on April 18, 1986 with the 1981 edition of the LSC, with or without waivers, is considered to be compliance with this standard as long as the facility continues to remain in compliance with that edition of the Code.

(ii) For facilities that meet the LSC definition of a residential board and care occupancy and that have more than 16 beds, the State survey agency may apply the State's fire and safety code as specified in paragraph (j)(2)(B) of this section.

(k) Standard: Paint. The facility must—

(1) Use lead-free paint inside the facility; and—

(2) Remove or cover interior paint or plaster containing lead so that it is not accessible to clients.

(I) Standard: Infection control.

(1) The facility must provide a sanitary environment to avoid sources and transmission of infections. There must be an active program for the prevention, control, and investigation of infection and communicable diseases.

(2) The facility must implement successful corrective action in affected problem areas.

(3) The facility must maintain a record of incidents and corrective actions related to infections.

(4) The facility must prohibit employees with symptoms or signs of a communicable disease from direct contact with clients and their food.

§ 483.480 Condition of participation: Dietetic services.

(a) Standard: Food and nutrition services. (1) Each client must receive a nourishing, well-balanced diet including modified and specially-prescribed diets.

(2) A qualified dietitian must be employed either full-time, part-time, or on a consultant basis at the facility's discretion.

(3) If a qualified dietitian is not employed full-time, the facility must designate a person to serve as the director of food services.

(4) The client's interdisciplinary team, including a qualified dietitian and physician, must prescribe all modified and special diets including those used as a part of a program to manage inappropriate client behavior.

(5) Foods proposed for use as a primary reinforcement of adaptive behavior are evaluated in light of the client's nutritional status and needs.

(6) Unless otherwise specified by medical needs, the diet must be prepared at least in accordance with the latest edition of the recommended dietary allowances of the Food and Nutrition Board of the National Research Council, National Academy of Sciences, adjusted for age, sex, disability and activity.

(b) Standard: Meal services. (1) Each client must receive at least three meals daily, at regular times comparable to normal mealtimes in the community with—

(i) Not more than 14 hours between a substantial evening meal and breakfast of the following day, except on weekends and holidays when a nourishing snack is provided at bedtime, 16 hours may elapse between a substantial evening meal and breakfast; and

(ii) Not less than 10 hours between breakfast and the evening meal of the same day, except as provided under paragraph (b)(1)(i) of this section.

(2) Food must be served—

(i) In appropriate quantity;

(ii) At appropriate temperature;

(iii) In a form consistent with the developmental level of the client; and

(iv) With appropriate utensils.

(3) Food served to clients individually and uneaten must be discarded.

(c) Standard: Menus. (1) Menus must—

(i) Be prepared in advance;

(ii) Provide a variety of foods at each meal;

(iii) Be different for the same days of each week and adjusted for seasonal changes; and

(iv) Include the average portion sizes for menu items.

(2) Menus for food actually served must be kept on file for 30 days.

(d) Standard: Dining areas and service. The facility must—

(1) Serve meals for all clients, including persons with ambulation deficits,
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§ 484.1 Basis and scope.

(a) Basis and scope. This part is based on the indicated provisions of the following sections of the Act:

(1) Sections 1861(o) and 1891 establish the conditions that an HHA must meet in order to participate in Medicare.

(2) Provide table service for all clients who can and will eat at a table, including clients in wheelchairs;

(3) Equip areas with tables, chairs, eating utensils, and dishes designed to meet the developmental needs of each client;

(4) Supervise and staff dining rooms adequately to direct self-help dining procedure, to assure that each client receives enough food and to assure that each client eats in a manner consistent with his or her developmental level; and

(5) Ensure that each client eats in an upright position, unless otherwise specified by the interdisciplinary team or a physician.
§ 484.2 Definitions.  
As used in this part, unless the context indicates otherwise—

Bylaws or equivalent means a set of rules adopted by an HHA for governing the agency’s operation.

Branch office means a location or site from which a home health agency provides services within a portion of the total geographic area served by the parent agency. The branch office is part of the home health agency and is located sufficiently close to share administration, supervision, and services in a manner that renders it unnecessary for the branch independently to meet the conditions of participation as a home health agency.

Clinical note means a notation of a contact with a patient that is written and dated by a member of the health team, and that describes signs and symptoms, treatment and drugs administered and the patient’s reaction, and any changes in physical or emotional condition.

HHA stands for home health agency.


Parent home health agency means the agency that develops and maintains administrative controls of subunits and/or branch offices.

Primary home health agency means the agency that is responsible for the services furnished to patients and for implementation of the plan of care.

Progress note means a written notation, dated and signed by a member of the health team, that summarizes facts about care furnished and the patient’s response during a given period of time.

Proprietary agency means a private profit-making agency licensed by the State.

Public agency means an agency operated by a State or local government.

Subdivision means a component of a multi-function health agency, such as the home care department of a hospital or the nursing division of a health department, which independently meets the conditions of participation for HHAs. A subdivision that has subunits or branch offices is considered a parent agency.

Subunit means a semi-autonomous organization that—

(1) Serves patients in a geographic area different from that of the parent agency; and

(2) Must independently meet the conditions of participation for HHAs because it is too far from the parent agency to share administration, supervision, and services on a daily basis.

Summary report means the compilation of the pertinent factors of a patient’s clinical notes and progress notes that is submitted to the patient’s physician.

Supervision means authoritative procedural guidance by a qualified person for the accomplishment of a function or activity. Unless otherwise specified in this part, the supervisor must be on the premises to supervise an individual who does not meet the qualifications specified in § 484.4.

§ 484.4 Personnel qualifications.  
Staff required to meet the conditions set forth in this part are staff who meet the qualifications specified in this section.

Administrator, home health agency. A person who:

(a) Is a licensed physician; or

(b) Is a registered nurse; or

(c) Has training and experience in health service administration and at least 1 year of supervisory or administrative experience in home health care or related health programs.

Audiologist. A person who:

(a) Meets the education and experience requirements for a Certificate of Clinical Competence in audiology granted by the American Speech-Language-Hearing Association; or
(b) Meets the educational requirements for certification and is in the process of accumulating the supervised experience required for certification.

Home health aide. Effective for services furnished after August 14, 1990, a person who has successfully completed a State-established or other training program that meets the requirements of §484.36(a) and a competency evaluation program or State licensure program that meets the requirements of §484.36(b) or (e), or a competency evaluation program or State licensure program that meets the requirements of §484.36(b) or (e). An individual is not considered to have completed a training and competency evaluation program, or a competency evaluation program if, since the individual’s most recent completion of this program(s), there has been a continuous period of 24 consecutive months during none of which the individual furnished services described in §409.40 of this chapter for compensation.

Occupational therapist. A person who:
(a) Is a graduate of an occupational therapy curriculum accredited jointly by the Committee on Allied Health Education and Accreditation of the American Medical Association and the American Occupational Therapy Association; or
(b) Is eligible for the National Registration Examination of the American Occupational Therapy Association; or
(c) Has 2 years of appropriate experience as an occupational therapist, and has achieved a satisfactory grade on a proficiency examination conducted, approved, or sponsored by the U.S. Public Health Service, except that such determinations of proficiency do not apply with respect to persons initially licensed by a State or seeking initial qualification as an occupational therapist after December 31, 1977.

(d) Was licensed or registered prior to January 1, 1966, and prior to January 1, 1970, had 15 years of full-time experience in the treatment of illness or injury through the practice of physical therapy in which services were rendered under the order and direction of attending and referring doctors of medicine or osteopathy; or
(e) If trained outside the United States,
(1) Was graduated since 1928 from a physical therapy curriculum approved in the country in which the curriculum was located and in which there is a
member organization of the World Confederation for Physical Therapy.

(2) Meets the requirements for membership in a member organization of the World Confederation for Physical Therapy.

Physical therapy assistant. A person who is licensed as a physical therapy assistant, if applicable, by the State in which practicing, and

(1) Has graduated from a 2-year college-level program approved by the American Physical Therapy Association; or

(2) Has 2 years of appropriate experience as a physical therapy assistant, and has achieved a satisfactory grade on a proficiency examination conducted, approved, or sponsored by the U.S. Public Health Service, except that these determinations of proficiency do not apply with respect to persons initially licensed by a State or seeking initial qualification as a social work assistant after December 31, 1977.

Social worker. A person who has a master’s degree from a school of social work accredited by the Council on Social Work Education, and has 1 year of social work experience in a health care setting.

Speech-language pathologist. A person who:

(1) Meets the education and experience requirements for a Certificate of Clinical Competence in (speech pathology or audiology) granted by the American Speech-Language-Hearing Association; or

(2) Meets the educational requirements for certification and is in the process of accumulating the supervised experience required for certification.

[54 FR 33367, August 14, 1989, as amended at 56 FR 32973, July 18, 1991]

Subpart B—Administration

§ 484.10 Condition of participation: Patient rights.

The patient has the right to be informed of his or her rights. The HHA must protect and promote the exercise of these rights.

(a) Standard: Notice of rights. (1) The HHA must provide the patient with a written notice of the patient’s rights in advance of furnishing care to the patient or during the initial evaluation visit before the initiation of treatment.

(2) The HHA must maintain documentation showing that it has complied with the requirements of this section.

(b) Standard: Exercise of rights and respect for property and person. (1) The patient has the right to exercise his or her rights as a patient of the HHA.

(2) The patient’s family or guardian may exercise the patient’s rights when the patient has been judged incompetent.

(3) The patient has the right to have his or her property treated with respect.

(4) The patient has the right to voice grievances regarding treatment or care that is (or fails to be) furnished, or regarding the lack of respect for property...
by anyone who is furnishing services on behalf of the HHA and must not be subjected to discrimination or reprisal for doing so.

(5) The HHA must investigate complaints made by a patient or the patient’s family or guardian regarding treatment or care that is (or fails to be) furnished, or regarding the lack of respect for the patient’s property by anyone furnishing services on behalf of the HHA, and must document both the existence of the complaint and the resolution of the complaint.

(c) Standard: Right to be informed and to participate in planning care and treatment. (1) The patient has the right to be informed, in advance about the care to be furnished, and of any changes in the care to be furnished.

(i) The HHA must advise the patient in advance of the disciplines that will furnish care, and the frequency of visits proposed to be furnished.

(ii) The HHA must advise the patient in advance of any change in the plan of care before the change is made.

(2) The patient has the right to participate in the planning of the care.

(i) The HHA must advise the patient in advance of the right to participate in planning the care or treatment and in planning changes in the care or treatment.

(ii) The HHA complies with the requirements of subpart I of part 489 of this chapter relating to maintaining written policies and procedures regarding advance directives. The HHA must inform and distribute written information to the patient, in advance, concerning its policies on advance directives, including a description of applicable State law. The HHA may furnish advance directives information to a patient at the time of the first home visit, as long as the information is furnished before care is provided.

(d) Standard: Confidentiality of medical records. The patient has the right to confidentiality of the clinical records maintained by the HHA. The HHA must advise the patient of the agency’s policies and procedures regarding disclosure of clinical records.

(e) Standard: Patient liability for payment. (1) The patient has the right to be advised, before care is initiated, of the extent to which payment for the HHA services may be expected from Medicare or other sources, and the extent to which payment may be required from the patient. Before the care is initiated, the HHA must inform the patient, orally and in writing, of—

(i) The extent to which payment may be expected from Medicare, Medicaid, or any other federally funded or aided program known to the HHA;

(ii) The charges for services that will not be covered by Medicare; and

(iii) The charges that the individual may have to pay.

(2) The patient has the right to be advised orally and in writing of any changes in the information provided in accordance with paragraph (e)(1) of this section when they occur. The HHA must advise the patient of these changes orally and in writing as soon as possible, but no later than 30 calendar days from the date that the HHA becomes aware of a change.

(f) Standard: Home health hotline. The patient has the right to be advised of the availability of the toll-free HHA hotline in the State. When the agency accepts the patient for treatment or care, the HHA must advise the patient in writing of the telephone number of the home health hotline established by the State, the hours of its operation, and that the purpose of the hotline is to receive complaints or questions about local HHAs. The patient also has the right to use this hotline to lodge complaints concerning the implementation of the advance directives requirements.

[54 FR 33367, August 14, 1989, as amended at 56 FR 32973, July 18, 1991; 57 FR 8203, Mar. 6, 1992; 60 FR 33293, June 27, 1995]

§ 484.11 Condition of participation: Release of patient identifiable OASIS information.

The HHA and agent acting on behalf of the HHA in accordance with a written contract must ensure the confidentiality of all patient identifiable information contained in the clinical record, including OASIS data, and may not release patient identifiable OASIS information to the public.

[64 FR 3763, Jan. 25, 1999]
§ 484.12 Condition of participation: Compliance with Federal, State, and local laws, disclosure and ownership information, and accepted professional standards and principles.

(a) Standard: Compliance with Federal, State, and local laws and regulations. The HHA and its staff must operate and furnish services in compliance with all applicable Federal, State, and local laws and regulations. If State or applicable local law provides for the licensure of HHAs, an agency not subject to licensure is approved by the licensing authority as meeting the standards established for licensure.

(b) Standard: Disclosure of ownership and management information. The HHA must comply with the requirements of Part 420, Subpart C of this chapter. The HHA also must disclose the following information to the State survey agency at the time of the HHA's initial request for certification, for each survey, and at the time of any change in ownership or management:

(1) The name and address of all persons with an ownership or control interest in the HHA as defined in §§ 420.201, 420.202, and 420.206 of this chapter.

(2) The name and address of each person who is an officer, a director, an agent or a managing employee of the HHA as defined in §§ 420.201, 420.202, and 420.206 of this chapter.

(3) The name and address of the corporation, association, or other company that is responsible for the management of the HHA, and the name and address of the chief executive officer and the chairman of the board of directors of that corporation, association, or other company responsible for the management of the HHA.

(c) Standard: Compliance with accepted professional standards and principles. The HHA and its staff must comply with accepted professional standards and principles that apply to professionals furnishing services in an HHA.

§ 484.14 Condition of participation: Organization, services, and administration.

Organization, services furnished, administrative control, and lines of authority for the delegation of responsibility down to the patient care level are clearly set forth in writing and are readily identifiable. Administrative and supervisory functions are not delegated to another agency or organization and all services not furnished directly, including services provided through subunits are monitored and controlled by the parent agency. If an agency has subunits, appropriate administrative records are maintained for each subunit.

(a) Standard: Services furnished. Part-time or intermittent skilled nursing services and at least one other therapeutic service (physical, speech, or occupational therapy; medical social services; or home health aide services) are made available on a visiting basis, in a place of residence used as a patient's home. An HHA must provide at least one of the qualifying services directly through agency employees, but may provide the second qualifying service and additional services under arrangements with another agency or organization.

(b) Standard: Governing body. A governing body (or designated persons so functioning) assumes full legal authority and responsibility for the operation of the agency. The governing body appoints a qualified administrator, arranges for professional advice as required under § 484.16, adopts and periodically reviews written bylaws or an acceptable equivalent, and oversees the management and fiscal affairs of the agency.

(c) Standard: Administrator. The administrator, who may also be the supervising physician or registered nurse required under paragraph (d) of this section, organizes and directs the agency's ongoing functions; maintains ongoing liaison among the governing body, the group of professional personnel, and the staff; employs qualified personnel and ensures adequate staff education and evaluations; ensures the accuracy of public information materials and activities; and implements an effective budgeting and accounting system. A qualified person is authorized in writing to act in the absence of the administrator.

(d) Standard: Supervising physician or registered nurse. The skilled nursing and other therapeutic services furnished
are under the supervision and direction of a physician or a registered nurse (who preferably has at least 1 year of nursing experience and is a public health nurse). This person, or similarly qualified alternate, is available at all times during operating hours and participates in all activities relevant to the professional services furnished, including the development of qualifications and the assignment of personnel.

(e) Standard: Personnel policies. Personnel practices and patient care are supported by appropriate, written personnel policies. Personnel records include qualifications and licensure that are kept current.

(f) Standard: Personnel under hourly or per visit contracts. If personnel under hourly or per visit contracts are used by the HHA, there is a written contract between those personnel and the agency that specifies the following:

1. Patients are accepted for care only by the primary HHA.
2. The services to be furnished.
3. The necessity to conform to all applicable agency policies, including personnel qualifications.
4. The responsibility for participating in developing plans of care.
5. The manner in which services will be controlled, coordinated, and evaluated by the primary HHA.
6. The procedures for submitting clinical and progress notes, scheduling of visits, periodic patient evaluation.
7. The procedures for payment for services furnished under the contract.

(g) Standard: Coordination of patient services. All personnel furnishing services maintain liaison to ensure that their efforts are coordinated effectively and support the objectives outlined in the plan of care. The clinical record or minutes of case conferences establish that effective interchange, reporting, and coordination of patient care does occur. A written summary report for each patient is sent to the attending physician at least every 62 days.

(h) Standard: Services under arrangements. Services furnished under arrangements are subject to a written contract conforming with the requirements specified in paragraph (f) of this section and with the requirements of section 1861(w) of the Act (42 U.S.C. 1495x(w)).

(i) Standard: Institutional planning. The HHA, under the direction of the governing body, prepares an overall plan and a budget that includes an annual operating budget and capital expenditure plan.

1. Annual operating budget. There is an annual operating budget that includes all anticipated income and expenses related to items that would, under generally accepted accounting principles, be considered income and expense items. However, it is not required that there be prepared, in connection with any budget, an item by item identification of the components of each type of anticipated income or expense.

2. Capital expenditure plan. There is a capital expenditure plan for at least a 3-year period, including the operating budget year. The plan includes and identifies in detail the anticipated sources of financing for, and the objectives of, each anticipated expenditure of more than $600,000 for items that would under generally accepted accounting principles, be considered capital items. In determining if a single capital expenditure exceeds $600,000, the cost of studies, surveys, designs, plans, working drawings, specifications, and other activities essential to the acquisition, improvement, modernization, expansion, or replacement of land, plant, building, and equipment are included. Expenditures directly or indirectly related to capital expenditures, such as grading, paving, broker commissions, taxes assessed during the construction period, and costs involved in demolishing or razing structures on land are also included. Transactions that are separated in time, but are components of an overall plan or patient care objective, are viewed in their entirety without regard to their timing. Other costs related to capital expenditures include title fees, permit and license fees, broker commissions, architect, legal, accounting, and appraisal fees; interest, finance, or carrying charges on bonds, notes and other costs incurred for borrowing funds.

(ii) If the anticipated source of financing is, in any part, the anticipated payment from title V (Maternal and Child Health and Crippled Children's
§ 484.16 Condition of participation: Group of professional personnel.

A group of professional personnel, which includes at least one physician and one registered nurse (preferably a public health nurse), and with appropriate representation from other professional disciplines, establishes and annually reviews the agency's policies governing scope of services offered, admission and discharge policies, medical supervision and plans of care, emergency care, clinical records, personnel qualifications, and program evaluation. At least one member of the group is neither an owner nor an employee of the agency.

(a) Standard: Advisory and evaluation function. The group of professional personnel meets frequently to advise the agency on professional issues, to participate in the evaluation of the agency's program, and to assist the agency in maintaining liaison with other health care providers in the community and in the agency's community information program. The meetings are documented by dated minutes.

[54 FR 33367, August 14, 1989, as amended at 56 FR 32974, July 18, 1991]

§ 484.18 Condition of participation: Acceptance of patients, plan of care, and medical supervision.

Patients are accepted for treatment on the basis of a reasonable expectation that the patient's medical, nursing, and social needs can be met adequately by the agency in the patient's place of residence. Care follows a written plan of care established and periodically reviewed by a doctor of medicine, osteopathy, or podiatric medicine.

(a) Standard: Plan of care. The plan of care developed in consultation with the agency staff covers all pertinent diagnoses, including mental status, types of services and equipment required, frequency of visits, prognosis, rehabilitation potential, functional limitations, activities permitted, nutritional requirements, medications and treatments, any safety measures to protect against injury, instructions for timely discharge or referral, and any other appropriate items. If a physician refers a
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patient under a plan of care that cannot be completed until after an evaluation visit, the physician is consulted to approve additions or modifications to the original plan. Orders for therapy services include the specific procedures and modalities to be used and the amount, frequency, and duration. The therapist and other agency personnel participate in developing the plan of care.

(b) Standard: Periodic review of plan of care. The total plan of care is reviewed by the attending physician and HHA personnel as often as the severity of the patient’s condition requires, but at least once every 60 days or more frequently when there is a beneficiary elected transfer; a significant change in condition resulting in a change in the case-mix assignment; or a discharge and return to the same HHA during the 60-day episode. Agency professional staff promptly alert the physician to any changes that suggest a need to alter the plan of care.

(c) Standard: Conformance with physician orders. Drugs and treatments are administered by agency staff only as ordered by the physician. Verbal orders are put in writing and 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. Verbal orders are only accepted by personnel authorized to do so by applicable State and Federal laws and regulations as well as by the HHA’s internal policies.

§ 484.20 Condition of participation: Reporting OASIS information.

HHAs must electronically report all OASIS data collected in accordance with § 484.55.

(a) Standard: Encoding OASIS data. The HHA must encode and be capable of transmitting OASIS data for each agency patient within 7 days of completing an OASIS data set.

(b) Standard: Accuracy of encoded OASIS data. The encoded OASIS data must accurately reflect the patient’s status at the time of assessment.

(c) Standard: Transmittal of OASIS data. The HHA must—

1. Electronically transmit accurate, completed, encoded and locked OASIS data for each patient to the State agency or HCFA OASIS contractor at least monthly;

2. For all assessments completed in the previous month, transmit OASIS data in a format that meets the requirements of paragraph (d) of this section;

3. Successfully transmit test data to the State agency or HCFA OASIS contractor beginning March 26, 1999, and no later than April 26, 1999; and

4. Transmit data using electronic communications software that provides a direct telephone connection from the HHA to the State agency or HCFA OASIS contractor.

(d) Standard: Data Format. The HHA must encode and transmit data using the software available from HCFA or software that conforms to HCFA standard electronic record layout, edit specifications, and data dictionary, and that includes the required OASIS data set.

[64 FR 3763, Jan. 25, 1999]

Subpart C—Furnishing of Services

§ 484.30 Condition of participation: Skilled nursing services.

The HHA furnishes skilled nursing services by or under the supervision of a registered nurse and in accordance with the plan of care.

(a) Standard: Duties of the registered nurse. The registered nurse makes the initial evaluation visit, regularly re-evaluates the patient’s nursing needs, initiates the plan of care and necessary revisions, furnishes those services requiring substantial and specialized nursing skill, initiates appropriate preventive and rehabilitative nursing procedures, prepares clinical and progress notes, coordinates services, informs the physician and other personnel of changes in the patient’s condition and needs, counsels the patient and family in meeting nursing and related needs, participates in in-service programs, and supervises and teaches other nursing personnel.

(b) Standard: Duties of the licensed practical nurse. The licensed practical nurse furnishes services in accordance
§ 484.32 Condition of participation: Therapy services.

Any therapy services offered by the HHA directly or under arrangement are given by a qualified therapist or by a qualified therapy assistant under the supervision of a qualified therapist and in accordance with the plan of care. The qualified therapist assists the physician in evaluating level of function, helps develop the plan of care (revising it as necessary), prepares clinical and progress notes, advises and consults with the family and other agency personnel, and participates in in-service programs.

(a) Standard: Supervision of physical therapy assistant and occupational therapy assistant. Services furnished by a qualified physical therapy assistant or qualified occupational therapy assistant may be furnished under the supervision of a qualified physical or occupational therapist. A physical therapy assistant or occupational therapy assistant performs services planned, delegated, and supervised by the therapist, assists in preparing clinical notes and progress reports, and participates in educating the patient and family, and in in-service programs.

(b) Standard: Supervision of speech therapy services. Speech therapy services are furnished only by or under supervision of a qualified speech pathologist or audiologist.

§ 484.36 Condition of participation: Home health aide services.

Home health aides are selected on the basis of such factors as a sympathetic attitude toward the care of the sick, ability to read, write, and carry out directions, and maturity and ability to deal effectively with the demands of the job. They are closely supervised to ensure their competence in providing care. For home health services furnished (either directly or through arrangements with other organizations) after August 14, 1990, the HHA must use individuals who meet the personnel qualifications specified in §484.4 for “home health aide”.

(a) Standard: Home health aide training—(1) Content and duration of training. The aide training program must address each of the following subject areas through classroom and supervised practical training totalling at least 75 hours, with at least 16 hours devoted to supervised practical training. The individual being trained must complete at least 16 hours of classroom training before beginning the supervised practical training.

(i) Communications skills.

(ii) Observation, reporting and documentation of patient status and the care or service furnished.

(iii) Reading and recording temperature, pulse, and respiration.

(iv) Basic infection control procedures.

(v) Basic elements of body functioning and changes in body function that must be reported to an aide’s supervisor.

(vi) Maintenance of a clean, safe, and healthy environment.

(vii) Recognizing emergencies and knowledge of emergency procedures.
work with the populations served by the HHA, including the need for respect for the patient, his or her privacy and his or her property.

(ix) Appropriate and safe techniques in personal hygiene and grooming that include:

(A) Bed bath.
(B) Sponge, tub, or shower bath.
(C) Shampoo, sink, tub, or bed.
(D) Nail and skin care.
(E) Oral hygiene.
(F) Toileting and elimination.
(x) Safe transfer techniques and ambulation.
(xi) Normal range of motion and positioning.
(xii) Adequate nutrition and fluid intake.
(xiii) Any other task that the HHA may choose to have the home health aide perform.

“Supervised practical training” means training in a laboratory or other setting in which the trainee demonstrates knowledge while performing tasks on an individual under the direct supervision of a registered nurse or licensed practical nurse.

(2) Conduct of training—(i) Organizations. A home health aide training program may be offered by any organization except an HHA that, within the previous 2 years has been found—

(A) Out of compliance with requirements of this paragraph (a) or paragraph (b) of this section;

(B) To permit an individual that does not meet the definition of “home health aide” as specified in §484.4 to furnish home health aide services (with the exception of licensed health professionals and volunteers);

(C) Has been subject to an extended (or partial extended) survey as a result of having been found to have furnished substandard care (or for other reasons at the discretion of the HCFA or the State);

(D) Has been assessed a civil monetary penalty of not less than $5,000 as an intermediate sanction;

(E) Has been found to have compliance deficiencies that endanger the health and safety of the HHA’s patients and has had a temporary management appointed to oversee the management of the HHA;

(F) Has had all or part of its Medicare payments suspended; or

(G) Under any Federal or State law within the 2-year period beginning on October 1, 1988—

(1) Has had its participation in the Medicare program terminated;

(2) Has been assessed a penalty of not less than $5,000 for deficiencies in Federal or State standards for HHAs;

(3) Was subject to a suspension of Medicare payments to which it otherwise would have been entitled;

(4) Had operated under a temporary management that was appointed to oversee the operation of the HHA and to ensure the health and safety of the HHA’s patients; or

(5) Was closed or had its residents transferred by the State.

(ii) Qualifications for instructors. The training of home health aides and the supervision of home health aides during the supervised practical portion of the training must be performed by or under the general supervision of a registered nurse who possesses a minimum of 2 years of nursing experience, at least 1 year of which must be in the provision of home health care. Other individuals may be used to provide instruction under the supervision of a qualified registered nurse.

(3) Documentation of training. The HHA must maintain sufficient documentation to demonstrate that the requirements of this standard are met.

(b) Standard: Competency evaluation and in-service training—(1) Applicability. An individual may furnish home health aide services on behalf of an HHA only after that individual has successfully completed a competency evaluation program as described in this paragraph. The HHA is responsible for ensuring that the individuals who furnish home health aide services on its behalf meet the competency evaluation requirements of this section.

(2) Content and frequency of evaluations and amount of in-service training. The competency evaluation must address each of the subjects listed in paragraph (a)(1)(i) through (xiii) of this section.

(ii) The HHA must complete a performance review of each home health aide no less frequently than every 12 months.
(iii) The home health aide must receive at least 12 hours of in-service training during each 12-month period. The in-service training may be furnished while the aide is furnishing care to the patient.

(3) Conduct of evaluation and training—(i) Organizations. A home health aide competency evaluation program may be offered by any organization except as specified in paragraph (a)(2)(i) of this section.

The in-service training may be offered by any organization.

(ii) Evaluators and instructors. The competency evaluation must be performed by a registered nurse. The in-service training generally must be supervised by a registered nurse who possesses a minimum of 2 years of nursing experience at least 1 year of which must be in the provision of home health care.

(iii) Subject areas. The subject areas listed at paragraphs (a)(1)(iii), (ix), (x), and (xi) of this section must be evaluated after observation of the aide's performance of the tasks with a patient. The other subject areas in paragraph (a)(1) of this section may be evaluated through written examination, oral examination, or after observation of a home health aide with a patient.

(4) Competency determination. (i) A home health aide is not considered competent in any task for which he or she is evaluated as “unsatisfactory.” The aide must not perform that task without direct supervision by a licensed nurse until after he or she receives training in the task for which he or she was evaluated as “unsatisfactory” and passes a subsequent evaluation with “satisfactory.”

(ii) A home health aide is not considered to have successfully passed a competency evaluation if the aide has an “unsatisfactory” rating in more than one of the required areas.

(5) Documentation of competency evaluation. The HHA must maintain documentation which demonstrates that the requirements of this standard are met.

(6) Effective date. The HHA must implement a competency evaluation program that meets the requirements of this paragraph before February 14, 1990. The HHA must provide the preparation necessary for the individual to successfully complete the competency evaluation program. After August 14, 1990, the HHA may use only those aides that have been found to be competent in accordance with §484.36(b).

(c) Standard: Assignment and duties of the home health aide—(1) Assignment. The home health aide is assigned to a specific patient by the registered nurse. Written patient care instructions for the home health aide must be prepared by the registered nurse or other appropriate professional who is responsible for the supervision of the home health aide under paragraph (d) of this section.

(2) Duties. The home health aide provides services that are ordered by the physician in the plan of care and that the aide is permitted to perform under State law. The duties of a home health aide include the provision of hands-on personal care, performance of simple procedures as an extension of therapy or nursing services, assistance in ambulation or exercises, and assistance in administering medications that are ordinarily self-administered. Any home health aide services offered by an HHA must be provided by a qualified home health aide.

(d) Standard: Supervision. (1) If the patient receives skilled nursing care, the registered nurse must perform the supervisory visit required by paragraph (d)(2) of this section. If the patient is not receiving skilled nursing care, but is receiving another skilled service (that is, physical therapy, occupational therapy, or speech-language pathology services), supervision may be provided by the appropriate therapist.

(2) The registered nurse (or another professional described in paragraph (d)(1) of this section) must make an on-site visit to the patient’s home no less frequently than every 2 weeks.

(3) If home health aide services are provided to a patient who is not receiving skilled nursing care, physical or occupational therapy or speech-language pathology services, the registered nurse must make a supervisory visit to the patient’s home no less frequently than every 62 days. In these cases, to ensure that the aide is properly caring for the patient, each supervisory visit...
must occur while the home health aide is providing patient care.

(4) If home health aide services are provided by an individual who is not employed directly by the HHA (or hospice), the services of the home health aide must be provided under arrangements, as defined in section 1861(w)(1) of the Act. If the HHA (or hospice) chooses to provide home health aide services under arrangements with another organization, the HHA’s (or hospice’s) responsibilities include, but are not limited to—

(i) Ensuring the overall quality of the care provided by the aide;

(ii) Supervision of the aide’s services as described in paragraphs (d)(1) and (d)(2) of this section; and

(iii) Ensuring that home health aides providing services under arrangements have met the training requirements of paragraphs (a) and (b) of this section.

(e) Personal care attendant: Evaluation requirements—

(1) Applicability. This paragraph applies to individuals who are employed by HHAs exclusively to furnish personal care attendant services under a Medicaid personal care benefit.

(2) Rule. An individual may furnish personal care services, as defined in §440.170 of this chapter, on behalf of an HHA after the individual has been found competent by the State to furnish those services for which a competency evaluation is required by paragraph (b) of this section and which the individual is required to perform. The individual need not be determined competent in those services listed in paragraph (a) of this section that the individual is not required to furnish.


§ 484.38 Condition of participation: Qualifying to furnish outpatient physical therapy or speech pathology services.

An HHA that wishes to furnish outpatient physical therapy or speech pathology services must meet all the pertinent conditions of this part and also meet the additional health and safety requirements set forth in §§485.711, 485.713, 485.715, 485.719, 485.723, and 485.727 of this chapter to implement section 1861(p) of the Act.

[54 FR 33367, Aug. 14, 1989, as amended at 60 FR 2239, Jan. 9, 1995; 60 FR 11632, Mar. 2, 1995]

§ 484.48 Condition of participation: Clinical records.

A clinical record containing pertinent past and current findings in accordance with accepted professional standards is maintained for every patient receiving home health services. In addition to the plan of care, the record contains appropriate identifying information; name of physician; drug, dietary, treatment, and activity orders; signed and dated clinical and progress notes; copies of summary reports sent to the attending physician; and a discharge summary. The HHA must inform the attending physician of the availability of a discharge summary. The discharge summary must be sent to the attending physician upon request and must include the patient’s medical and health status at discharge.

(a) Standards: Retention of records. Clinical records are retained for 5 years after the month the cost report to which the records apply is filed with the intermediary, unless State law stipulates a longer period of time. Policies provide for retention even if the HHA discontinues operations. If a patient is transferred to another health facility, a copy of the record or abstract is sent with the patient.

(b) Standards: Protection of records. Clinical record information is safeguarded against loss or unauthorized use. Written procedures govern use and removal of records and the conditions for release of information. Patient’s written consent is required for release of information not authorized by law.


§ 484.52 Condition of participation: Evaluation of the agency’s program.

The HHA has written policies requiring an overall evaluation of the agency’s total program at least once a year by the group of professional personnel (or a committee of this group), HHA staff, and consumers, or by professional people outside the agency working in conjunction with consumers. The evaluation consists of an overall policy and
administrative review and a clinical record review. The evaluation assesses the extent to which the agency's program is appropriate, adequate, effective, and efficient. Results of the evaluation are reported to and acted upon by those responsible for the operation of the agency and are maintained separately as administrative records.

(a) Standard: Policy and administrative review. As a part of the evaluation process the policies and administrative practices of the agency are reviewed to determine the extent to which they promote patient care that is appropriate, adequate, effective, and efficient. Mechanisms are established in writing for the collection of pertinent data to assist in evaluation.

(b) Standard: Clinical record review. At least quarterly, appropriate health professionals, representing at least the scope of the program, review a sample of both active and closed clinical records to determine whether established policies are followed in furnishing services directly or under arrangement. There is a continuing review of clinical records for each 62-day period that a patient receives home health services to determine adequacy of the plan of care and appropriateness of continuation of care.

§ 484.55 Condition of participation: Comprehensive assessment of patients.

Each patient must receive, and an HHA must provide, a patient-specific, comprehensive assessment that accurately reflects the patient’s current health status and includes information that may be used to demonstrate the patient's progress toward achievement of desired outcomes. The comprehensive assessment must identify the patient’s continuing need for home care and meet the patient’s medical, nursing, rehabilitative, social, and discharge planning needs. For Medicare beneficiaries, the HHA must verify the patient’s eligibility for the Medicare home health benefit including homebound status, both at the time of the initial assessment visit and at the time of the comprehensive assessment. The comprehensive assessment must also incorporate the use of the current version of the Outcome and Assessment Information Set (OASIS) items, using the language and groupings of the OASIS items, as specified by the Secretary.

(a) Standard: Initial assessment visit.

(1) A registered nurse must conduct an initial assessment visit to determine the immediate care and support needs of the patient; and, for Medicare patients, to determine eligibility for the Medicare home health benefit, including homebound status. The initial assessment visit must be held either within 48 hours of referral, or within 48 hours of the patient’s return home, or on the physician-ordered start of care date.

(2) When rehabilitation therapy service (speech language pathology, physical therapy, or occupational therapy) is the only service ordered by the physician, and if the need for that service establishes program eligibility, the initial assessment visit may be made by the appropriate rehabilitation skilled professional.

(b) Standard: Completion of the comprehensive assessment.

(1) The comprehensive assessment must be completed in a timely manner, consistent with the patient’s immediate needs, but no later than 5 calendar days after the start of care.

(2) Except as provided in paragraph (b)(3) of this section, a registered nurse must complete the comprehensive assessment and for Medicare patients, determine eligibility for the Medicare home health benefit, including homebound status.

(3) When physical therapy, speech-language pathology, or occupational therapy is the only service ordered by the physician, a physical therapist, speech-language pathologist or occupational therapist may complete the comprehensive assessment, and for Medicare patients, determine eligibility for the Medicare home health benefit, including homebound status. The occupational therapist may complete the comprehensive assessment if the need for occupational therapy establishes program eligibility.

(c) Standard: Drug regimen review. The comprehensive assessment must include a review of all medications the patient is currently using in order to identify any potential adverse effects.
Health Care Financing Administration, HHS § 484.205

(a) Basis. This subpart implements section 1895 of the Act, which provides for the implementation of a prospective payment system (PPS) for HHAs for portions of cost reporting periods occurring on or after October 1, 2000.

(b) Scope. This subpart sets forth the framework for the HHA PPS, including the methodology used for the development of the payment rates, associated adjustments, and related rules.

§ 484.202 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 clinical model.

Discipline means one of the six home health disciplines covered under the Medicare home health benefit (skilled nursing services, home health aide services, physical therapy services, occupational therapy services, speech-language pathology services, and medical social services).

Home health market basket index means an index that reflects changes over time in the prices of an appropriate mix of goods and services included in home health services.

§ 484.205 Basis of payment.

(a) Method of payment. An HHA receives a national prospective 60-day episode payment of a predetermined rate for a home health service previously paid on a reasonable cost basis (except the osteoporosis drug defined in section 1861(kk) of the Act) as of August 5, 1997. The national 60-day episode payment is determined in accordance with §484.215. The national prospective 60-day episode payment is subject to the following adjustments and additional payments:

(1) A low-utilization payment adjustment (LUPA) of a predetermined per-visit rate as specified in §484.230.

(2) A partial episode payment (PEP) adjustment due to an intervening event defined as a beneficiary elected transfer or a discharge and return to the same HHA during the 60-day episode, that warrants a new 60-day episode payment during an existing 60-day episode, that initiates the start of a new 60-day episode payment and a new physician certification of the new plan of care. The PEP adjustment is determined in accordance with §484.235.
(3) A significant change in condition (SCIC) payment adjustment due to the intervening event defined as a significant change in the patient's condition during an existing 60-day episode. The SCIC adjustment occurs when a beneficiary experiences a significant change in condition during a 60-day episode that was not envisioned in the original plan of care. The SCIC adjustment is determined in accordance with § 484.237.

(4) An outlier payment is determined in accordance with § 484.240.

(b) Episode payment. The national prospective 60-day episode payment represents payment in full for all costs associated with furnishing home health services previously paid on a reasonable cost basis (except the osteoporosis drug listed in section 1861(m) of the Act as defined in section 1861(kk) of the Act) as of August 5, 1997 unless the national 60-day episode payment is subject to a low-utilization payment adjustment set forth in § 484.230, a partial episode payment adjustment set forth at § 484.235, a significant change in condition payment set forth at § 484.237, or an additional outlier payment set forth in § 484.240. All payments under this system may be subject to a medical review adjustment reflecting beneficiary eligibility, medical necessity determinations, and HHRG assignment. DME provided as a home health service as defined in section 1861(m) of the Act continues to be paid the fee schedule amount.

(1) Split percentage payment for initial episodes. The initial percentage payment for initial episodes is paid to an HHA at 60 percent of the case-mix and wage adjusted 60-day episode rate. The residual final payment for initial episodes is paid at 40 percent of the case-mix and wage adjusted 60-day episode rate. Split percentage payments are made in accordance with requirements at § 409.43(c) of this chapter.

(2) Split percentage payment for subsequent episodes. The initial percentage payment for subsequent episodes is paid to an HHA at 50 percent of the case-mix and wage adjusted 60-day episode rate. The residual final payment for subsequent episodes is paid at 50 percent of the case-mix and wage adjusted 60-day episode rate. Split percentage payments are made in accordance with requirements at § 409.43(c) of this chapter.

(c) Low-utilization payment. An HHA receives a national 60-day episode payment of a predetermined rate for home health services previously paid on a reasonable cost basis as of August 5, 1997 unless HCFA determines at the end of the 60-day episode that the HHA furnished minimal services to a patient during the 60-day episode. A low-utilization payment adjustment is determined in accordance with § 484.230.

(d) Partial episode payment adjustment. An HHA receives a national 60-day episode payment of a predetermined rate for home health services previously paid on a reasonable cost basis as of August 5, 1997, unless HCFA determines an intervening event, defined as a beneficiary elected transfer, or discharge and return to the same HHA during a 60-day episode, warrants a new 60-day episode payment. The PEP adjustment would not apply in situations of transfers among HHAs of common ownership as defined in § 424.22 of this chapter. Those situations would be considered services provided under arrangement on behalf of the originating HHA by the receiving HHA with the common ownership interest for the balance of the 60-day episode. The common ownership exception to the transfer PEP adjustment does not apply if the beneficiary moves to a different MSA or Non-MSA during the 60-day episode before the transfer to the receiving HHA. The transferring HHA in situations of common ownership not only serves as a billing agent, but must also exercise professional responsibility over the arranged-for services in order for services provided under arrangements to be paid. The discharge and return to the same HHA during the 60-day episode is only recognized in those circumstances when a beneficiary reached the goals in the original plan of care. The original plan of care must have been terminated with no anticipated need for additional home health services for the balance of the 60-day episode. If the intervening event warrants a new 60-day episode payment and the new physician certification of a new plan of care, the initial HHA receives a partial episode payment adjustment reflecting the length of time the patient remained under its care.
§ 484.215 Initial establishment of the calculation of the national 60-day episode payment.

(a) Determining an HHA's costs. In calculating the initial unadjusted national 60-day episode payment applicable for a service furnished by an HHA using data on the most recent available audited cost reports, HCFA determines each HHA's costs by summing its allowable costs for the period. HCFA determines the national mean cost per visit.

(b) Determining HHA utilization. In calculating the initial unadjusted national 60-day episode payment, HCFA determines the national mean utilization for each of the six disciplines using home health claims data.

(c) Use of the market basket index. HCFA uses the HHA market basket index to adjust the HHA cost data to reflect cost increases occurring between October 1, 1996 through September 30, 2001.

(d) Calculation of the unadjusted national average prospective payment amount for the 60-day episode. HCFA calculates the unadjusted national 60-day episode payment in the following manner:

1. By computing the mean national cost per visit.
2. By computing the national mean utilization for each discipline.
3. By multiplying the mean national cost per visit by the national mean utilization summed in the aggregate for the six disciplines.
4. By adding to the amount derived in paragraph (d)(3) of this section, amounts for nonroutine medical supplies, an OASIS adjustment for estimated ongoing reporting costs, an OASIS adjustment for the one-time implementation costs associated with assessment scheduling form changes and amounts for Part B therapies that could have been unbundled to Part B prior to October 1, 2000. The resulting
§ 484.220 Calculation of the adjusted national prospective 60-day episode payment rate for case-mix and area wage levels.

(a) HCA adjusts the national prospective 60-day episode payment rate to account for—
   (1) HHA case-mix using a case-mix index to explain the relative resource utilization of different patients; and
   (2) Geographic differences in wage levels using an appropriate wage index based on the site of service for the beneficiary.

(b) The cost data described in paragraph (a) of this section to remove the effects of geographic variation in wage levels and variation in case-mix;
   (1) The cost data for geographic variation in wage levels using the hospital wage index; and
   (2) The cost data for HHA variation in case-mix using the case-mix indices and other data that indicate HHA case-mix.

§ 484.225 Annual update of the unadjusted national prospective 60-day episode payment rate.

(a) HCFA updates the unadjusted national 60-day episode payment rate on a fiscal year basis.

(b) For fiscal year 2001, the unadjusted national 60-day episode payment rate is adjusted using the latest available home health market basket index factors.

(c) For fiscal years 2002 and 2003, the unadjusted national prospective 60-day episode payment rate is updated by the proportion of the applicable home health market basket minus 11 percentage points.

(d) For subsequent fiscal years, the unadjusted national rate is equal to the rate for the previous fiscal year increased by the applicable home health market basket index amount.

§ 484.230 Methodology used for the calculation of the low-utilization payment adjustment.

An episode with four or fewer visits is paid the national per-visit amount by discipline updated annually by the applicable market basket for each visit type. The national per-visit amount is determined by using cost data set forth in §484.210(a) and adjusting by the appropriate wage index based on the site of service for the beneficiary.

§ 484.235 Methodology used for the calculation of the partial episode payment adjustment.

(a) HCFA makes a PEP adjustment to the original 60-day episode payment that is interrupted by an intervening event described in §484.205(d).

(b) The original 60-day episode payment is adjusted to reflect the length of time the beneficiary remained under the care of the original HHA based on the first billable visit date through and including the last billable visit date.

(c) The partial episode payment is calculated by determining the actual days served by the original HHA as a proportion of 60 multiplied by the initial 60-day episode payment.

§ 484.237 Methodology used for the calculation of the significant change in condition payment adjustment.

(a) HCFA makes a SCIC payment adjustment to the original 60-day episode payment that is interrupted by the intervening event defined in §484.205(e).

(b) The SCIC payment adjustment is calculated in two parts.

(1) The first part of the SCIC payment adjustment reflects the adjustment to the level of payment prior to the significant change in the patient’s condition during the 60-day episode. The first part of the SCIC adjustment is determined by taking the span of days (the first billable visit date through and including the last billable visit date) prior to the patient’s significant change in condition as a proportion of 60 multiplied by the original episode amount.

(2) The second part of the SCIC payment adjustment reflects the adjustment to the level of payment after the significant change in the patient’s condition occurs during the 60-day episode. The second part of the SCIC adjustment is calculated by using the span of days (the first billable visit date through and including the last billable
§ 484.240 Methodology used for the calculation of the outlier payment.

(a) HCFA makes an outlier payment for an episode whose estimated cost exceeds a threshold amount for each case-mix group.

(b) The outlier threshold for each case-mix group is the episode payment amount for that group, the PEP adjustment amount for the episode or the total significant change in condition adjustment amount for the episode plus a fixed dollar loss amount that is the same for all case-mix groups.

(c) The outlier payment is a proportion of the amount of estimated cost beyond the threshold.

(d) HCFA imputes the cost for each episode by multiplying the national per-visit amount of each discipline by the number of visits in the discipline and computing the total imputed cost for all disciplines.

(e) The fixed dollar loss amount and the loss sharing proportion are chosen so that the estimated total outlier payment is no more than 5 percent of total payment under home health PPS.

§ 484.245 Accelerated payments for home health agencies.

(a) General rule. Upon request, an accelerated payment may be made to an HHA that is receiving payment under the home health prospective payment system if the HHA is experiencing financial difficulties because there is a delay by the intermediary in making payment to the HHA.

(b) Approval of payment. An HHA’s request for an accelerated payment must be approved by the intermediary and HCFA.

(c) Amount of payment. The amount of the accelerated payment is computed as a percentage of the net payment for unbilled or unpaid covered services.

(d) Recovery of payment. Recovery of the accelerated payment is made by recoupment as HHA bills are processed or by direct payment by the HHA.

§ 484.250 Patient assessment data.

An HHA must submit to HCFA the OASIS data described at § 484.55(b)(1) and (d)(1) in order for HCFA to administer the payment rate methodologies described in §§ 484.215, 484.230, 484.235, and 484.237.

§ 484.260 Limitation on review.

An HHA is not entitled to judicial or administrative review under sections 1869 or 1878 of the Act, or otherwise, with regard to the establishment of the payment unit, including the national 60-day prospective episode payment rate, adjustments and outlier payments. An HHA is not entitled to the review regarding the establishment of the transition period, definition and application of the unit of payments, the computation of initial standard prospective payment amounts, the establishment of the adjustment for outliers, and the establishment of case-mix and area wage adjustment factors.

PART 485—CONDITIONS OF PARTICIPATION: SPECIALIZED PROVIDERS

Subpart A [Reserved]

Subpart B—Conditions of Participation: Comprehensive Outpatient Rehabilitation Facilities

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Subparts C-E [Reserved]
§ 485.50 Basis and scope.

This subpart sets forth the conditions that facilities must meet to be certified as comprehensive outpatient rehabilitation facilities (CORFs) under section 1861(cc)(2) of the Social Security Act and be accepted for participation in Medicare in accordance with part 489 of this chapter.

§ 485.51 Definition.

As used in this subpart, unless the context indicates otherwise, “comprehensive outpatient rehabilitation facility”, “CORF”, or “facility” means a nonresidential facility that—

(a) Is established and operated exclusively for the purpose of providing diagnostic, therapeutic, and restorative services to outpatients for the rehabilitation of injured, disabled, or sick persons, at a single fixed location, by or under the supervision of a physician; and

(b) Meets all the requirements of this subpart.

§ 485.54 Condition of participation: Compliance with State and local laws.

The facility and all personnel who provide services must be in compliance
with applicable State and local laws and regulations.

(a) Standard: Licensure of facility. If State or local law provides for licensing, the facility must be currently licensed or approved as meeting the standards established for licensure.

(b) Standard: Licensure of personnel. Personnel that provide service must be licensed, certified, or registered in accordance with applicable State and local laws.

§ 485.56 Condition of participation: Governing body and administration.

The facility must have a governing body that assumes full legal responsibility for establishing and implementing policies regarding the management and operation of the facility.

(a) Standard: Disclosure of ownership. The facility must comply with the provisions of part 420, subpart C of this chapter that require health care providers and fiscal agents to disclose certain information about ownership and control.

(b) Standard: Administrator. The governing body must appoint an administrator who—

1. Is responsible for the overall management of the facility under the authority delegated by the governing body;

2. Implements and enforces the facility's policies and procedures;

3. Designates, in writing, an individual who, in the absence of the administrator, acts on behalf of the administrator; and

4. Retains professional and administrative responsibility for all personnel providing facility services.

(c) Standard: Group of professional personnel. The facility must have a group of professional personnel associated with the facility that—

1. Develops and periodically reviews policies to govern the services provided by the facility; and

2. Consists of at least one physician and one professional representing each of the services provided by the facility.

(d) Standard: Institutional budget plan. The facility must have an institutional budget plan that meets the following conditions:

1. It is prepared, under the direction of the governing body, by a committee consisting of representatives of the governing body and the administrative staff.

2. It provides for—
   i. An annual operating budget prepared according to generally accepted accounting principles;
   ii. A 3-year capital expenditure plan if expenditures in excess of $100,000 are anticipated, for that period, for the acquisition of land; the improvement of land, buildings, and equipment; and the replacement, modernization, and expansion of buildings and equipment; and
   iii. Annual review and updating by the governing body.

(e) Standard: Patient care policies. The facility must have written patient care policies that govern the services it furnishes. The patient care policies must include the following:

1. A description of the services the facility furnishes through employees and those furnished under arrangements.

2. Rules for and personnel responsibilities in handling medical emergencies.

3. Rules for the storage, handling, and administration of drugs and biologicals.


5. Procedures for preparing and maintaining clinical records on all patients.

6. A procedure for explaining to the patient and the patient’s family the extent and purpose of the services to be provided.

7. A procedure to assist the referring physician in locating another level of care for—patients whose treatment has terminated and who are discharged.

8. A requirement that patients accepted by the facility must be under the care of a physician.

9. A requirement that there be a plan of treatment established by a physician for each patient.

10. A procedure to ensure that the group of professional personnel reviews and takes appropriate action on recommendations from the utilization review committee regarding patient care policies.
§ 485.58 Condition of participation: Comprehensive rehabilitation program.

The facility must provide a coordinated rehabilitation program that includes, at a minimum, physicians' services, physical therapy services, and social or psychological services. The services must be furnished by personnel that meet the qualifications set forth in § 485.70 and must be consistent with the plan of treatment and the results of comprehensive patient assessments.

(a) Standard: Physician services. (1) A facility physician must be present in the facility for a sufficient time to—

(i) Provide, in accordance with accepted principles of medical practice, medical direction, medical care services, and consultation;

(ii) Establish the plan of treatment in cases where a plan has not been established by the referring physician;

(iii) Assist in establishing and implementing the facility's patient care policies; and

(iv) Participate in plan of treatment reviews, patient case review conferences, comprehensive patient assessment and reassessments, and utilization review.

(2) The facility must provide for emergency physician services during the facility operating hours.

(b) Standard: Plan of treatment. For each patient, a physician must establish a plan of treatment before the facility initiates treatment. The plan of treatment must meet the following requirements:

(1) It must delineate anticipated goals and specify the type, amount, frequency and duration of services to be provided.

(2) It must be promptly evaluated after changes in the patient's condition and revised when necessary.

(3) It must, if appropriate, be developed in consultation with the facility physician and the appropriate facility professional personnel.

(4) It must be reviewed at least every 60 days by a facility physician who, when appropriate, consults with the professional personnel providing services. The results of this review must be communicated to the patient's referring physician for concurrence before treatment is continued or discontinued.

(5) It must be revised if the comprehensive reassessment of the patient's status or the results of the patient case review conference indicate the need for revision.

(c) Standard: Coordination of services. The facility must designate, in writing, a qualified professional to ensure that professional personnel coordinate their related activities and exchange information about each patient under their
care. Mechanisms to assist in the coordination of services must include—

(1) Providing to all personnel associated with the facility, a schedule indicating the frequency and type of services provided at the facility;

(2) A procedure for communicating to all patient care personnel pertinent information concerning significant changes in the patient's status;

(3) Periodic clinical record entries, noting at least the patient's status in relationship to goal attainment; and

(4) Scheduling patient case review conferences for purposes of determining appropriateness of treatment, when indicated by the results of the initial comprehensive patient assessment, reassessment(s), the recommendation of the facility physician (or other physician who established the plan of treatment), or upon the recommendation of one of the professionals providing services.

(d) Standard: Provision of services. (1) All patients must be referred to the facility by a physician who provides the following information to the facility before treatment is initiated:

(i) The patient's significant medical history.

(ii) Current medical findings.

(iii) Diagnosis(es) and contraindications to any treatment modality.

(iv) Rehabilitation goals, if determined.

(2) Services may be provided by facility employees or by others under arrangements made by the facility.

(3) The facility must have on its premises the necessary equipment to implement the plan of treatment and sufficient space to allow adequate care.

(4) The services must be furnished by personnel that meet the qualifications of §485.70 and the number of qualified personnel must be adequate for the volume and diversity of services offered.

Personnel that do not meet the qualifications specified in §485.70 may be used by the facility in assisting qualified staff. When a qualified individual is assisted by these personnel, the qualified individual must be on the premises, and must instruct these personnel in appropriate patient care service techniques and retain responsibility for their activities.

(5) A qualified professional must initiate and coordinate the appropriate portions of the plan of treatment, monitor the patient's progress, and recommend changes, in the plan, if necessary.

(6) A qualified professional representing each service made available at the facility must be either on the premises of the facility or must be available through direct telecommunication for consultation and assistance during the facility's operating hours. At least one qualified professional must be on the premises during the facility's operating hours.

(7) All services must be provided consistent with accepted professional standards and practice.

(e) Standard: Scope and site of services—(1) Basic requirements. The facility must provide all the CORF services required in the plan of treatment and, except as provided in paragraph (e)(2) of this section, must provide the services on its premises.

(2) Exceptions. Physical therapy, occupational therapy, and speech pathology services furnished away from the premises of the CORF may be covered as CORF services if Medicare payment is not otherwise made for these services. In addition, a single home visit is covered if there is need to evaluate the potential impact of the home environment on the rehabilitation goals.

(f) Standard: Patient assessment. Each qualified professional involved in the patient's care, as specified in the plan of treatment, must—

(1) Carry out an initial patient assessment; and

(2) In order to identify whether or not the current plan of treatment is appropriate, perform a patient reassessment after significant changes in the patient's status.

(g) Standard: Laboratory services. (1) If the facility provides its own laboratory services, the services must meet the applicable requirements for laboratories specified in part 493 of this chapter.

(2) If the facility chooses to refer specimens for laboratory testing, the referral laboratory must be certified in the appropriate specialties and sub-specialties of services in accordance
§485.60 Condition of participation: Clinical records.

The facility must maintain clinical records on all patients in accordance with accepted professional standards and practice. The clinical records must be completely, promptly, and accurately documented, readily accessible, and systematically organized to facilitate retrieval and compilation of information.

(a) Standard: Content. Each clinical record must contain sufficient information to identify the patient clearly and to justify the diagnosis and treatment. Entries in the clinical record must be made as frequently as is necessary to insure effective treatment and must be signed by personnel providing services. All entries made by assistant level personnel must be countersigned by the corresponding professional. Documentation on each patient must be consolidated into one clinical record that must contain—

(1) The initial assessment and subsequent reassessments of the patient’s needs;
(2) Current plan of treatment;
(3) Identification data and consent or authorization forms;
(4) Pertinent medical history, past and present;
(5) A report of pertinent physical examinations if any;
(6) Progress notes or other documentation that reflect patient reaction to treatment, tests, or injury, or the need to change the established plan of treatment; and
(7) Upon discharge, a discharge summary including patient status relative to goal achievement, prognosis, and future treatment considerations.

(b) Standard: Protection of clinical record information. The facility must safeguard clinical record information against loss, destruction, or unauthorized use. The facility must have procedures that govern the use and removal of records and the conditions for release of information. The facility must obtain the patient’s written consent before releasing information not required to be released by law.

(c) Standard: Retention and preservation. The facility must retain clinical record information for 5 years after patient discharge and must make provision for the maintenance of such records in the event that it is no longer able to treat patients.

§485.62 Condition of participation: Physical environment.

The facility must provide a physical environment that protects the health and safety of patients, personnel, and the public.

(a) Standard: Safety and comfort of patients. The physical premises of the facility and those areas of its surrounding physical structure that are used by the patients (including at least all stairwells, corridors and passageways) must meet the following requirements:

(1) Applicable Federal, State, and local building, fire, and safety codes must be met.
(2) Fire extinguishers must be easily accessible and fire regulations must be prominently posted.
(3) A fire alarm system with local (in-house) capability must be functional, and where power is generated by electricity, an alternate power source with automatic triggering must be present.
(4) Lights, supported by an emergency power source, must be placed at exits.
(5) A sufficient number of staff to evacuate patients during a disaster must be on the premises of the facility whenever patients are being treated.
(6) Lighting must be sufficient to carry out services safely; room temperature must be maintained at comfortable levels; and ventilation through windows, mechanical means, or a combination of both must be provided.
(7) Safe and sufficient space must be available for the scope of services offered.

(b) Standard: Sanitary environment. The facility must maintain a sanitary environment and establish a program to identify, investigate, prevent, and control the cause of patient infections.

(1) The facility must establish written policies and procedures designed to
control and prevent infection in the facility and to investigate and identify possible causes of infection.

(2) The facility must monitor the infection control program to ensure that the staff implement the policies and procedures and that the policies and procedures are consistent with current practices in the field.

(3) The facility must make available at all times a quantity of laundered linen adequate for proper care and comfort of patients. Linens must be handled, stored, and processed in a manner that prevents the spread of infection.

(4) Provisions must be in effect to ensure that the facility’s premises are maintained free of rodent and insect infestation.

(c) Standard: Maintenance of equipment, physical location, and grounds. The facility must establish a written preventive maintenance program to ensure that—

(1) All equipment is properly maintained and equipment needing periodic calibration is calibrated consistent with the manufacturer’s recommendations; and

(2) The interior of the facility, the exterior of the physical structure housing the facility, and the exterior walkways and parking areas are clean and orderly and maintained free of any defects that are a hazard to patients, personnel, and the public.

(d) Standard: Access for the physically impaired. The facility must ensure the following:

(1) Doorways, stairwells, corridors, and passageways used by patients are—

(i) Of adequate width to allow for easy movement of all patients (including those on stretchers or in wheelchairs); and

(ii) In the case of stairwells, equipped with firmly attached handrails on at least one side.

(2) At least one toilet facility is accessible and constructed to allow utilization by ambulatory and nonambulatory individuals.

(3) At least one entrance is usable by individuals in wheelchairs.

(4) In multi-story buildings, elevators are accessible to and usable by the physically impaired on the level that they use to enter the building and all levels normally used by the patients of the facility.

(5) Parking spaces are large enough and close enough to the facility to allow safe access by the physically impaired.

§ 485.64 Condition of participation: Disaster procedures.

The facility must have written policies and procedures that specifically define the handling of patients, personnel, records, and the public during disasters. All personnel associated with the facility must be knowledgeable with respect to these procedures, be trained in their application, and be assigned specific responsibilities.

(a) Standard: Disaster plan. The facility’s written disaster plan must be developed and maintained with assistance of qualified fire, safety, and other appropriate experts. The plan must include—

(1) Procedures for prompt transfer of casualties and records;

(2) Procedures for notifying community emergency personnel (for example, fire department, ambulance, etc.);

(3) Instructions regarding the location and use of alarm systems and signals and fire fighting equipment; and

(4) Specification of evacuation routes and procedures for leaving the facility.

(b) Standard: Drills and staff training. (1) The facility must provide ongoing training and drills for all personnel associated with the facility in all aspects of disaster preparedness.

(2) All new personnel must be oriented and assigned specific responsibilities regarding the facility’s disaster plan within two weeks of their first workday.

§ 485.66 Condition of participation: Utilization review plan.

The facility must have in effect a written utilization review plan that is implemented at least each quarter, to assess the necessity of services and promotes the most efficient use of services provided by the facility.

(a) Standard: Utilization review committee. The utilization review committee, consisting of the group of professional personnel specified in §485.56(c), a committee of this group, or
§ 485.70 Personnel qualifications.

This section sets forth the qualifications that must be met, as a condition of participation, under §485.58, and as a condition of coverage of services under §410.100 of this chapter.

(a) A facility physician must be a doctor of medicine or osteopathy who—

(1) Is licensed under State law to practice medicine or surgery; and

(2) Has had, subsequent to completing a 1-year hospital internship, at least 1 year of training in the medical management of patients requiring rehabilitation services; or

(3) Has had at least 1 year of full-time or part-time experience in a rehabilitation setting providing physicians' services similar to those required in this subpart.

(b) A licensed practical nurse must be licensed as a practical or vocational nurse by the State in which practicing, if applicable.

(c) An occupational therapist and an occupational therapist assistant must meet the qualifications set forth in §405.1202(f) and (g) of this chapter.

(d) An orthotist must—

(1) Be licensed by the State in which practicing, if applicable;

(2) Have successfully completed a training program in orthotics that is jointly recognized by the American Council on Education and the American Board for Certification in Orthotics and Prosthetics; and

(3) Be eligible to take that Board's certification examination in orthotics.

(e) A physical therapist and a physical therapist assistant must meet the qualifications set forth in paragraphs (b) and (c) of §485.705.

(f) A prosthetist must—

(1) Be licensed by the State in which practicing, if applicable;

(2) Have successfully completed a training program in prosthetics that is jointly recognized by the American Council on Education and the American Board for Certification in Orthotics and Prosthetics; and

(3) Be eligible to take that Board's certification examination in prosthetics.

(g) A psychologist must be certified or licensed by the State in which he or she is practicing, if that State requires certification or licensing, and must hold a masters degree in psychology from and educational institution approved by the State in which the institution is located.

(h) A registered nurse must be a graduate of an approved school of nursing and be licensed as a registered nurse by the State in which practicing, if applicable.

(i) A rehabilitation counselor must—

(1) Be licensed by the State in which practicing, if applicable;

(2) Hold at least a bachelor's degree; and

(3) Be eligible to take the certification examination administered by the Commission on Rehabilitation Counselor Certification.

(j) A respiratory therapist must—

(1) Be licensed by the State in which practicing, if applicable;

(2) Have successfully completed a training program accredited by the Committee on Allied Health Education and Accreditation (CAHEA) in collaboration with the Joint Review Committee for Respiratory Therapy Education; and

(3) Either—

(i) Be eligible to take the registry examination for respiratory therapists administered by the National Board for Respiratory Therapy, Inc.; or
§ 485.603 Rural health network.

A rural health network is an organization that meets the following specifications:

(a) It includes—
(1) At least one hospital that the State has designated or plans to designate as a CAH; and
(2) At least one hospital that furnishes acute care services.

(b) The members of the organization have entered into agreements regarding—
(1) Patient referral and transfer;
(2) The development and use of communications systems, including, where feasible, telemetry systems and systems for electronic sharing of patient data; and
(3) The provision of emergency and nonemergency transportation among members.

(c) Each CAH has an agreement with respect to credentialing and quality assurance with at least—
(1) One hospital that is a member of the network when applicable;
(2) One PRO or equivalent entity; or
(3) One other appropriate and qualified entity identified in the State rural health care plan.

§ 485.604 Personnel qualifications.

Staff that furnish services in a CAH must meet the applicable requirements of this section.

(a) Clinical nurse specialist. A clinical nurse specialist must be a person who performs the services of a clinical nurse specialist as authorized by the State, in accordance with State law or the State regulatory mechanism provided by State law.

(b) Nurse practitioner. A nurse practitioner must be a registered professional nurse who is currently licensed to practice in the State, who meets the State's requirements governing the qualification of nurse practitioners, and who meets one of the following conditions:

(1) Is currently certified as a primary care nurse practitioner by the American Nurses' Association or by the National Board of Pediatric Nurse Practitioners and Associates.

(2) Has satisfactorily completed a 1 academic year program that—

   (i) Prepares registered nurses to perform an expanded role in the delivery of primary care;

   (ii) Includes at least 4 months (in the aggregate) of classroom instruction and a component of supervised clinical practice; and

   (iii) Awards a degree, diploma, or certificate to persons who successfully complete the program.

(3) Has satisfactorily completed a formal educational program (for preparing physician assistants) that does not meet the requirements of paragraph (c)(2) of this section and has been assisting primary care physicians for a total of 12 months during the 18-month period immediately preceding June 25, 1993.

(c) Physician assistant. A physician assistant must be a person who meets the applicable State requirements governing the qualifications for assistants to primary care physicians, and who meets at least one of the following conditions:

(1) Is currently certified by the National Commission on Certification of Physician Assistants to assist primary care physicians.

(2) Has satisfactorily completed a program for preparing physician assistants that—

   (i) Was at least one academic year in length;

   (ii) Consisted of supervised clinical practice and at least 4 months (in the aggregate) of classroom instruction directed toward preparing students to deliver health care; and

   (iii) Was accredited by the American Medical Association's Committee on Allied Health Education and Accreditation.

(3) Has satisfactorily completed a formal educational program (for preparing physician assistants) that does not meet the requirements of paragraph (c)(2) of this section and has been assisting primary care physicians for a total of 12 months during the 18-month period immediately preceding June 25, 1993.

§ 485.606 Designation and certification of CAHs.

(a) Criteria for State designation. (1) A State that has established a Medicare rural hospital flexibility program described in section 1820(c) of the Act may designate one or more facilities as CAHs if each facility meets the CAH conditions of participation in this subpart F.

(2) The State must not deny any hospital that is otherwise eligible for designation as a CAH under this paragraph (a) solely because the hospital has entered into an agreement under which the hospital may provide posthospital SNF care as described in §482.66 of this chapter.

(b) Criteria for HCFA certification. HCFA certifies a facility as a CAH if—

(1) The facility is designated as a CAH by the State in which it is located and has been surveyed by the State survey agency or by HCFA and found to meet all conditions of participation in this Part and all other applicable requirements for participation in Part 489 of this chapter.

(2) The facility is a medical assistance facility operating in Montana or a rural primary care hospital designated by HCFA before August 5, 1997, and is otherwise eligible to be designated as a
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CAH by the State under the rules in this subpart.


§ 485.608 Condition of participation: Compliance with Federal, State, and local laws and regulations.

The CAH and its staff are in compliance with applicable Federal, State and local laws and regulations.

(a) Standard: Compliance with Federal laws and regulations. The CAH is in compliance with applicable Federal laws and regulations related to the health and safety of patients.

(b) Standard: Compliance with State and local laws and regulations. All patient care services are furnished in accordance with applicable State and local laws and regulations.

(c) Standard: Licensure of CAH. The CAH is licensed in accordance with applicable Federal, State and local laws and regulations.

(d) Standard: Licensure, certification or registration of personnel. Staff of the CAH are licensed, certified, or registered in accordance with applicable Federal, State, and local laws and regulations.


§ 485.610 Condition of participation: Status and location.

(a) Standard: Status. The facility is—

(1) A currently participating hospital that meets all conditions of participation set forth in this subpart;

(2) A recently closed facility, provided that the facility—

(i) Was a hospital that ceased operations on or after the date that is 10 years before November 29, 1999; and

(ii) Meets the criteria for designation under this subpart as of November 29, 1999; or

(3) A health clinic or a health center (as defined by the State) that—

(i) Is licensed by the State as a health clinic or a health center;

(ii) Was a hospital that was downsized to a health clinic or a health center; and

(iii) As of the effective date of its designation, meets the criteria for designation set forth in this subpart.

(b) Standard: Location. The CAH meets the following requirements:

(1) The CAH 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 the regulations in §412.62(f) of this chapter.

(2) The CAH is not deemed to be located in an urban area under §412.63(b) of this chapter.

(3) The CAH has not been classified as an urban hospital for purposes of the standardized payment amount by HCFA or the Medicare Geographic Classification Review Board under §412.230(e) of this chapter, and is not among a group of hospitals that have been redesignated to an adjacent urban area under §412.232 of this chapter.

(4) The CAH is being treated as being located in a rural area in accordance with §412.103 of this chapter.

(5) The CAH is located more than a 35-mile drive (or, in the case of mountainous terrain or in areas with only secondary roads available, a 15-mile drive) from a hospital or another CAH, or the CAH is certified by the State as being a necessary provider of health care services to residents in the area.


§ 485.612 Condition of participation: Compliance with hospital requirements at time of application.

The hospital has a provider agreement to participate in the Medicare program as a hospital at the time the hospital applies for designation as a CAH.


§ 485.616 Condition of participation: Agreements.

(a) Standard: Agreements with network hospitals. In the case of a CAH that is a member of a rural health network as defined in §485.603 of this chapter, the CAH has in effect an agreement with at least one hospital that is a member of the network for—

(1) Patient referral and transfer;

(2) The development and use of communications systems of the network, including the network’s system for the electronic sharing of patient data, and telemetry and medical records, if the
§ 485.618 Condition of participation: Emergency services.

The CAH provides emergency care necessary to meet the needs of its inpatients and outpatients.

(a) Standard: Availability. Emergency services are available on a 24-hours a day basis.

(b) Standard: Equipment, supplies, and medication. Equipment, supplies, and medication used in treating emergency cases are kept at the CAH and are readily available for treating emergency cases. The items available must include the following:

1. Drugs and biologicals commonly used in life-saving procedures, including analgesics, local anesthetics, antibiotics, anticonvulsants, antidotes and emetics, serums and toxoids, antiarrhythmics, cardiac glycosides, antihypertensives, diuretics, and electrolytes and replacement solutions.

2. Equipment and supplies commonly used in life-saving procedures, including airways, endotracheal tubes, ambu bag/valve/mask, oxygen, tourniquets, immobilization devices, nasogastric tubes, splints, IV therapy supplies, suction machine, defibrillator, cardiac monitor, chest tubes, and indwelling urinary catheters.

(c) Standard: Blood and blood products. The facility provides, either directly or under arrangements, the following:

1. Services for the procurement, safekeeping, and transfusion of blood, including the availability of blood products needed for emergencies on a 24-hours a day basis.

2. Blood storage facilities that meet the requirements of 42 CFR part 493, subpart K, and are under the control and supervision of a pathologist or other qualified doctor of medicine or osteopathy. If blood banking services are provided under an arrangement, the arrangement is approved by the facility’s medical staff and by the persons directly responsible for the operation of the facility.

(d) Standard: Personnel. There must be a doctor of medicine or osteopathy, a physician assistant, or a nurse practitioner with training or experience in emergency care on call and immediately available by telephone or radio contact, and available on site within the following timeframes:

1. Within 30 minutes, on a 24-hour a day basis, if the CAH is located in an area other than an area described in paragraph (d)(2) of this section; or

2. Within 60 minutes, on a 24-hour a day basis, if all of the following requirements are met:

   (i) The CAH is located in an area designated as a frontier area (that is, an area with fewer than six residents per square mile based on the latest population data published by the Bureau of the Census) or in an area that meets criteria for a remote location adopted by the State in its rural health care plan, and approved by HCFA, under section 1820(b) of the Act.

   (ii) The State has determined under criteria in its rural health care plan that allowing an emergency response time longer than 30 minutes is the only feasible method of providing emergency care to residents of the area served by the CAH.

   (iii) The State maintains documentation showing that the response time of up to 60 minutes at a particular CAH designated is justified because other available alternatives would increase the time needed to stabilize a patient in an emergency.

(e) Standard: Coordination with emergency response systems. The CAH must, in coordination with emergency response systems in the area, establish procedures under which a doctor of medicine or osteopathy is immediately available by telephone or radio contact on a 24-hours a day basis to receive emergency calls, provide information
§ 485.620 Condition of participation: Number of beds and length of stay.

(a) Standard: Number of beds. Except as permitted for CAHs having swing-bed agreements under §485.645 of this chapter, the CAH maintains no more than 15 inpatient beds.

(b) Standard: Length of stay. The CAH provides acute inpatient care for a period that does not exceed, on an annual average basis, 96 hours per patient.

§ 485.623 Condition of participation: Physical plant and environment.

(a) Standard: Construction. The CAH is constructed, arranged, and maintained to ensure access to and safety of patients, and provides adequate space for the provision of direct services.

(b) Standard: Maintenance. The CAH has housekeeping and preventive maintenance programs to ensure that—

(1) All essential mechanical, electrical, and patient-care equipment is maintained in safe operating condition;

(2) There is proper routine storage and prompt disposal of trash;

(3) Drugs and biologicals are appropriately stored;

(4) The premises are clean and orderly; and

(5) There is proper ventilation, lighting, and temperature control in all pharmaceutical, patient care, and food preparation areas.

(c) Standard: Emergency procedures. The CAH assures the safety of patients in non-medical emergencies by—

(1) Training staff in handling emergencies, including prompt reporting of fires, extinguishing of fires, protection and, where necessary, evacuation of patients, personnel, and guests, and cooperation with fire fighting and disaster authorities;

(2) Providing for emergency power and lighting in the emergency room and for battery lamps and flashlights in other areas;

(3) Providing for an emergency fuel and water supply; and

(4) Taking other appropriate measures that are consistent with the particular conditions of the area in which the CAH is located.

(d) Standard: Life safety from fire—(1) Except as provided in paragraphs (d)(2) and (d)(3) of this section, the CAH must meet the requirements of chapter 12, New Health Care Occupancy, or chapter 13, Existing Health Care Occupancy, of the 1985 edition of the Life Safety Code of the National Fire Protection Association. Incorporation by reference of the 1985 edition of the National Fire Protection Association’s Life Safety Code (published February 7, 1985; ANSI/NFPA 101) was approved by the Director of the Federal Register in accordance with 5 U.S.C. 552(a) and 1 CFR part 51. The Code is available for inspection at the HCFA Information Resource Center, 7500 Security Boulevard, Room C2-07-13, Central Building, Baltimore, MD 21244-1850, and the Office of the Federal Register, 800 North Capitol Street, NW., suite 700, Washington, DC. Copies may be obtained from the National Fire Protection Association, Batterymarch Park, Quincy, Mass. 02269. If any changes in this code are also to be incorporated by reference, a document to that effect will be published in the FEDERAL REGISTER.

(2) Any CAH that as a hospital on or before November 26, 1982, complied, with or without waivers, with the requirements of the 1967 edition of the Life Safety Code, or after November 26, 1982 and on or before May 9, 1988, complied with the 1981 edition of the Life Safety Code, is considered to be in compliance with this standard as long as the CAH continues to remain in compliance with that edition of the Code. The 1967 and 1981 Life Safety Codes are available for inspection at the HCFA Information Resource Center, 7500 Security Boulevard, Room C2-07-13, Central Building, Baltimore, MD 21244-1850.

(3) After consideration of State survey agency findings, HCFA may waive specific provisions of the Life Safety Code that, if rigidly applied, would result in unreasonable hardship on the CAH, but only if the waiver does not
§ 485.627 Condition of participation: Organizational structure.

(a) Standard: Governing body or responsible individual. The CAH has a governing body or an individual that assumes full legal responsibility for determining, implementing and monitoring policies governing the CAH's total operation and for ensuring that those policies are administered so as to provide quality health care in a safe environment.

(b) Standard: Disclosure. The CAH discloses the names and addresses of—

(1) Its owners, or those with a controlling interest in the CAH or in any subcontractor in which the CAH directly or indirectly has a 5 percent or more ownership interest, in accordance with subpart C of part 420 of this chapter;

(2) The person principally responsible for the operation of the CAH; and

(3) The person responsible for medical direction.

§ 485.631 Condition of participation: Staffing and staff responsibilities.

(a) Standard: Staffing—(1) The CAH has a professional health care staff that includes one or more doctors of medicine or osteopathy, and may include one or more physician assistants, nurse practitioners, or clinical nurse specialists.

(2) Any ancillary personnel are supervised by the professional staff.

(3) The staff is sufficient to provide the services essential to the operation of the CAH.

(4) A doctor of medicine or osteopathy, nurse practitioner, clinical nurse specialist, or physician assistant is available to furnish patient care services at all times the CAH operates.

(5) A registered nurse, clinical nurse specialist, or licensed practical nurse is on duty whenever the CAH has one or more inpatients.

(b) Standard: Responsibilities of the doctor of medicine or osteopathy—

(i) Provides medical direction for the CAH's health care activities and consultation for, and medical supervision of, the health care staff;

(ii) In conjunction with the physician assistant and/or nurse practitioner member(s), participates in developing, executing, and periodically reviewing the CAH's written policies governing the services it furnishes;

(iii) In conjunction with the physician assistant and/or nurse practitioner members, periodically reviews the CAH's patient records, provides medical orders, and provides medical care services to the patients of the CAH; and

(iv) Periodically reviews and signs the records of patients cared for by nurse practitioners, clinical nurse specialists, or physician assistants.

(2) A doctor of medicine or osteopathy is present for sufficient periods of time, at least once in every 2 week period (except in extraordinary circumstances) to provide the medical direction, medical care services, consultation, and supervision described in this paragraph, and is available through direct radio or telephone communication for consultation, assistance with medical emergencies, or patient referral. The extraordinary circumstances are documented in the records of the CAH. A site visit is not required if no patients have been treated since the latest site visit.

(c) Standard: Physician assistant, nurse practitioner, and clinical nurse specialist responsibilities.

(1) The physician assistant, the nurse practitioner, or clinical nurse specialist members of the CAH's staff—

(i) Participate in the development, execution and periodic review of the written policies governing the services the CAH furnishes; and

(ii) Participate with a doctor of medicine or osteopathy in a periodic review of the patients' health records.

(2) The physician assistant, nurse practitioner, or clinical nurse specialist performs the following functions to the extent they are not being

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performed by a doctor of medicine or osteopathy:

(i) Provides services in accordance with the CAH’s policies.

(ii) Arranges for, or refers patients to, needed services that cannot be furnished at the CAH, and assures that adequate patient health records are maintained and transferred as required when patients are referred.

(3) Whenever a patient is admitted to the CAH by a nurse practitioner, physician assistant, or clinical nurse specialist, a doctor of medicine or osteopathy on the staff of the CAH is notified of the admission.


§ 485.635 Condition of participation: Provision of services.

(a) Standard: Patient care policies. (1) The CAH’s health care services are furnished in accordance with appropriate written policies that are consistent with applicable State law.

(2) The policies are developed with the advice of a group of professional personnel that includes one or more doctors of medicine or osteopathy and one or more physician assistants, nurse practitioners, or clinical nurse specialists, if they are on staff under the provisions of § 485.631(a)(1); at least one member is not a member of the CAH staff.

(3) The policies include the following:

(i) A description of the services the CAH furnishes directly and those furnished through agreement or arrangement.

(ii) Policies and procedures for emergency medical services.

(iii) Guidelines for the medical management of health problems that include the conditions requiring medical consultation and/or patient referral, the maintenance of health care records, and procedures for the periodic review and evaluation of the services furnished by the CAH.

(iv) Rules for the storage, handling, dispensation, and administration of drugs and biologicals. These rules must provide that there is a drug storage area that is administered in accordance with accepted professional principles, that current and accurate records are kept of the receipt and disposition of all scheduled drugs, and that outdated, mislabeled, or otherwise unusable drugs are not available for patient use.

(v) Procedures for reporting adverse drug reactions and errors in the administration of drugs.

(vi) A system for identifying, reporting, investigating and controlling infections and communicable diseases of patients and personnel.

(vii) If the CAH furnishes inpatient services, procedures that ensure that the nutritional needs of inpatients are met in accordance with recognized dietary practices and the orders of the practitioner responsible for the care of the patients, and that the requirement of § 483.25(i) is met with respect to inpatients receiving posthospital SNF care.

(4) These policies are reviewed at least annually by the group of professional personnel required under paragraph (a)(2) of this section, and reviewed as necessary by the CAH.

(b) Standard: Direct services—(1) General. The CAH staff furnishes, as direct services, those diagnostic and therapeutic services and supplies that are commonly furnished in a physician’s office or at another entry point into the health care delivery system, such as a low intensity hospital outpatient department or emergency department. These direct services include medical history, physical examination, specimen collection, assessment of health status, and treatment for a variety of medical conditions.

(2) Laboratory services. The CAH provides, as direct services, basic laboratory services essential to the immediate diagnosis and treatment of the patient that meet the standards imposed under section 353 of the Public Health Service Act (42 U.S.C. 236a). (See the laboratory requirements specified in part 493 of this chapter.) The services provided include:

(i) Chemical examination of urine by stick or tablet method or both (including urine ketones);

(ii) Hemoglobin or hematocrit;

(iii) Blood glucose;

(iv) Examination of stool specimens for occult blood;

(v) Pregnancy tests; and

(vi) Primary culturing for transmittal to a certified laboratory.
§ 485.638 Conditions of participation: Clinical records.

(a) Standard: Records system.—(1) The CAH maintains a clinical records system in accordance with written policies and procedures.

(2) The records are legible, complete, accurately documented, readily accessible, and systematically organized.

(3) A designated member of the professional staff is responsible for maintaining the records and for ensuring that they are completely and accurately documented, readily accessible, and systematically organized.

(4) For each patient receiving health care services, the CAH maintains a record that includes, as applicable—

(i) Identification and social data, evidence of properly executed informed consent forms, pertinent medical history, assessment of the health status and health care needs of the patient, and a brief summary of the episode, disposition, and instructions to the patient;

(ii) Reports of physical examinations, diagnostic and laboratory test results, including clinical laboratory services, and consultative findings;

(iii) All orders of doctors of medicine or osteopathy or other practitioners,

(iv) Inpatient hospital care;

(v) Services of doctors of medicine or osteopathy; and

(vi) Additional or specialized diagnostic and clinical laboratory services that are not available at the CAH.

(5) The person principally responsible for the operation of the CAH under § 485.627(b)(2) of this chapter is also responsible for the following:

(i) Services furnished in the CAH whether or not they are furnished under arrangements or agreements.

(ii) Ensuring that a contractor of services (including one for shared services and joint ventures) furnishes services that enable the CAH to comply with all applicable conditions of participation and standards for the contracted services.

(d) Standard: Nursing services. Nursing services must meet the needs of patients.

(1) A registered nurse must provide (or assign to other personnel) the nursing care of each patient, including patients at a SNF level of care in a swing-bed CAH. The care must be provided in accordance with the patient’s needs and the specialized qualifications and competence of the staff available.

(2) A registered nurse or, where permitted by State law, a physician assistant, must supervise and evaluate the nursing care for each patient, including patients at a SNF level of care in a swing-bed CAH.

(3) All drugs, biologicals, and intravenous medications must be administered by or under the supervision of a registered nurse, a doctor of medicine or osteopathy, or, where permitted by State law, a physician assistant, in accordance with written and signed orders, accepted standards of practice, and Federal and State laws.

(4) A nursing care plan must be developed and kept current for each inpatient.
§ 485.641 Condition of participation: Periodic evaluation and quality assurance review.

(a) Standard: Periodic evaluation—(1) The CAH carries out or arranges for a periodic evaluation of its total program. The evaluation is done at least once a year and includes review of—

(i) A qualified anesthesiologist;

(ii) A doctor of medicine or osteopathy other than an anesthesiologist, including an osteopathic practitioner recognized under section 1101(a)(7) of the Act;

(iii) A doctor of dental surgery or dental medicine;

(iv) A doctor of podiatric medicine;

(v) A certified registered nurse anesthetist, as defined in §410.69(b) of this chapter;

(vi) An anesthesiologist’s assistant, as defined in §410.69(b) of this chapter; or

(vii) A supervised trainee in an approved educational program, as described in §§413.85 or 413.86 of this chapter.

(2) In those cases in which a certified registered nurse anesthetist administers the anesthesia, the anesthetist must be under the supervision of the operating practitioner. An anesthesiologist’s assistant who administers anesthesia must be under the supervision of an anesthesiologist.

(d) Discharge. All patients are discharged in the company of a responsible adult, except those exempted by the practitioner who performed the surgical procedure.


§ 485.639 Condition of participation: Surgical services.

Surgical procedures must be performed in a safe manner by qualified practitioners who have been granted clinical privileges by the governing body of the CAH in accordance with the designation requirements under paragraph (a) of this section.

(a) Designation of qualified practitioners. The CAH designates the practitioners who are allowed to perform surgery for CAH patients, in accordance with its approved policies and procedures, and with State scope of practice laws. Surgery is performed only by—

(1) A doctor of medicine or osteopathy, including an osteopathic practitioner recognized under section 1101(a)(7) of the Act;

(2) A doctor of dental surgery or dental medicine; or

(3) A doctor of podiatric medicine.

(b) Anesthetic risk and evaluation. A qualified practitioner, as described in paragraph (a) of this section, must examine the patient immediately before surgery to evaluate the risk of anesthesia and of the procedure to be performed. Before discharge from the CAH, each patient must be evaluated for proper anesthesia recovery by a qualified practitioner as described in paragraph (a) of this section.

(c) Administration of anesthesia. The CAH designates the person who is allowed to administer anesthesia to CAH patients in accordance with its approved policies and procedures and with State scope of practice laws.

(1) Anesthetics must be administered only by—

(i) A qualified anesthesiologist;

(ii) A doctor of medicine or osteopathy other than an anesthesiologist, including an osteopathic practitioner recognized under section 1101(a)(7) of the Act;

(iii) A doctor of dental surgery or dental medicine;

(iv) A doctor of podiatric medicine;

(v) A certified registered nurse anesthetist, as defined in §410.69(b) of this chapter;

(vi) An anesthesiologist’s assistant, as defined in §410.69(b) of this chapter; or

(vii) A supervised trainee in an approved educational program, as described in §§413.85 or 413.86 of this chapter.

§ 485.643 Condition of participation: Organ, tissue, and eye procurement.

The CAH must have and implement written protocols that:

(a) Incorporate an agreement with an OPO designated under part 486 of this chapter, under which it must notify, in a timely manner, the OPO or a third party designated by the OPO of individuals whose death is imminent or who have died in the CAH. The OPO determines medical suitability for organ donation and, in the absence of alternative arrangements by the CAH, the OPO determines medical suitability for tissue and eye donation, using the definition of potential tissue and eye donor and the notification protocol developed in consultation with the tissue and eye banks identified by the CAH for this purpose;

(b) Incorporate an agreement with at least one tissue bank and at least one eye bank to cooperate in the retrieval, processing, preservation, storage and distribution of tissues and eyes, as may be appropriate to assure that all usable tissues and eyes are obtained from potential donors, insofar as such an agreement does not interfere with organ procurement;

(c) Ensure, in collaboration with the designated OPO, that the family of each potential donor is informed of its option to either donate or not donate organs, tissues, or eyes. The individual designated by the CAH to initiate the request to the family must be a designated requestor. A designated requestor is an individual who has completed a course offered or approved by the OPO and designed in conjunction with the tissue and eye bank community in the methodology for approaching potential donor families and requesting organ or tissue donation;

(d) Encourage discretion and sensitivity with respect to the circumstances, views, and beliefs of the families of potential donors;

(e) Ensure that the CAH works cooperatively with the designated OPO, tissue bank and eye bank in educating staff on donation issues, reviewing death records to improve identification of potential donors, and maintaining potential donors while necessary testing and placement of potential donated organs, tissues, and eyes take place.

§ 485.643 Organ, tissue, and eye procurement.

The CAH must have and implement written protocols that:

(i) The utilization of CAH services, including at least the number of patients served and the volume of services;

(ii) A representative sample of both active and closed clinical records; and

(iii) The CAH's health care policies.

(2) The purpose of the evaluation is to determine whether the utilization of services was appropriate, the established policies were followed, and any changes are needed.

(b) Standard: Quality assurance. The CAH has an effective quality assurance program to evaluate the quality and appropriateness of the diagnosis and treatment furnished in the CAH and of the treatment outcomes. The program requires that—

(1) All patient care services and other services affecting patient health and safety, are evaluated;

(2) Nosocomial infections and medication therapy are evaluated;

(3) The quality and appropriateness of the diagnosis and treatment furnished by nurse practitioners, clinical nurse specialists, and physician assistants at the CAH are evaluated by a member of the CAH staff who is a doctor of medicine or osteopathy or by another doctor of medicine or osteopathy under contract with the CAH;

(4) The quality and appropriateness of the diagnosis and treatment furnished by doctors of medicine or osteopathy at the CAH are evaluated by—

(i) One hospital that is a member of the network, when applicable;

(ii) One PRO or equivalent entity; or

(iii) One other appropriate and qualified entity identified in the State rural health care plan; and

(5)(i) The CAH staff considers the findings of the evaluations, including any findings or recommendations of the PRO, and takes corrective action if necessary.

(ii) The CAH also takes appropriate remedial action to address deficiencies found through the quality assurance program.

(iii) The CAH documents the outcome of all remedial action.

(f) For purposes of these standards, the term "Organ" means a human kidney, liver, heart, lung, or pancreas.

§ 485.645 Special requirements for CAH providers of long-term care services ("swing-beds")

A CAH must meet the following requirements in order to be granted an approval from HCFA to provide post-hospital SNF care, as specified in § 409.30 of this chapter, and to be paid for SNF-level services, in accordance with paragraph (c) of this section.

(a) Eligibility. A CAH must meet the following eligibility requirements:

(1) The facility has been certified as a CAH by HCFA under § 485.606(b) of this subpart; and

(2) The facility provides not more than 25 inpatient beds, and the number of beds used at any time for acute care inpatient services does not exceed 15 beds. Any bed of a unit of the facility that is licensed as distinct-part SNF at the time the facility applies to the State for designation as a CAH is not counted under paragraph (a) of this section.

(b) Facilities participating as rural primary care hospitals (R P C H s) on September 30, 1997. These facilities must meet the following requirements:

(1) Notwithstanding paragraph (a) of this section, a CAH that participated in Medicare as a R P C H on September 30, 1997, and on that date had in effect an approval from HCFA to use its inpatient facilities to provide post-hospital SNF care may continue in that status under the same terms, conditions and limitations that were applicable at the time those approvals were granted.

(2) A CAH that was granted swing-bed approval under paragraph (b)(1) of this section may request that its application to be a CAH and swing-bed provider be reevaluated under paragraph (a) of this section. If this request is approved, the approval is effective not earlier than October 1, 1997. As of the date of approval, the CAH no longer has any status under paragraph (b)(1) of this section and may not request reinstatement under paragraph (b)(1) of this section.

(c) Payment. Payment for inpatient RPCH services to a CAH that has qualified as a CAH under the provisions in paragraph (a) of this section is made in accordance with § 413.70 of this chapter. Payment for post-hospital SNF-level of care services is made in accordance with the payment provisions in § 413.114 of this chapter.

(d) SNF services. The CAH is substantially in compliance with the following SNF requirements contained in subpart B of part 483 of this chapter:

(1) Residents rights (§ 483.10(b)(3) through (b)(6), (d) (e), (h), (l), (j)(1)(vii) and (viii), (l), and (m) of this chapter).

(2) Admission, transfer, and discharge rights (§ 483.12(a) of this chapter).

(3) Resident behavior and facility practices (§ 483.13 of this chapter).

(4) Patient activities (§ 483.15(f) of this chapter), except that the services may be directed either by a qualified professional meeting the requirements of § 483.15(f)(2), or by an individual on the facility staff who is designated as the activities director and who serves in consultation with a therapeutic recreation specialist, occupational therapist, or other professional with experience or education in recreational therapy.

(5) Social services (§ 483.15(g) of this chapter).

(6) Comprehensive assessment, comprehensive care plan, and discharge planning (§ 483.20(b), (d), and (e) of this chapter, except that the CAH is not required to comply with the requirements for frequency, scope and number of assessments prescribed in § 413.343(b)).

(7) Specialized rehabilitative services (§ 483.45 of this chapter).

(8) Dental services (§ 483.55 of this chapter).

(9) Nutrition (§ 483.25(i) of this chapter).

[63 FR 26359, May 12, 1998 as amended at 64 FR 41544, July 30, 1999]
§ 485.701 Basis and scope.

This subpart implements section 1861(p)(4) of the Act, which—
(a) Defines outpatient physical therapy and speech pathology services;
(b) Imposes requirements with respect to adequate program, facilities, policies, staffing, and clinical records; and
(c) Authorizes the Secretary to establish by regulation other health and safety requirements.
[60 FR 2327, Jan. 9, 1995]

§ 485.703 Definitions.

Clinic. A facility that is established primarily to furnish outpatient physician services and that meets the following tests of physician involvement:
(1) The medical services are furnished by a group of three or more physicians practicing medicine together.
(2) A physician is present during all hours of operation of the clinic to furnish medical services, as distinguished from purely administrative services.

Organization. A clinic, rehabilitation agency, or public health agency.

Public health agency. An official agency established by a State or local government, the primary function of which is to maintain the health of the population served by performing environmental health services, preventive medical services, and in certain cases, therapeutic services.

Rehabilitation agency. An agency that—
(1) Provides an integrated multidisciplinary rehabilitation program designed to upgrade the physical functioning of handicapped disabled individuals by bringing specialized rehabilitation staff together to perform as a team; and
(2) Provides at least the following services:
(i) Physical therapy or speech-language pathology services.
(ii) Social or vocational adjustment services.

Supervision. Authoritative procedural guidance that is for the accomplishment of a function or activity and that—
(1) Includes initial direction and periodic observation of the actual performance of the function or activity; and
(2) Is furnished by a qualified person—
(i) Whose sphere of competence encompasses the particular function or activity; and
(ii) Who (unless otherwise provided in this subpart) is on the premises if the person performing the function or activity does not meet the assistant-level practitioner qualifications specified in §485.705.

§ 485.705 Personnel qualifications.

(a) General qualification requirements. Except as specified in paragraphs (b) and (c) of this section, all personnel who are involved in the furnishing of outpatient physical therapy, occupational therapy, and speech-language pathology services directly by or under arrangements with an organization must be legally authorized (licensed or, if applicable, certified or registered) to practice by the State in which they perform the functions or actions, and must act only within the scope of their State license or State certification or registration.

(b) Exception for Federally defined qualifications. The following Federally defined qualifications must be met:
(1) For a physician, the qualifications and conditions as defined in section 1861(r) of the Act and the requirements in part 484 of this chapter.
(2) For a speech-language pathologist, the qualifications specified in section 1861(11)(1) of the Act and the requirements in part 484 of this chapter.
(c) Exceptions when no State licensing laws or State certification or registration requirements exist. If no State licensing
laws or State certification or registration requirements exist for the profession, the following requirements must be met—

(1) An administrator is a person who has a bachelor's degree and:
   (i) has experience or specialized training in the administration of health institutions or agencies or;
   (ii) is qualified and has experience in one of the professional health disciplines.

(2) An occupational therapist must meet the requirements in part 484 of this chapter.

(3) An occupational therapy assistant must meet the requirements in part 484 of this chapter.

(4) A physical therapist must meet the requirements in part 484 of this chapter.

(5) A physical therapist assistant must meet the requirements in part 484 of this chapter.

(6) A social worker must meet the requirements in part 484 of this chapter.

(7) A vocational specialist is a person who has a baccalaureate degree and—
   (i) Two years experience in vocational counseling in a rehabilitation setting such as a sheltered workshop, State employment service agency, etc.; or
   (ii) At least 18 semester hours in vocational rehabilitation, educational or vocational guidance, psychology, social work, special education or personnel administration, and 1 year of experience in vocational counseling in a rehabilitation setting; or
   (iii) A master's degree in vocational counseling.

(8) A nurse practitioner is a person who must:
   (i) 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
   (ii) Be certified as a nurse practitioner by December 31, 2000; or
   (iv) Be a nurse practitioner who on or after January 1, 2001, applies for a Medicare billing number for the first time and meets the standards for nurse practitioners in paragraphs (c)(8)(i) and (c)(8)(ii) of this section; or
   (v) 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.

(9) A clinical nurse specialist is a person who must:
   (i) 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; or
   (ii) Have a master's degree in a defined clinical area of nursing from an accredited educational institution; and
   (iii) Be certified as a clinical nurse specialist by the American Nurses Credentialing Center.

(10) A physician assistant is a person who:
   (i) Has graduated from a physician assistant educational program that is accredited by the Commission on Accreditation of Allied Health Education Programs; or
   (ii) Has passed the national certification examination that is administered by the National Commission on Certification of Physician Assistants; and
   (iii) Is licensed by the State to practice as a physician assistant.

§ 485.707 Condition of participation: Compliance with Federal, State, and local laws.

The organization and its staff are in compliance with all applicable Federal, State, and local laws and regulations.

(a) Standard: Licensure of organization. In any State in which State or applicable local law provides for the licensing of organizations, a clinic, rehabilitation agency, or public health agency is
licensed in accordance with applicable laws.

(b) Standard: Licensure or registration of personnel. Staff of the organization are licensed or registered in accordance with applicable laws.

§ 485.709 Condition of participation: Administrative management.

The clinic or rehabilitation agency has an effective governing body that is legally responsible for the conduct of the clinic or rehabilitation agency. The governing body designates an administrator, and establishes administrative policies.

(a) Standard: Governing body. There is a governing body (or designated person(s) so functioning) which assumes full legal responsibility for the overall conduct of the clinic or rehabilitation agency and for compliance with applicable laws and regulations. The name of the owner(s) of the clinic or rehabilitation agency is fully disclosed to the State agency. In the case of corporations, the names of the corporate officers are made known.

(b) Standard: Administrator. The governing body—

(1) Appoints a qualified full-time administrator;
(2) Delegates to the administrator the internal operation of the clinic or rehabilitation agency in accordance with written policies;
(3) Defines clearly the administrator’s responsibilities for procurement and direction of personnel; and
(4) Designates a competent individual to act during temporary absence of the administrator.

(c) Standard: Personnel policies. Personnel practices are supported by appropriate written personnel policies that are kept current. Personnel records include the qualifications of all professional and assistant level personnel, as well as evidence of State licensure if applicable.

(d) Standard: Patient care policies. Patient care practices and procedures are supported by written policies established by a group of professional personnel including one or more physicians associated with the clinic or rehabilitation agency, one or more qualified physical therapists (if physical therapy services are provided), and one or more qualified speech pathologists (if speech pathology services are provided). The policies govern the outpatient physical therapy and/or speech pathology services and related services that are provided. These policies are evaluated at least annually by the group of professional personnel, and revised as necessary based upon this evaluation.

§ 485.711 Condition of participation: Plan of care and physician involvement.

For each patient in need of outpatient physical therapy or speech pathology services there is a written plan of care established and periodically reviewed by a physician, or by a physical therapist or speech pathologist respectively. The organization has a physician available to furnish necessary medical care in case of emergency.

(a) Standard: Medical history and prior treatment. The following are obtained by the organization before or at the time of initiation of treatment:

(1) The patient’s significant past history.
(2) Current medical findings, if any.
(3) Diagnosis(es), if established.
(4) Physician’s orders, if any.
(5) Rehabilitation goals, if determined.
(6) Contraindications, if any.
(7) The extent to which the patient is aware of the diagnosis(es) and prognosis.
(8) If appropriate, the summary of treatment furnished and results achieved during previous periods of rehabilitation services or institutionalization.

(b) Standard: Plan of care. (1) For each patient there is a written plan of care established by the physician or by the physical therapist or speech-language pathologist who furnishes the services.
(2) The plan of care for physical therapy or speech pathology services indicates anticipated goals and specifies for those services the—
   (i) Type;
   (ii) Amount;
   (iii) Frequency; and
   (iv) Duration.

(3) The plan of care and results of treatment are reviewed by the physician or by the individual who established the plan at least as often as the patient’s condition requires, and the indicated action is taken. (For Medicare patients, the plan must be reviewed by a physician, nurse practitioner, clinical nurse specialist, or physician assistant at least every 30 days, in accordance with §410.61(e) of this chapter.)

(4) Changes in the plan of care are noted in the clinical record. If the patient has an attending physician, the therapist or speech-language pathologist who furnishes the services promptly notifies him or her of any change in the patient’s condition or in the plan of care.

(c) Standard: Emergency care. The organization provides for one or more doctors of medicine or osteopathy to be available on call to furnish necessary medical care in case of emergency. The established procedures to be followed by personnel in an emergency cover immediate care of the patient, persons to be notified, and reports to be prepared.

§ 485.713 Condition of participation: Physical therapy services

If the organization offers physical therapy services, it provides an adequate program of physical therapy and has an adequate number of qualified personnel and the equipment necessary to carry out its program and to fulfill its objectives.

(a) Standard: Adequate program. (1) The organization is considered to have an adequate outpatient physical therapy program if it can:
   (i) Provide services using therapeutic exercise and the modalities of heat, cold, water, and electricity; (ii) Conduct patient evaluations; and (iii) Administer tests and measurements of strength, balance, endurance, range of motion, and activities of daily living.

(2) A qualified physical therapist is present or readily available to offer supervision when a physical therapist assistant furnishes services.
   (i) If a qualified physical therapist is not on the premises during all hours of operation, patients are scheduled so as to ensure that the therapist is present when special skills are needed, for example, for evaluation and reevaluation.
   (ii) When a physical therapist assistant furnishes services off the organization’s premises, those services are supervised by a qualified physical therapist who makes an onsite supervisory visit at least once every 30 days.

(b) Standard: Facilities and equipment. The organization has the equipment and facilities required to provide the range of services necessary in the treatment of the types of disabilities it accepts for service.

(c) Standard: Personnel qualified to provide physical therapy services. Physical therapy services are provided by, or under the supervision of, a qualified physical therapist. The number of qualified physical therapists and qualified physical therapist assistants is adequate for the volume and diversity of physical therapy services offered. A qualified physical therapist is on the premises or readily available during the operating hours of the organization.

(d) Standard: Supportive personnel. If personnel are available to assist qualified physical therapists by performing services incident to physical therapy that do not require professional knowledge and skill, these personnel are instructed in appropriate patient care services by qualified physical therapists who retain responsibility for the treatment prescribed by the attending physician.
§ 485.715 Condition of participation: Speech pathology services.

If speech pathology services are offered, the organization provides an adequate program of speech pathology and has an adequate number of qualified personnel and the equipment necessary to carry out its program and to fulfill its objectives.

(a) Standard: Adequate program. The organization is considered to have an adequate outpatient speech pathology program if it can provide the diagnostic and treatment services to effectively treat speech disorders.

(b) Standard: Facilities and equipment. The organization has the equipment and facilities required to provide the range of services necessary in the treatment of the types of speech disorders it accepts for service.

(c) Standard: Personnel qualified to provide speech pathology services. Speech pathology services are given or supervised by a qualified speech pathologist and the number of qualified speech pathologists is adequate for the volume and diversity of speech pathology services offered. At least one qualified speech pathologist is present at all times when speech pathology services are furnished.


§ 485.717 Condition of participation: Rehabilitation program.

This condition and its standards apply only to a rehabilitation agency's own patients, not to patients of hospitals, skilled nursing facilities (SNFs), or Medicaid nursing facilities (NFs) to whom the agency furnishes services. (The hospital, SNF, or NF is responsible for ensuring that qualified staff furnish services for which they arrange or contract for their patients.) The rehabilitation agency provides, in addition to physical therapy and speech-language pathology services, social or vocational adjustment services to all of its patients who need them. The agency provides for special qualified staff to evaluate the social and vocational factors, to counsel and advise on the social or vocational problems that arise from the patient's illness or injury, and to make appropriate referrals for needed services.

(a) Standard: Qualification of staff. The agency's social or vocational adjustment services are furnished as appropriate, by qualified psychologists, qualified social workers, or qualified vocational specialists. Social or vocational adjustment services may be performed by a qualified psychologist or qualified social worker. Vocational adjustment services may be furnished by a qualified vocational specialist.

(b) Standard: Arrangements for social or vocational adjustment services. (1) If a rehabilitation agency does not provide social or vocational adjustment services through salaried employees, it may provide those services through a written contract with others who meet the requirements and responsibilities set forth in this subpart for salaried personnel.

(2) The contract must specify the term of the contract and the manner of termination or renewal and provide that the agency retains responsibility for the control and supervision of the services.


§ 485.719 Condition of participation: Arrangements for physical therapy and speech pathology services to be performed by other than salaried organization personnel.

(a) Conditions. If an organization provides outpatient physical therapy or speech pathology services under an arrangement with others, the services are to be furnished in accordance with the terms of a written contract, which provides that the organization retains professional and administrative responsibility for, and control and supervision of, the services.

(b) Standard: Contract provisions. The contract—

(1) Specifies the term of the contract and the manner of termination or renewal;

(2) Requires that personnel who furnish the services meet the requirements that are set forth in this subpart for salaried personnel; and
§ 485.723 Condition of participation: Clinical records.

The organization maintains clinical records on all patients in accordance with accepted professional standards, and practices. The clinical records are completely and accurately documented, readily accessible, and systematically organized to facilitate retrieving and compiling information.

(a) Standard: Protection of clinical record information. The organization recognizes the confidentiality of clinical record information and provides safeguards against loss, destruction, or unauthorized use. Written procedures govern the use and removal of records and the conditions for release of information. The patient's written consent is required for release of information not authorized by law.

(b) Standard: Content. The clinical record contains sufficient information to identify the patient clearly, to justify the diagnosis(es) and treatment, and to document the results accurately. All clinical records contain the following general categories of data:

(1) Documented evidence of the assessment of the needs of the patient, of an appropriate plan of care, and of the care and services furnished.

(2) Identification data and consent forms.

(3) Medical history.

(4) Report of physical examinations, if any.

(5) Observations and progress notes.

(6) Reports of treatments and clinical findings.

(7) Discharge summary including final diagnosis(es) and prognosis.

(c) Standard: Completion of records and centralization of reports. Current clinical records and those of discharged patients are completed promptly. All clinical information pertaining to a patient is centralized in the patient's clinical record. Each physician signs the entries that he or she makes in the clinical record.

(d) Standard: Retention and preservation. Clinical records are retained for at least:

(1) The period determined by the respective State statute, or the statute of limitations in the State; or

(2) In the absence of a State statute—

(i) Five years after the date of discharge; or

(ii) In the case of a minor, 3 years after the patient becomes of age under State law or 5 years after the date of discharge, whichever is longer.

(e) Standard: Indexes. Clinical records are indexed at least according to name of patient to facilitate acquisition of statistical medical information and retrieval of records for research or administrative action.

(f) Standard: Location and facilities. The organization maintains adequate facilities and equipment, conveniently located, to provide efficient processing of clinical records (reviewing, indexing, filing, and prompt retrieval).


§ 485.723 Condition of participation: Physical environment.

The building housing the organization is constructed, equipped, and maintained to protect the health and safety of patients, personnel, and the public and provides a functional, sanitary, and comfortable environment.

(a) Standard: Safety of patients. The organization satisfies the following requirements:

(1) It complies with all applicable State and local building, fire, and safety codes.

(2) Permanently attached automatic fire-extinguishing systems of adequate capacity are installed in all areas of the premises considered to have special fire hazards. Fire extinguishers are conveniently located on each floor of
§ 485.725 Condition of participation: Infection control.

The organization that provides outpatient physical therapy services establishes an infection-control committee of representative professional staff with responsibility for overall infection control. All necessary housekeeping and maintenance services are provided to maintain a sanitary and comfortable environment and to help prevent the development and transmission of infection.

(a) Standard: Infection-control committee. The infection-control committee establishes policies and procedures for investigating, controlling, and preventing infections in the organization and monitors staff performance to ensure that the policies and procedures are executed.

(b) All personnel follow written procedures for effective aseptic techniques. The procedures are reviewed annually and revised if necessary to improve them.

(c) Standard: Housekeeping. (1) The organization employs sufficient housekeeping personnel and provides all necessary equipment to maintain a safe, clean, and orderly interior. A full-time employee is designated as the one responsible for the housekeeping services and for supervision and training of housekeeping personnel.

(2) An organization that has a contract with an outside resource for housekeeping services may be found to be in compliance with this standard provided the organization or outside resource or both meet the requirements of the standard.

(d) Standard: Linen. The organization 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.

(e) Standard: Pest control. The organization's premises are maintained free...
§ 485.727 Condition of participation: Disaster preparedness.

The organization has a written plan, periodically rehearsed, 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 a disaster.

(a) Standard: Disaster plan. The organization has a written plan in operation, with procedures to be followed in the event of fire, explosion, or other disaster. The plan is developed and maintained with the assistance of qualified fire, safety, and other appropriate experts, and includes:

1. Transfer of casualties and records;
2. The location and use of alarm systems and signals;
3. Methods of containing fire;
4. Notification of appropriate persons; and
5. Evacuation routes and procedures.

(b) Standard: Staff training and drills. All employees are trained, as part of their employment orientation, in all aspects of preparedness for any disaster. The disaster program includes orientation and ongoing training and drills for all personnel in all procedures so that each employee promptly and correctly carries out his assigned role in case of a disaster.

§ 485.729 Condition of participation: Program evaluation.

The organization has procedures that provide for a systematic evaluation of its total program to ensure appropriate utilization of services and to determine whether the organization’s policies are followed in providing services to patients through employees or under arrangements with others.

(a) Standard: Clinical-record review. A sample of active and closed clinical records is reviewed quarterly by the appropriate health professionals to ensure that established policies are followed in providing services.

(b) Standard: Annual statistical evaluation. An evaluation is conducted annually of statistical data such as number of different patients treated, number of patient visits, condition on admission and discharge, number of new patients, number of patients by diagnosis(es), sources of referral, number and cost of units of service by treatment given, and total staff days or work hours by discipline.

PART 486—CONDITIONS FOR COVERAGE OF SPECIALIZED SERVICES FURNISHED BY SUPPLIERS

Subpart A—General Provisions

Sec. 486.1 Basis and scope.

Subpart B [Reserved]

Subpart C—Conditions for Coverage: Portable X-Ray Services

486.100 Condition for coverage: Compliance with Federal, State, and local laws and regulations.
486.102 Condition for coverage: Supervision by a qualified physician.
486.104 Condition for coverage: Qualifications, orientation, and health of technical personnel.
486.106 Condition for coverage: Referral for service and preservation of records.
486.108 Condition for coverage: Safety standards.
486.110 Condition for coverage: Inspection of equipment.

Subpart D—Conditions for Coverage: Outpatient Physical Therapy Services Furnished by Physical Therapists in Independent Practice

486.150 Condition for coverage: General requirements.
486.151 Condition for coverage: Supervision.
486.153 Condition for coverage: Compliance with Federal, State, and local laws.
486.155 Condition for coverage: Plan of care.
486.157 Condition for coverage: Physical therapy services.
486.159 Condition for coverage: Coordination of services with other organizations, agencies, or individuals.
§ 486.1 Basis and scope.

(a) Statutory basis. This part is based on the following sections of the Act:

1138(b)—for coverage of organ procurement services.
1861(p)—for coverage of outpatient physical therapy services furnished by physical therapists in independent practice.
1861(s) (3), (15), and (17)—for coverage of portable X-ray services.

(b) Scope. (1) This part sets forth the conditions for coverage of certain specialized services that are furnished by suppliers and that are not specified in other portions of this chapter.

(2) The conditions for coverage of other specialized services furnished by suppliers are set forth in the following regulations which, unless otherwise indicated, are part of this chapter:

(i) Ambulatory surgical center (ASC) services—Part 416.
(ii) Ambulance services—Part 410, subpart B.
(iii) ESRD services—Part 405, subpart U.
(iv) Laboratory services—Part 493.
(v) Mammography services—Part 410, subpart B (§410.34) and 21 CFR Part 900, subpart B, of the Food and Drug Administration regulations.
(vi) Rural health clinic and Federally qualified health center services—Part 491, subpart A.

[60 FR 50447, Sept. 29, 1995]

Subpart B [Reserved]

Subpart C—Conditions for Coverage: Portable X-Ray Services

AUTHORITY: Secs. 1102, 1861(s) (3), (11) and (12), 1864, and 1871 of the Social Security Act (42 U.S.C. 1302, 1395x(s) (3), (11), and (12), 1395aa and 1395hh).


§ 486.100 Condition for coverage: Compliance with Federal, State, and local laws and regulations.

The supplier of portable X-ray services is in conformity with all applicable Federal, State, and local laws and regulations.

(a) Standard—licensure or registration of supplier. In any State in which State or applicable local law provides for the licensure or registration of suppliers of X-ray services, the supplier is (1) licensed or registered pursuant to such law, or (2) approved by the agency of the State or locality responsible for licensure or registration as meeting the standards established for such licensure or registration.

(b) Standard—licensure or registration of personnel. All personnel engaged in operating portable X-ray equipment are currently licensed or registered in accordance with all applicable State and local laws.

(c) Standard—licensure or registration of equipment. All portable X-ray equipment used in providing portable X-ray
services is licensed or registered in accordance with all applicable State and local laws.

(d) Standard—conformity with other Federal, State, and local laws and regulations. The supplier of portable X-ray services agrees to render such services in conformity with Federal, State, and local laws relating to safety standards.

§ 486.102 Condition for coverage: Supervision by a qualified physician.

Portable X-ray services are provided under the supervision of a qualified physician.

(a) Standard—physician supervision. The performance of the roentgenologic procedures is subject to the supervision of a physician who meets the requirements of paragraph (b) of this section and one of the following requirements is met:

(1) The supervising physician owns the equipment and it is operated only by his employees, or

(2) The supervising physician certifies annually that he periodically checks the procedural manuals and observes the operators' performance, that he has verified that equipment and personnel meet applicable Federal, State, and local licensure and registration requirements and that safe operating procedures are used.

(b) Standard—qualifications of the physician supervisor. Portable X-ray services are provided under the supervision of a licensed doctor of medicine or licensed doctor of osteopathy who is qualified by advanced training and experience in the use of X-rays for diagnostic purposes, i.e., he (1) is certified in radiology by the American Board of Radiology or by the American Osteopathic Board of Radiology or possesses qualifications which are equivalent to those required for such certification, or (2) is certified or meets the requirements for certification in a medical specialty in which he has become qualified by experience and training in the use of X-rays for diagnostic purposes, or (3) specializes in radiology and is recognized by the medical community as a specialist in radiology.

§ 486.104 Condition for coverage: Qualifications, orientation and health of technical personnel.

Portable X-ray services are provided by qualified technologists.

(a) Standard—qualifications of technologists. All operators of the portable X-ray equipment meet the requirements of paragraph (a) (1), (2), or (3) of this section:

(1) Successful completion of a program of formal training in X-ray technology of not less than 24 months' duration in a school approved by the Council on Education of the American Medical Association or by the American Osteopathic Association, or have earned a bachelor's or associate degree in radiologic technology from an accredited college or university.

(2) For those whose training was completed prior to July 1, 1966, but on or after July 1, 1960: Successful completion of 24 full months of training and/or experience under the direct supervision of a physician who is certified in radiology by the American College of Radiology or who possesses qualifications which are equivalent to those required for such certification, and at least 12 full months of pertinent portable X-ray equipment operation experience in the 5 years prior to January 1, 1968.

(3) For those whose training was completed prior to July 1, 1960: Successful completion of 24 full months of training and/or experience of which at least 12 full months were under the direct supervision of a physician who is certified in radiology by the American College of Radiology or who possesses qualifications which are equivalent to those required for such certification, and at least 12 full months of pertinent portable X-ray equipment operation experience in the 5 years prior to January 1, 1968.

(b) Standard—personnel orientation. The supplier of portable X-ray services has an orientation program for personnel, based on a procedural manual.
which is: Available to all members of the staff, incorporates relevant portions of professionally recognized documents, and includes instruction in all of the following:

1. Precautions to be followed to protect the patient from unnecessary exposure to radiation;
2. Precautions to be followed to protect an individual supporting the patient during X-ray procedures from unnecessary exposure to radiation;
3. Precautions to be followed to protect other individuals in the surrounding environment from exposure to radiation;
4. Precautions to be followed to protect the operator of portable X-ray equipment from unnecessary exposure to radiation;
5. Considerations in determining the area which will receive the primary beam;
6. Determination of the time interval at which to check personnel radiation monitors;
7. Use of the personnel radiation monitor in providing an additional check on safety of equipment;
8. Proper use and maintenance of equipment;
9. Proper maintenance of records;
10. Technical problems which may arise and methods of solution;
11. Protection against electrical hazards;
12. Hazards of excessive exposure to radiation.

(a) Standard—referral by a physician. Portable X-ray examinations are performed only on the order of a doctor of medicine or doctor of osteopathy licensed to practice in the State. The supplier’s records show that:

1. The X-ray test was ordered by a licensed doctor of medicine or doctor of osteopathy, and
2. Such physician’s written, signed order specifies the reason an X-ray test is required, the area of the body to be exposed, the number of radiographs to be obtained, and the views needed; it also includes a statement concerning the condition of the patient which indicates why portable X-ray services are necessary.

(b) Standard—records of examinations performed. The supplier makes for each patient a record of the date of the X-ray examination, the name of the patient, a description of the procedures ordered and performed, the referring physician, the operator(s) of the portable X-ray equipment who performed the examination, the physician to whom the radiograph was sent, and the date it was sent.

(c) Standard—preservation of records. Such reports are maintained for a period of at least 2 years, or for the period of time required by State law for such records (as distinguished from requirements as to the radiograph itself), whichever is longer.


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§ 486.106 Condition for coverage: Referral for service and preservation of records.

All portable X-ray services performed for Medicare beneficiaries are ordered by a doctor of medicine or doctor of osteopathy and records are properly preserved.

(a) Standard—tubes housing and devices to restrict the useful beam. The tube housing is of diagnostic type. Diaphragms, cones, or adjustable collimators capable of restricting the useful beam to the area of clinical interest are used and provide the same degree of
protection as is required of the housing.

(b) Standard—total filtration. (1) The aluminum equivalent of the total filtration in the primary beam is not less than that shown in the following table except when contraindicated for a particular diagnostic procedure.

<table>
<thead>
<tr>
<th>Operating kVp</th>
<th>Total filtration (inherent plus added)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Below 50 kVp</td>
<td>0.5 millimeters aluminum.</td>
</tr>
<tr>
<td>50–70 kVp</td>
<td>1.5 millimeters aluminum.</td>
</tr>
<tr>
<td>Above 70 kVp</td>
<td>2.5 millimeters aluminum.</td>
</tr>
</tbody>
</table>

(2) If the filter in the machine is not accessible for examination or the total filtration is unknown, it can be assumed that the requirements are met if the half-value layer is not less than that shown in the following table:

<table>
<thead>
<tr>
<th>Operating kVp</th>
<th>Half-value layer</th>
</tr>
</thead>
<tbody>
<tr>
<td>50 kVp</td>
<td>0.6 millimeters aluminum.</td>
</tr>
<tr>
<td>70 kVp</td>
<td>1.6 millimeters aluminum.</td>
</tr>
<tr>
<td>90 kVp</td>
<td>2.6 millimeters aluminum.</td>
</tr>
<tr>
<td>100 kVp</td>
<td>2.8 millimeters aluminum.</td>
</tr>
<tr>
<td>110 kVp</td>
<td>3.0 millimeters aluminum.</td>
</tr>
<tr>
<td>120 kVp</td>
<td>3.3 millimeters aluminum.</td>
</tr>
</tbody>
</table>

(c) Standard—termination of exposure. A device is provided to terminate the exposure after a preset time or exposure.

(d) Standard—control panel. The control panel provides a device (usually a milliammeter or a means for an audible signal to give positive indication of the production of X-rays whenever the X-ray tube is energized. The control panel includes appropriate indicators (labelled control settings and/or meters) which show the physical factors (such as kVp, mA, exposure time or whether timing is automatic) used for the exposure.

(e) Standard—exposure control switch. The exposure control switch is of the dead-man type and is so arranged that the operator can stand at least 6 feet from the patient and well away from the useful beam.

(f) Standard—protection against electrical hazards. Only shockproof equipment is used. All electrical equipment is grounded.

(g) Standard—mechanical supporting or restraining devices. Mechanical supporting or restraining devices are provided so that such devices can be used when a patient must be held in position for radiography.

(h) Standard—protective gloves and aprons. Protective gloves and aprons are provided so that when the patient must be held by an individual, that individual is protected with these shielding devices.

(i) Standard—restriction of the useful beam. Diaphragms, cones, or adjustable collimators are used to restrict the useful beam to the area of clinical interest.

(j) Standard—personnel monitoring. A device which can be worn to monitor radiation exposure (e.g., a film badge) is provided to each individual who operates portable X-ray equipment. The device is evaluated for radiation exposure to the operator at least monthly and appropriate records are maintained by the supplier of portable X-ray services of radiation exposure measured by such a device for each individual.

(k) Standard—personnel and public protection. No individual occupationally exposed to radiation is permitted to hold patients during exposures except during emergencies, nor is any other individual regularly used for this service. Care is taken to assure that pregnant women do not assist in portable X-ray examinations.

§ 486.110 Condition for coverage: Inspection of equipment.

Inspections of all X-ray equipment and shielding are made by qualified individuals at intervals not greater than every 24 months.

(a) Standard—qualified inspectors. Inspections are made at least every 24 months by a radiation health specialist who is on the staff of or approved by an appropriate State or local government agency.

(b) Standard—records of inspection and scope of inspection. The supplier maintains records of current inspections which include the extent to which
§ 486.150 Condition for coverage: General requirements.

In order to be covered under Medicare as a supplier of outpatient physical therapy services, a physical therapist in independent practice must meet the following requirements:

(a) Be licensed in the State in which he or she practices.

(b) Meet one of the personnel qualifications specified in § 485.705(b).

(c) Furnish services under the circumstances described in § 410.60 of this chapter.

(d) Meet the requirements of this subpart.

[60 FR 2329, Jan. 9, 1995]

§ 486.151 Condition for coverage: Supervision.

The services are furnished by or under the direct supervision of a qualified physical therapist in independent practice.

[60 FR 2329, Jan. 9, 1995]

§ 486.153 Condition for coverage: Compliance with Federal, State, and local laws.

The physical therapist in independent practice and staff, if any, are in compliance with all applicable Federal, State, and local laws and regulations:

(a) Standard: Licensure of facility. In any State in which State or applicable local law provides for the licensing of the facility of a physical therapist, such facility is:

(1) Licensed pursuant to such law; or

(2) If not subject to licensure, is approved (by the agency of such State or locality responsible for licensing) as meeting the standards established for such licensing.

(b) Standard: Licensure or registration of personnel. The physical therapist in independent practice and staff, if any, are licensed or registered in accordance with applicable laws.

[41 FR 20065, May 21, 1976, unless otherwise noted. Redesignated at 42 FR 52826, Sept. 30, 1977. Redesignated and amended at 60 FR 2326, 2329, Jan. 9, 1995]

§ 486.155 Condition for coverage: Plan of care.

For each patient, a written plan of care is established and periodically reviewed by the individual who established it.

(a) Standard: Medical history and prior treatment. The physical therapist obtains the following information before or at the time of initiation of treatment:

(1) The patient's significant past history.

(2) Diagnosis(es), if established.

(3) Physician's orders, if any.

(4) Rehabilitation goals and potential for their achievement.

(5) Contraindications, if any.

(6) The extent to which the patient is aware of the diagnosis(es) and prognosis.

(7) If appropriate, the summary of treatment provided and results achieved during previous periods of physical therapy services or institutionalization.

(b) Standard: Plan of care. (1) For each patient there is a written plan of care that is established by the physician or by the physical therapist who furnishes the services.

(2) The plan indicates anticipated goals and specifies for physical therapy services the:

(i) Type;

(ii) Amount;

(iii) Frequency; and

(iv) Duration.

(3) The plan of care and results of treatment are reviewed by the physician or by the therapist at least as often as the patient's condition requires, and the indicated action is taken.

(4) Changes in the plan of care are noted in the clinical record. If the patient has an attending physician, the
therapist who furnishes the services promptly notifies him or her of any change in the patient's condition or in the plan of care. (For Medicare patients, the plan must be reviewed by a physician in accordance with §410.61(e).)

[54 FR 36879, Sept. 20, 1989. Redesignated and amended at 60 FR 2326, 2329, Jan. 9, 1995]

§ 486.157 Condition for coverage: Physical therapy services.

The physical therapist in independent practice provides an adequate program of physical therapy services and has the facilities and equipment necessary to carry out the services offered.

(a) Standard: Adequate program. The physical therapist will be considered to have an adequate physical therapy program when services can be provided, utilizing therapeutic exercise and the modalities of heat, cold, water, and electricity; patient evaluations are conducted; and tests and measurements of strength, balance, endurance, range of motion, and activities of daily living are administered.

(b) Standard: Supervision of physical therapy services. Physical therapy services are provided by, or under the supervision of, a qualified physical therapist.

[41 FR 20865, May 21, 1976, unless otherwise noted. Redesignated at 42 FR 52826, Sept. 30, 1977. Redesignated and amended at 60 FR 2326, 2329, Jan. 9, 1995]

§ 486.159 Condition for coverage: Coordination of services with other organizations, agencies, or individuals.

The physical therapist coordinates her physical therapy services with the health and medical services the patient receives from organizations or agencies or other individual practitioners through exchange of information that meets the following standard:

If a patient is receiving or has recently received, from other sources, services related to the physical therapy program, the physical therapist exchanges pertinent documented information with those other sources—

(a) On a regular basis;

(b) Subject to the requirements for protection of the confidentiality of

§ 486.161 Condition for coverage: Clinical records.

The physical therapist in independent practice maintains clinical records on all patients in accordance with accepted professional standards and practices. The clinical records are completely and accurately documented, readily accessible, and systematically organized to facilitate retrieving and compiling information.

(a) Standard: Protection of clinical record information. Clinical-record information is recognized as confidential and is safeguarded against loss, destruction, or unauthorized use. Written procedures govern use and removal of records and include conditions for release of information. A patient's written consent is required for release of information not authorized by law.

(b) Standard: Content. The clinical record contains sufficient information to identify the patient clearly, to justify the diagnosis(es) and treatment, and to document the results accurately. All clinical records contain the following general categories of data:

(1) Documented evidence of the assessment of the needs of the patient, of an appropriate plan of care, and of the care and services provided,

(2) Identification data and consent forms,

(3) Medical history,

(4) Report of physical examination(s), if any,

(5) Observations and progress notes,

(6) Reports of treatments and clinical findings, and

(7) Discharge summary including final diagnosis(es) and prognosis.

(c) Standard: Completion of records and centralization of reports. Current clinical records and those of discharged patients are completed promptly. All clinical information pertaining to a patient is centralized in the patient’s clinical record.

(d) Standard: Retention and preservation. Clinical records are retained for a period of time not less than:
§ 486.163 Condition for coverage—physical environment.

The physical environment of the office or facility of the physical therapist in independent practice affords a functional, sanitary, safe, and comfortable surrounding for patients, personnel, and the public.

(a) Standard: Building construction. The construction of the building housing the physical therapy office meets all applicable State and local building, fire, and safety codes.

(b) Standard: Maintenance of the physical therapy office and equipment. There is a written preventive-maintenance program to ensure that equipment is operative and that the physical therapy office is clean and orderly. All essential mechanical, electrical, and patient-care equipment is maintained in safe operating condition, and is properly calibrated.

(c) Standard: Other environmental considerations. The building housing the physical therapy office is accessible to, and functional for, patients, personnel, and the public. Written effective procedures in aseptic techniques are followed by all personnel and the procedures are reviewed annually, and when necessary, revised.

(d) The physical therapist is alert to the possibility of fire and other nonmedical emergencies and has written plans that include—

(1) The means for leaving the office and the building safely, demonstrated, for example, by fire exit signs; and

(2) Other provisions necessary to ensure the safety of patients.

[41 FR 20865, May 21, 1976, unless otherwise noted. Redesignated at 42 FR 52826, Sept. 30, 1977. Redesignated and amended at 60 FR 2326, 2329, Jan. 9, 1995]

§ 486.301 Basis and scope.

(a) Statutory Basis. (1) Section 1138(b) of the Act sets forth the requirements that an organ procurement organization must meet to have its organ procurement services to hospitals covered under Medicare and Medicaid. These include certification as a “qualified” organ procurement organization (OPO) and designation as the OPO for a particular service area.

(2) Section 371(b) of the PHS Act sets forth the requirements for certification and the functions that a qualified OPO is expected to perform.

(b) Scope. This subpart sets forth—

(1) The conditions and requirements that an OPO must meet;

(2) The procedures for certification and designation of OPOs; and

(3) The terms of the agreement with HCFA, and the basis for, and the effect of, termination of the agreement.

[61 FR 19743, May 2, 1996]

§ 486.302 Definitions.

As used in this subpart, the following definitions apply:

Certification or recertification means a HCFA determination that an entity meets the standards for a qualified OPO at §486.304 of this subpart and is eligible for designation if it meets the additional conditions for designation at §§486.306 and 486.308. No payment ensues from certification alone.

Designation or redesignation means HCFA approval of an OPO for Medicare and Medicaid payment purposes under section 1138(b)(1)(F) of the Act. The
Health Care Financing Administration, HHS

§ 486.304 General requirements.

(a) Designation—a condition for payment. Payment may be made under the Medicare and Medicaid programs for organ procurement costs attributable to payments made by an OPO only if the organization has been designated by the Secretary as an OPO, payment to which may be treated as organ procurement costs for reimbursement of hospitals under Medicare and Medicaid.

(b) Requirements for designated status. To be the designated OPO for a service area, an entity must do the following:

(1) Submit to HCFA a written application for designation, using the application form prescribed by HCFA.

(2) Be certified as a qualified OPO.

(3) Participate in the Organ Procurement and Transplantation Network as specified in § 486.308.

(4) Enter into an agreement with HCFA that meets the requirements set forth in paragraph (c) of this section.

(5) Upon its initial designation, meet the requirements at § 486.310(a)(3) or § 486.310(b)(4), as appropriate, concerning working relationships with hospitals or transplant centers. During the initial designation period, the OPO is not required to demonstrate compliance with § 486.310(a)(1) and (a)(2) or § 486.310(b)(1), which set forth performance standards for OPoS.

(6) To be redesignated after an initial designation period, comply with all the requirements of this subpart, including those at § 486.310, which set forth performance standards for OPoS.

(7) Obtain HCFA approval before entering into any change of ownership, merger, consolidation, or change in its service area (see § 486.318, which sets forth requirements concerning approval for changes in ownership and service area). Failure to do so could result in termination.

(8) Enter into a working relationship with any hospitals, including transplant centers, in the OPO's service area that request a working relationship.

(c) Agreement with HCFA. An OPO must enter into an agreement with HCFA. The agreement is effective upon submission by the OPO and acceptance by HCFA, but may be terminated by either party. If an OPO agreement is terminated, payment for organ procurement services attributable to that OPO will not be made for services furnished on or after the effective date of termination. In the agreement, the OPO must agree to do the following:

(1) Maintain compliance with the requirements of titles XVIII and XIX of the Act, section 1138 of the Act, and applicable regulations, including the conditions set forth in this subpart, and the regulations of the OPTN approved...
§ 486.306 Qualifications for designation as an OPO.

To be designated as the OPO for a service area, an organization must, at the time of application and throughout the period of its designation, meet the following requirements:

(a) Be a nonprofit entity that is exempt from Federal income taxation under section 501 of the Internal Revenue Code of 1986.

(b) Have accounting and other fiscal procedures necessary to assure the fiscal stability of the organization, including procedures to obtain payment for kidneys and non-renal organs provided to transplant centers.

(c) Have an agreement with the Secretary to be reimbursed under Medicare for the procurement of covered organs.

(d) When OPUs may apply for designation. Entities may apply for designation whenever a service area becomes an open area.

(e) Designation periods—(1) General. An OPO is normally designated for 2 years. A designation period may not exceed 2 years but may be shorter.

(2) Redesignation. Redesignation must occur at least every 2 years and be completed before the end of an existing designation period.

(3) Interim designation. HCFA may designate an organization for an interim designation period if the period is needed in order for HCFA to make a final designation determination.

(i) The interim designee may be either the OPO previously designated for the service area or another organization.

(ii) The interim designation period does not exceed 180 days after the normal designation period has expired.

(iii) The interim designee must meet all requirements of section 371(b) of the Public Health Service Act (42 U.S.C. 273(b)) regarding qualified OPUs and must not be out of compliance with the requirements of section 1138(b)(1) (B) through (E) of the Act regarding requirements for payment of organ procurement costs under title XVIII or title XIX of the Act.
(d) Document that it has a defined service area that meets the requirements of §486.307.

(e) Have a director and such other staff, including an organ donation coordinator and an organ procurement specialist, necessary to obtain organs effectively from donors in its service area.

(f) Have a board of directors or an advisory board that has the authority to recommend policies relating to the donation, procurement, and distribution of organs. While an OPO may have more than one board, the members specified in paragraphs (f)(1) through (f)(5) of this section must be members of a single board. The board of directors or advisory board must be composed of the following:

1. Members who represent hospital administrators, tissue banks, voluntary health associations in its service area and either intensive care or emergency room personnel.
2. Members who represent the public residing in that area.
3. A physician with knowledge, experience, or skill in the field of human histocompatibility, or an individual with a doctorate degree in a biological science and with knowledge, experience, or skills in the field of human histocompatibility.
4. A neurosurgeon or another physician with knowledge or skills in the field of neurology.
5. A transplant surgeon from each transplant center in its service area with which the OPO has arrangements to coordinate its activities.

(g) To identify potential organ donors, have documented evidence that—
1. It has a working relationship with at least 75 percent of the hospitals that participate in the Medicare and Medicaid programs in its service area and that have an operating room and the equipment and personnel for retrieving organs; and
2. It conducts systematic efforts intended to acquire all usable organs from potential donors.

(h) Arrange for the appropriate tissue typing of donated organs.

(i) Have a system to equitably allocate donated organs among transplant patients that is consistent with—

1. “Guidelines for Preventing Transmission of Human Immunodeficiency Virus Through Transplantation of Human Tissue and Organs” issued by the Centers for Disease Control and Prevention (CDC) that are appended to this subpart; and
2. Rules of the Organ Procurement and Transplantation Network (OPTN), see §486.308.

(j) Provide or arrange for the transportation of donated organs to transplant centers.

(k) Have arrangements to coordinate its activities with transplant centers in the area.

(l) Have arrangements to cooperate with tissue banks for the retrieval, processing, preservation, storage and distribution of tissues as may be appropriate to assure that all usable tissues are obtained from potential donors.

(m) Maintain and make available upon request of the Secretary, the Comptroller General, or their designee data that relate to the performance standards.

(n) Maintain data in a format that can be readily used by a successor OPO and agree to turn over to the Secretary copies of all records and data necessary to assure uninterrupted service by a successor OPO newly designated by HCFA.

(o) Have a procedure for ensuring the confidentiality of patient records. Information from or copies of records may be released only to authorized individuals and the OPO must ensure that unauthorized individuals cannot gain access to or alter patient records. Original medical records may be released by the OPO only in accordance with Federal or State laws, court orders, or subpoenas.

(p) Conduct and participate in professional education concerning organ procurement.

(q) Ensure that appropriate donor screening and infection tests, consistent with OPTN standards and the CDC guidelines that are appended to this subpart, are performed by a laboratory that is certified in the appropriate specialty or subspecialty of service in accordance with part 493 of this chapter, including tests to prevent the acquisition of organs that are infected. 
with the etiologic agent for acquired immune deficiency syndrome.

(r) Assist hospitals in establishing and implementing protocols for making routine inquiries about organ donations by potential donors.

(s) Ensure that donors are tested for human immunodeficiency viral markers consistent with OPTN rules and the CDC guidelines appended to this subpart for solid organ donation.

(t) Submit accurate data to HCFA within 15 days following the end of a calendar year (unless otherwise notified) giving information on the following:

1. Population of designated service area based on the most recent U.S. Bureau of the Census data.
2. Number of actual donors.
3. Number of kidneys procured.
4. Number of kidneys transplanted.
5. Number of extrarenal organs by type procured.
6. Number of extrarenal organs by type transplanted.

§ 486.307 OPO service area size designation and documentation requirements.

(a) General documentation requirement. An OPO must make available to HCFA documentation verifying that the OPO meets the requirements of paragraphs (b) through (d) of this section at the time of application and throughout the period of its designation.

(b) Boundary designation. The defined service area either includes an entire Metropolitan Statistical Area or a New England County Metropolitan Area as specified by the Director of the Office of Management and Budget or does not include any part of such an area.

(c) Service area location and characteristics. An OPO must precisely define and document a proposed service area’s location through the following information:

1. The names of counties (or parishes in Louisiana) served or, if the service area includes an entire State, the name of the State.

2. Geographic boundaries of the service area for which U.S. population statistics are available.

3. Total population in service area.

4. The number of and the names of acute care hospitals in the service area with an operating room and the equipment and personnel to retrieve organs.

5. Sufficient size requirements.

(i) Before January 1, 1996, an OPO must demonstrate that it can procure organs from at least 50 potential donors per calendar year or that its service area comprises an entire State.

(ii) Beginning January 1, 1996, an OPO must meet at least one of the following requirements:

(1) Its service area must include an entire State or official U.S. territory.

(2) It must either procure organs from an average of at least 24 donors per calendar year in the 2 years before the year of redesignation or request and be granted an exception to this requirement under paragraph (d)(3) or (d)(4) of this section.

(iii) In the case of an OPO operating exclusively in a noncontiguous U.S. State, a U.S. territory, or a U.S. commonwealth, such as Hawaii or Puerto Rico, it must procure organs at the rate of 50 percent of the national average of all OPOs for kidney procurement per million population and for kidney transplantation per million population.

(iv) If it is an entity that has not been previously designated as an OPO, it must demonstrate that it can procure organs from at least 50 potential donors per calendar year.

3. HCFA may grant an OPO an exception to paragraph (d)(2)(ii) of this section if the OPO can demonstrate that—

(i) It failed to meet the requirement because of unusual circumstances beyond its control;

(ii) It has historically maintained a service area of sufficient size to meet the criterion in paragraph (d)(2)(ii) of this section; and

(iii) It has a specific plan to meet the size criterion in paragraph (d)(2)(ii) of this section in the future.

4. During the 1996 redesignation process only, HCFA may grant an exception to paragraph (d)(2)(ii) of this section to an OPO that can demonstrate that—

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§ 486.310 Condition: Adherence to performance standards.

(a) Standards before January 1, 1996.
Before January 1, 1996, OPOs must meet the following performance standards:

(1) Each OPO must procure within its service area a minimum ratio of 23 cadaveric kidneys per million population of its service area for each 12-month period surveyed.

(2) Each OPO must provide a minimum ratio of cadaveric kidneys procured in its service area and transplanted (either locally or exported and transplanted) of 19 cadaveric kidneys per million population of its service area for each 12-month period surveyed.

(b) Standards beginning on January 1, 1996.
Except as specified in paragraph (c) of this section, each OPO must achieve at least 75 percent of the national mean for four of the following five performance categories, averaged over the 2 calendar years before the year of redesignation:

(1) Number of actual donors per million population.

(2) Number of kidneys recovered per million population.

(3) Number of extrarenal organs recovered per million population.

(4) Number of kidneys transplanted per million population.

(5) Number of extrarenal organs transplanted per million population.

(3) Exception for 1996 transition period.
During the 1996 designation period only, HCFA may continue to designate for a service area an OPO that does not meet the standards under paragraph (b) of this section if the OPO:

(i) Meets three of the criteria in paragraphs (b)(1) through (b)(5) of this section; and

(ii) Submits an acceptable corrective action plan in accordance with paragraph (d) of this section.

(d) Corrective action plans and corrected information—(1) Corrective action plans.
(i) If a designated OPO does not meet the standards of paragraph (a) of
§ 486.314 Effect of failure to meet requirements.

Failure to continue to meet any of the requirements in §§486.306 and 486.308 or to meet the performance standards in §486.310 may result in termination of the OPO’s agreement with HCFA.


§ 486.316 Designation of one OPO for each service area.

(a) HCFA designates only one OPO per service area. Applications for designation are accepted only during a period when the service area is an open area. A service area is open for competition once the existing designation period has expired, when the existing designated status of the OPO for that service area has been terminated, or when no OPO has been designated for the area. HCFA may also declare the service area open in the event an OPO ceases to operate or HCFA has reasonable ground for anticipating it will cease to operate. In cases of urgent need (such as evidence of medically or ethically unsound practices), HCFA may terminate its agreement with an OPO immediately. The service area remains open until an OPO is designated for it. If more than one organization applies and substantially meets the requirements of §486.306 in a given service area, HCFA considers other factors in reaching a decision concerning which organization to designate. These factors follow:

(1) Prior performance, including the previous year’s experience in terms of the number of organs retrieved and wasted and the average cost per organ;
(2) Actual number of donors compared to the number of potential donors;
(3) The nature of relationships and degree of involvement with hospitals in the organization’s service area;
(4) Bed capacity associated with the hospitals with which the organizations have a working relationship;
(5) Willingness and ability to place organs within the service area; and
(6) Proximity of the organization to the donor hospitals.

(b) An organization that applies to HCFA to be the designated OPO for its service area and that is not designated may appeal its nondesignation under part 498 of this chapter.

(c) After January 1, 1996, a hospital must enter into an agreement only with the OPO designated to serve the area in which the hospital is located unless HCFA has granted the hospital a waiver under paragraphs (d) through (g) of this section to be serviced by another OPO.

(d) If HCFA changes the OPO designated for an area, hospitals located in that area must enter into agreements with the newly designated OPO or submit a request for a waiver in accordance with paragraph (e) of this section within 30 days of notice of the change in designation.

§ 486.325 Terminations of agreement with HCFA.

(a) Types—(1) Voluntary termination. If an OPO wishes to terminate its agreement, it must send written notice of its intention with the proposed effective date to HCFA. HCFA may approve the proposed date, set a different date no later than 6 months after the proposed effective date, or set a date less than 6 months after the proposed date if it determines that it would not disrupt services to the service area or otherwise interfere with the effective and efficient administration of the Medicare and Medicaid programs. If HCFA determines that a designated OPO has ceased to furnish organ procurement services to its service area, the cessation of services is deemed to constitute a voluntary termination by the OPO, effective on a date determined by HCFA.

(2) Involuntary termination. HCFA may terminate an agreement if it finds that an OPO no longer meets the conditions for coverage in this subpart, or is

§ 486.318 Changes in ownership or service area.

(a) OPO requirements. (1) A designated OPO considering a change in ownership or in its service area must notify HCFA before putting it into effect. This notification is required to ensure that the entity, as changed, will continue to satisfy Medicare and Medicaid requirements. A change in ownership takes place if there is the merger of one entity into another or the consolidation of one entity with another.

(2) A designated OPO considering a change in its service area must obtain prior HCFA approval. In the case of a service area change that results from a change of ownership due to merger or consolidation, the entities must submit the information required in an application for designation, or other written documentation HCFA determines to be necessary for designation.

(b) HCFA requirements. (1) If HCFA finds that the entity has changed to such an extent that it no longer satisfies the prerequisites for OPO designation, HCFA may terminate the OPO's agreement and declare the OPO's service area to be an open area.

(2) If HCFA finds that the changed entity continues to satisfy the prerequisites for OPO designation, the period of designation of the changed entity is the remaining designation term of the OPO that was reorganized. If more than one designated OPO is involved in the reorganization, the remaining designation term is ordinarily the longest of the remaining periods. HCFA may determine, however, that a shorter period applies if it decides that a shorter period is in the best interest of the Medicare and Medicaid programs. The performance standards of §486.310 apply at the end of this remaining period.

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not in substantial compliance with any other applicable Federal regulations or provisions of titles XI, XVIII, or title XIX of the Act. HCFA may also terminate an agreement immediately in cases of urgent need, such as the discovery of unsound medical practices.

(b) Notice to OPO. HCFA gives notice of termination to an OPO at least 90 days before the effective date stated in the notice.

(c) Appeal right. The OPO may appeal the termination in accordance with the provisions set forth in part 498, which sets forth appeals procedures for determinations that affect participation in the Medicare and Medicaid programs.

(d) Effects of termination. When an OPO agreement is terminated—

(1) Medicare and Medicaid payments may not be made for organ procurement services the OPO furnishes on or after the effective date of termination; and

(2) HCFA will accept applications from any entity to be the designated OPO for that area.

(e) Public notice. In the case of voluntary termination, the OPO must give prompt public notice of the date of termination, and such information regarding the effect of that termination as HCFA may require, through publication in local newspapers in the service area. In the case of involuntary termination, HCFA gives notice of the date of termination.

(f) Reinstatement. HCFA may, at its discretion, designate an OPO whose agreement was previously terminated if HCFA finds that the cause for termination has been removed, is satisfied that it is not likely to recur, has not designated another OPO for the service area, and finds that the OPO meets all the necessary requirements for designation.

Guidelines for Preventing Transmission of Human Immunodeficiency Virus Through Transplantation of Human Tissue and Organs

Summary

Although previous recommendations for preventing transmission of human immunodeficiency virus (HIV) through transplantation of human tissue and organs have markedly reduced the risk for this type of transmission, a case of HIV transmission from a screened, antibody-negative donor to several recipients raised questions about the need for additional federal oversight of transplantation of organs and tissues. A working group formed by the Public Health Service (PHS) in 1991 to address these issues concluded that further recommendations should be made to reduce the already low risk of HIV transmission by transplantation of organs and tissues. In revising these recommendations, the PHS sought assistance from public and private health professionals and representatives of transplant, public health, and other organizations. The revised guidelines address issues such as donor screening, testing, and exclusionary criteria; quarantine of tissue from living donors; inactivation or elimination of infectious organisms in organs and tissues before transplantation; timely detection, reporting, and tracking of potentially infected tissues, organs, and recipients; and recall of stored tissues from donors found after donation to have been infected. Factors considered in the development of these guidelines include differences between the screening of living and cadaveric donors; time constraints due to organ/tissue viability that may preclude performing certain screening procedures; differences in the risk of HIV transmission from various organs and tissues; differences between systems for procuring and distributing organs and tissues; the effect of screening practices on the limited availability of organs and some tissues; and the benefit of the transplant to the recipient.

INTRODUCTION

Exclusion of prospective blood donors based on their acknowledged risk behaviors for human immunodeficiency virus (HIV) infection began in 1983 (1). In 1985, when tests for HIV antibody became available, screening prospective donors of blood, organs, and other tissues also began (2,3). Both measures have reduced markedly the transmission of HIV via these routes.

A 1991 investigation determined that several recipients had been infected with HIV by an organ/tissue donor who had tested negative for HIV antibody at the time of donation (4). This occurrence raised questions about the need for additional federal oversight of transplantation of organs and tissues. To address these questions, the Public Health Service (PHS) formed a working group comprising representatives from several federal agencies. The working group concluded that, although existing recommendations are largely sufficient, revisions should be made to reduce the already low risk of HIV transmission via transplantation of organs and tissues. Adequate federal
regulations, recommendations, and guidelines for blood and plasma are already established and are not addressed in this document.

Those developing guidelines for other organs and tissues should consider donor screening and testing; quarantine of tissue from living donors; inactivation or elimination of infectious organisms in organs and tissues before transplantation; timely detection, reporting, and tracking of potentially infected organs, tissues, and recipients; and recall of stored tissue from donors found after donation to have been infected.

These guidelines apply largely to donation and transplantation of organs and solid tissues. Although they also apply generally to donation of human milk and semen, some modifications may be needed because donors of human milk and semen are living and often donate repeatedly. Additionally, donor milk should be pasteurized (a heating procedure that inactivates HIV) before dispensing. This document can serve as a general guide to facilities that bank breast milk or semen and should be followed where feasible.

In revising these recommendations for transplantation of organs and tissues, PHS sought assistance from public and private health professionals and representatives of transplant, public health, and other organizations (see pages iii-v). These guidelines do not supersede existing state laws but are to be implemented in accordance with existing statutes.

BACKGROUND

Epidemiology of HIV Infection in Recipients of Organs and Tissues

Most transmission of HIV to organ/tissue recipients occurred before 1985, before the implementation of donor screening. In addition to HIV transmission through blood and blood products, reports of HIV infection following transplantation have implicated the kidney, liver, heart, pancreas, bone, and possibly skin as sources of infection (4). HIV has also been transmitted from infected semen during artificial insemination (5).

Several studies and case reports indicate that HIV can be transmitted through breast milk from HIV-infected women to their children (6,7); these investigations include several prospective studies indicating that breast-fed infants are at greater risk of acquiring HIV from their infected mothers than are bottle-fed infants (8,9).

Reports of transmission from screened, HIV-antibody-negative donors of organs or tissues have been rare. In one instance, hemodilution from multiple transfusions given to the organ/tissue donor before collection of the blood sample resulted in an HIV-antibody test result that was initially false negative (10). Serum samples taken on admission, before the transfusions, and 2 days after the transfusions later tested positive for antibody to HIV. In another instance, a kidney donor tested HIV-antibody negative 8 months before donation but seroconverted between the time of testing and donation (11). The donor was not retested at the time of donation. In a third instance, an organ from an HIV-infected donor was transplanted under emergency conditions before results of the HIV-antibody test were known (12).

A fourth case involved transmission from an organ/tissue donor whose HIV-antibody test was negative at the time of donation (4). Most likely, the donation occurred sometime between infection and antibody seroconversion, which, for most
infected persons, ranges from 4 weeks to 6 months (13). Six years after the donor's death and ensuing donation, HIV infection in the stored donor material was confirmed by virus culture and polymerase chain reaction (PCR) of stored donor lymphocytes (41). Among the 41 recipients identified and tested, those who received the solid organs and unprocessed, fresh-frozen bone acquired HIV infection from the allografts (one recipient of a heart, two recipients of kidneys, one recipient of a liver, and three recipients of fresh-frozen bone). The recipients of other processed bone and relatively avascular soft tissue (fascia lata, tendons, ligaments, dura mater, and corneas) did not become HIV infected (4).

Current Use of Organ and Tissue Transplants

The number of transplants has grown considerably over the last several years, a phenomenon attributable to many factors, including the availability of improved immunosuppressant drugs. Approximately 66 Organ Procurement Organizations (OPOs) and 260 organ transplant centers are members of the Organ Procurement and Transplantation Network (OPTN). In 1990 these centers recovered approximately 15,000 organs (e.g., kidney, liver, heart, lung, and pancreas) from 4,500 donors.

OPOs and tissue banks also recovered tissues (other than the organs listed above) from an estimated 7,500–10,000 donors in 1990. These tissues were used in approximately 250,000–300,000 (mostly bone) allografts.

In 1990, member banks of the Eye Bank Association of America (EBAA) retrieved ocular tissue from more than 40,000 donors. These tissues are used for corneal transplantation and are also processed into epikeratophakia lenticules (EBAA Statistical Report, 1990).

More than 400 establishments either bank or commercially process one or more human tissues. Approximately 100 eye banks and 125 bone banks operate in the United States (although the number of hospitals that store bone for future transplantation is difficult to estimate). Also, several hospitals may retrieve and store bone from living donors. Seven human milk banks operating in the United States process donor breast milk.

The American Fertility Society is aware of approximately 100 semen banks in the United States. Slightly fewer than half of artificial inseminations performed in the United States involve unrelated-donor semen used to inseminate approximately 75,000 women per year. In addition to these 100 semen banks, an undetermined number of smaller banks are hospital based or located in the offices of individual physicians.

The National Heart, Lung, and Blood Institute (NHLBI) within the National Institutes of Health (NIH) is aware of 99 bone marrow transplant centers, of which 41 participate in programs involving bone marrow transplants from unrelated donors. Many additional facilities are equipped to obtain marrow from donors. About 2,200 bone marrow transplants involving allogeneic marrow took place in the United States in 1991. Of those, approximately 435 were provided by donors who were not related to the recipients. Peripheral blood stem cells are being used for autologous transplantation and, in the future, may be useful for allogeneic use. Furthermore, cord blood stem cells are being used for both related- and unrelated-donor allogeneic transplantation.
Current Guidelines and Recommendations

Procedures for procurement and transplantation of organs and tissues are addressed by a) federal laws, regulations, and guidelines; b) state laws and regulations; and c) voluntary industry standards. Several federal agencies either directly or indirectly regulate procurement and transplantation of organs and tissues. These activities range from the publication of guidelines that address the transmission of communicable diseases through transplantation to regulatory requirements for registration and premarket product licensure or approval (blood and certain other tissue products).

The Health Resources and Services Administration (HRSA), through the United Network for Organ Sharing (UNOS), administers the contract for OPTN as required by Section 372 of the Public Health Service Act and as amended (42 USC 274). The contract covers specified solid organs (kidney, liver, heart, lung, and pancreas) but does not cover corneas, eyes, or other tissues. Technically, all UNOS policies are voluntary; however, HRSA is currently developing regulations dealing with OPTN membership and operation.

Under a separate contract with HRSA, UNOS maintains a Scientific Registry for Transplant Recipients that includes information on all solid-organ transplant recipients (since October 1, 1987) from the date of transplantation until failure of the graft or death of the patient. In addition, HRSA informally conveys recommendations to organizations involved in procurement and transplantation of organs. Through OPTN and the Scientific Registry for Transplant Recipients, HRSA has the capacity to link organ donors and their recipients.

FDA regulates a limited number of specific tissues as either "biological products" or "medical devices." Examples of tissues include blood, dura mater, corneal lenticules, umbilical veins, nonautologous cultured skin, and heart valves. In addition, FDA has recently published regulations regarding behavioral screening and infectious-disease testing (HIV-1, HIV-2, hepatitis B virus, and hepatitis C virus) for donors of human tissue for transplantation (14). FDA also regulates certain agents and devices for processing bone marrow, although bone marrow transplants from unrelated donors are under the auspices of NHLBI.

NHLBI manages the federal contract for the National Marrow Donor Program. Two bone marrow donor registries currently exist: one independent registry and one registry managed through the NHLBI contractor. Each registry group has voluntary guidelines/standards that resemble blood-banking standards. Although federal regulations have not yet been promulgated, the current practice of bone marrow acquisition and transplantation includes procedures to reduce the risk of HIV transmission. NHLBI is preparing regulations that will set forth criteria, standards, and procedures for entities involved in bone marrow collection, processing, and transplantation. These entities include the National Marrow Donor Registry, individual donor centers, donor registries, marrow-collection centers, and marrow-transplant centers. The regulations will include donor-selection criteria to prevent the transmission of infectious diseases, including HIV infection.

Donor Screening

PHS has made recommendations for preventing HIV transmission through organ/tissue transplantation and artificial insemination (1-3,15,16). These
recommendations include screening for behaviors that are associated with acquisition of HIV infection, a physical examination for signs and symptoms related to HIV infection, and laboratory screening for antibody to HIV.

PHS has made no specific recommendations for donation and banking of human milk, although HIV-infected women in the United States are advised to avoid breast feeding their infants because of the risk of HIV transmission through breast milk (17). The Human Milk Banking Association of North America has issued guidelines for the establishment and operation of human milk banks (18). These guidelines state that all human milk donors should be screened according to the American Association of Blood Banks' standards for screening blood donors. All milk accepted for donation should be pasteurized unless the recipient's condition requires fresh-frozen milk, in which case the milk bank director should consult with the medical director and advisory board to approve the dispensing of microbiologically screened, fresh-frozen milk from suitable donors.

Since March 1985, the FDA has licensed a number of screening and supplemental tests for detection and confirmation of HIV antibody. All these tests are intended for use on either fresh or freezer-stored samples of serum or plasma. The FDA has not required manufacturers to submit data showing that HIV-1 antigen and antibody-detection kits produce accurate results when applied to postmortem blood samples. Postmortem blood samples are often hemolyzed, which may affect the specificity of screening assays for HIV antibody (19,20).

The screening tests include enzyme immunoassays (EIA), several of which are also approved for testing blood spots dried onto a specific filter paper, which may provide a method for storing samples. Rapid screening assays for HIV antibody that use a latex-agglutination or EIA (microparticle-based) format have also been approved for screening serum, plasma, or whole blood. A licensed EIA for detecting antibodies to HIV-2 is also commercially available, as are "combination tests" that simultaneously detect antibodies to HIV-1 and HIV-2 (21). FDA has also licensed one manufacturer to make and distribute a test for detection of HIV-1 p24 antigen for patient diagnosis and prognosis of HIV infection but not for screening blood donors.

Western blot tests and an immunofluorescence assay for HIV-1 are approved for supplemental, more specific testing of serum, plasma, and whole-blood samples found reactive by HIV-1 antibody screening tests. No additional, more specific test is approved that confirms either antibodies to HIV-2 (21) or eluted, dried blood-spot results. The licensed p24-antigen test includes a neutralization procedure that is to be used for specific testing of samples with repeatedly reactive test results.

Federal regulations already require that all donations of blood, blood components, and plasma intended for further processing into injectable products ("source plasma") be screened with a licensed test that detects HIV antibody. Since June 1992, PHS has also required that all blood and plasma donations be screened for HIV-2 antibody.

PHS has not recommended the use of the licensed HIV-1 p24-antigen assay for screening donated blood or source plasma, nor has the kit been approved for use in donor screening. This position is based on findings from several studies indicating that a blood donor with a positive test for antigen and a negative test for antibody is rare (22,23). Such rarity is probably attributable to the effectiveness of the donor-qualification procedures, including donor education, voluntary exclusion, and
antibody testing that together operate to prevent donation by persons at increased risk for HIV infection.

Limited studies have been conducted to examine the use of the p24-antigen assay to screen organ/tissue donors (19,20,24). Among approximately 1,000 samples from HIV-1 antibody-negative donors, no donors had detectable HIV-1 p24 antigen.

**Recipient Screening**

Until recently, PHS had made no recommendations regarding routine testing of recipients of organs, tissues, semen, or donated human milk. However, in response to the July 18, 1991, report of the PHS Workgroup on Organ and Tissue Transplantation, HRSA asked UNOS to request that transplant centers implement an interim voluntary HIV-testing policy for organ recipients. HRSA has requested that recipients be tested for HIV-1 antibody immediately before transplantation and at 3, 6, and 12 months after transplantation. If HIV infection is diagnosed in an organ recipient, the results of the HIV test are reported by the transplant center to the Scientific Registry for Transplant Recipients and to the procuring OPO, in accordance with existing state laws. No comparable registry exists for recipients of tissues, semen, or donated human milk. However, the National Marrow Donor Program routinely tracks both donors and recipients of bone marrow for unrelated-donor transplants. This program reports no known seroconversions among either donors or recipients, although recipients are not routinely screened for HIV.

Routine testing of recipients after transplantation has several potential benefits. First, early identification of HIV infection in a recipient allows for early intervention before signs and symptoms develop. Both antiviral therapy to prevent progression to acquired immunodeficiency syndrome (AIDS) (25) and prophylactic therapy to prevent opportunistic infections (26,27) have been recommended for HIV-infected patients, based on CD4+ T-lymphocyte levels. Second, early identification of HIV infection in a transplant recipient allows for early intervention to prevent further transmission from the recipient to sex or needle-sharing partners and to future offspring (through vertical transmission from mother to infant). Third, early identification of HIV infection in a recipient potentially identifies an infectious donor. Should further investigation indicate that the donor is the source of the HIV infection in the recipient, other recipients of tissue from that same donor can be notified and stored tissue can be retrieved, preventing further transmission through transplantation.

Concern has been expressed that linking HIV infection in a transplant recipient to the transplantation may be difficult because many recipients may have also received blood or blood products or have other risk factors. However, identification of multiple HIV-infected recipients of tissue from the same donor strongly implicates the donor as the source of the HIV infection in the recipients. In addition, stored blood or lymphoid samples from the donor (when available) can be tested for the presence of virus to confirm the HIV-infection status of the donor (4).

Questions have been raised about whether transplant recipients who may be receiving immunosuppressive therapy to prevent rejection are capable of producing antibody against HIV if transmission occurs. Several reports now indicate that the HIV-antibody response is not delayed in transplant recipients receiving antirejection therapy, which primarily affects cellular immunity (4).
The additional costs of routine screening for HIV in recipients must be considered as well. The Institute of Medicine has estimated that laboratory costs are approximately $4 for a patient who tests negative and $35 for a patient who tests positive. (The latter cost includes the added expense of repeat EIAAs and Western blot or other supplemental tests.) These costs may be underestimates, however. The time required for pretest and posttest counseling was estimated to be approximately 0.5–1.0 hour for an HIV-seronegative patient and 1.5–2.0 hours for an HIV-seropositive patient (28).

**Inactivation of HIV in Tissues**

Thorough donor screening is considered the most effective method for preventing HIV transmission through transplantation; however, the use of chemical or physical inactivating or sterilizing agents to reduce further the already low risk of transmission has been considered. If such agents are to be useful, they must either inactivate or eliminate the virus while maintaining the functional integrity of the tissue or organ.

No mechanism for inactivating virus in whole organs currently exists. However, several agents have been suggested as possible disinfectants for tissues such as bone fragments (4). Pasteurization has been shown to inactivate HIV in human milk without substantially compromising nutritional and immunologic characteristics (29).

Although some physical and chemical agents have been shown to reduce the likelihood of isolating virus from treated solid tissues, conclusive evidence that those processes render solid tissue completely safe yet structurally intact is lacking. In the recent case of an HIV-infected donor who was antibody negative (4), tissues that had been processed in a variety of ways did not transmit HIV. These tissues included a) lyophilized fascia lata, tendons, or ligaments; b) dura mater that was lyophilized and irradiated with 3.0–3.4 Mrad of gamma radiation through a cobalt-60 source; c) bone fragments that were treated with ethanol and lyophilized; and d) one sample of fresh-frozen long bone with the marrow elements evacuated (4). However, because most of these tissues were relatively avascular, it is unclear whether the absence of HIV transmission was due to processing, avascularity, or both.

**General Considerations**

In developing guidelines for preventing HIV transmission from organ/tissue donors to recipients, several factors were considered: a) differences between the screening of living, brain-dead, and cadaveric donors; b) time constraints due to organ/tissue viability that may preclude performing certain screening procedures; c) differences in the risk for HIV transmission from various organs and tissues; d) differences between systems in place for procuring and distributing organs and tissues; e) the effect of screening practices on the limited availability of organs and some tissues; and f) the benefit of the transplant to the recipient (i.e., some transplants are lifesaving, whereas others are life enhancing).

Living donors can be interviewed about potential high-risk behavior, whereas deceased donors cannot. In the case of brain-dead or cadaveric donors, family members and others may be unable to provide an accurate risk history. Therefore, exclusion of potentially infected brain-dead or cadaveric donors relies even more heavily on laboratory screening and physical examinations than on interviews regarding high-risk behavior.
Screening procedures that require more than 24 hours to complete may not be feasible for brain-dead or cadaveric donors of organs and certain tissues. Most tissues must be recovered and most organs must be recovered and transplanted shortly after cessation of circulatory function of the donor. Whereas some tissues can be stored for months, others must be transplanted within a few days after procurement. These time constraints may limit the ability to interview certain family members or significant life partners who are not nearby and may preclude the use of certain laboratory screening tests that cannot be performed within these time constraints.

The precise risk of HIV transmission from various tissues is not known, yet some organs and tissues clearly present a higher risk for HIV transmission than others (4). For example, studies indicate that the risk for transmission from an organ of an HIV-infected donor is nearly 100%. Fresh-frozen, unprocessed bone also appears to carry a high risk for transmission, particularly if marrow elements and adherent tissue are not removed. Relatively avascular solid tissue, some of which is also processed by using techniques that might inactivate HIV, appears to carry a lower risk for HIV transmission.

As noted earlier in these guidelines, there is considerable variability in the role of federal agencies regarding transplantation of organs and tissues and the procurement and distribution systems. Oversight for, existence of, and compliance with recommendations also vary between these systems. When organs and tissues are procured from a single donor, tracking systems must involve multiple distribution systems that may be difficult to link.

Donor-screening practices must also consider the already inadequate supply of most organs and tissues needed for transplantation. However, even though attempts should be made to ensure the highest level of safety, donor-screening practices should not unnecessarily exclude acceptable potential donors.

Those involved in developing guidelines should consider that some transplants are lifesaving (e.g., a heart transplant), whereas others are life enhancing. Some physicians may be willing to offer the patient a transplant of a lifesaving organ from a donor whose HIV risk status is questionable but would not use life-enhancing tissue from such a donor.

RECOMMENDATIONS

Donor Screening

1. All prospective living donors or next of kin or significant life partners accompanying brain-dead or cadaveric donors should be informed of the general nature of the donor-evaluation process, including a review of medical and behavioral history, physical examination, and blood tests to exclude infectious agents that might be transmitted by organ or tissue transplant.

2. Prospective living donors or next of kin or significant life partners accompanying brain-dead or cadaveric donors should be informed about modes of transmission and risk factors for HIV infection, emphasizing that HIV can be transmitted via transplanted organs and tissues. They should be told that a negative test for HIV antibody does not guarantee that the donor is free of HIV infection because of the
rare situation of donation after infection but before seroconversion. Therefore, organs and tissue must not be transplanted from persons who may have engaged in activities that placed them at increased risk for HIV infection. This information should be presented in simple language to ensure that the donor, next of kin, or significant life partner understands what is considered high-risk behavior and the importance of excluding persons who have engaged in this behavior. Persons soliciting the donation should not place undue pressure to donate on potential living donors and those persons providing permission for potential brain-dead or cadaveric donors who might otherwise decline to donate or give permission because of high-risk behavior.

3. To ascertain risk factors, all prospective living donors should be interviewed in a confidential and sensitive manner by a health-care professional competent to elicit information about behaviors that place persons at risk for HIV infection. Interviewers should ask direct questions about high-risk behavior.

4. For potential pediatric donors for whom maternal transmission of HIV is a consideration, the mother and, if possible, the father should be interviewed about behaviors that may have placed them at risk for acquiring HIV infection that could have been transmitted to their child.

5. Except where retrieval occurs by legal authorization, the next of kin or significant life partner of brain-dead or cadaveric donors should be interviewed in a confidential and sensitive manner by a health-care professional regarding potential HIV risk factors in the donor. Other family members, friends, and sex partners may also need to be interviewed, if available. When consent for removal of organs/tissue is required, at least the person signing the consent form should be interviewed. Other possible sources of information about behavioral risk factors may include hospital, police, and coroner's records, if available. When an interview is not performed, as allowed by legal authorization, the transplant surgeon should be fully informed that the donation was accepted, even though a direct interview with the next of kin or significant life partner was not performed.

6. If available, the medical records, including autopsy reports of all donors, should be reviewed for signs and symptoms associated with HIV infection and for evidence of high-risk behavior (e.g., male-to-male sexual contact, acquisition of sexually transmitted diseases, exchange of sex for money or drugs, injecting-drug use, or birth to a mother either at risk for or infected with HIV).

7. All prospective donors of organs, solid tissue, and semen should undergo a physical examination as close as possible before donation, with special attention to physical signs of HIV disease and injecting-drug use. The extent of the physical examination should be determined by the responsible medical officials according to the context of organ/tissue donation. Human milk banks should obtain a release from the primary health-care provider certifying that the prospective donor is in good health and does not constitute a risk to potential recipients.

8. As with donors of blood and plasma, prospective living organ, tissue, semen, and milk donors found after careful screening to be acceptable donors should sign
a consent statement indicating that they have reviewed and understand the information provided regarding the spread of HIV and have agreed not to donate should they be at potential risk for spreading HIV. The statement should also indicate that prospective donors understand that they must be tested for HIV as part of the donor-screening process and will be notified of positive results as specified by any existing state statutes, regulations, or guidelines. For acceptable brain-dead or cadaveric donors, procurement personnel should document that a careful attempt has been made to eliminate persons at high risk through available information, including interview of family members or significant life partners, physical examination, review of medical records, autopsy findings, and any other records that might provide information about high-risk behavior or possible HIV infection. For either type of donor, the statement should be included as part of a general checklist or donor evaluation form covering all important aspects of the donor evaluation and should be included in the transplant records or record of the procuring agency. All records generated by the interview should be kept confidential.

Donor Testing

1. For all prospective donors, a blood sample obtained before any transfusions were administered (during the current hospital admission for inpatients) should be collected as close to the time of retrieval of tissue as possible. Bone marrow donors must provide blood samples far enough in advance of marrow harvest to permit the tests to be performed and results reported before the recipient's preparative regimen (marrow ablation) is begun. Samples should be tested for antibodies to both HIV-1 and HIV-2 by using FDA-licensed tests. Separate tests or a combination test for HIV-1 and HIV-2 may be used. All antibody-screening tests should be performed by EIA unless the condition of the recipient or donor dictates the use of a more rapid screening assay.

2. Transfusions and infusion of other fluids to the prospective donor might produce false-negative results because of hemodilution. Efforts should be made to perform HIV-antibody testing on the most recent pretransfusion/infusion specimen for which identity and quality can be ensured. Specimens should not be drawn immediately downstream from an intravenous site to prevent dilution with intravenous fluids.

Posttransfusion/infusion specimens may be considered for testing after efforts to obtain a pretransfusion/infusion sample have been exhausted and posttransfusion/infusion samples have been assessed for evidence of dilution. The suitability of posttransfusion/infusion samples must consider a) the volume of the material transfused as a percentage of the patient's total blood volume and b) the amount of time between the last transfusion/infusion and the collection of the sample to be tested. An exchange of one total blood volume will reduce the concentration of an intravascular substance such as IgG to 35% of initial levels if there is no replacement from the extravascular space. More than 50% of total body IgG is extravascular, and reequilibration to normal levels of IgG should be nearly complete within 24 hours of a total blood volume exchange of albumin (30).
3. The HIV p24-antigen assay may identify a few of the rare donors who are HIV-infected, yet antibody-negative; however, studies examining the utility of this assay for screening organ/tissue donors are limited and currently do not allow a definitive recommendation on the use of this test (19,24). The utility of other assays such as PCR, which are currently experimental, should be considered for evaluation as they become available for clinical use. Those institutions choosing to use the HIV-1 p24-antigen assay should be aware that in populations with low prevalence (e.g., organ/tissue donors), a large percentage of persons who test repeatedly reactive (without confirmation with the neutralization assay) will be false positive. Consideration should also be given to the potential problems with decreased specificity when the assay is used to test postmortem samples (19).

4. The testing algorithm for HIV-antibody assays should be performed as described in the package insert with an initial test and, if reactive, a retest on the same specimen. However, the time constraints of some situations may not accommodate the delay of repeat testing by EIA as described in the package insert. In such extreme cases of lifesaving organ transplantation, the sample should be set up in triplicate in the initial EIA. A repeatedly reactive result (positive screening test) is defined as reactivity above the test cutoff in two or more of the three assays. When testing by EIA is impractical, a more rapid licensed test should be performed in triplicate. Testing by the conventional algorithm should be performed as early as possible, even if it follows the procurement and/or transplant of the organs or tissues.

5. Results of HIV testing for organ/tissue donors should be handled confidentially, in accordance with general medical practices and applicable federal and state statutes, regulations, and guidelines.

6. Prospective living donors should be notified if they are found through the screening process to be HIV-infected. Because of the possibility of sexual or parenteral transmission, the spouse or known sex partners of brain-dead or cadaveric donors should be notified in accordance with state law. All notifications should be handled in a manner congruent with current recommendations regarding counseling, testing, and partner notification (31,32). Before the notification of these persons, transplant and procurement organizations should consult with their state health department concerning local notification policies.

Also before notification, the repeatedly reactive screening assay should be confirmed with more specific supplemental tests. An aliquot of the original sample should be analyzed by using the following, more specific tests. For repeatedly reactive HIV-1 antibody EIAs, an HIV-1 Western blot or immunofluorescence assay should be performed. For repeatedly reactive HIV-1 antigen assays (if performed), a neutralization procedure must be performed. For HIV-2, no licensed supplemental test is available; however, consideration may be given to the use of research assays such as Western blot, immunofluorescence, radioimmune precipitation, and synthetic peptide-based EIA. Arrangements for HIV-2 supplemental testing may need to be made with either the state or local health department. For repeatedly reactive combination HIV-1 and HIV-2 assays, the published testing algorithm should be followed (21). When the results of any supplemental tests are unclear, the use of research assays should be considered.
Notification of HIV-infected prospective living donors or spouses/known sex partners of cadaveric donors should be done in accordance with state law and in a confidential and sensitive manner by staff competent in counseling and discussing positive HIV results and their implications. If such staff are not available in the organ/tissue procurement organization, arrangements should be made with other organizations such as health departments or clinics to provide appropriate notification.

7. When it is possible to properly obtain and store samples, one or more of the following samples from the donor should be saved for at least 5 years after the expiration date of the tissue: dried blood spots, a frozen buffy coat, spleen cells, lymph node cells, bone marrow, and an aliquot of serum. These samples can be examined if subsequent information indicates that the donor may have donated during the period after infection but before antibody seroconversion.

8. Confirmed positive HIV test results in a prospective organ/tissue donor should be reported to state health agencies if required by state law or regulation.

Donor Exclusion Criteria

Regardless of their HIV antibody test results, persons who meet any of the criteria listed below should be excluded from donation of organs or tissues unless the risk to the recipient of not performing the transplant is deemed to be greater than the risk of HIV transmission and disease (e.g., emergent, life-threatening illness requiring transplantation when no other organs/tissues are available and no other lifesaving therapies exist). In such a case, informed consent regarding the possibility of HIV transmission should be obtained from the recipient.

Behavior/History Exclusionary Criteria

1. Men who have had sex with another man in the preceding 5 years.

2. Persons who report nonmedical intravenous, intramuscular, or subcutaneous injection of drugs in the preceding 5 years.

3. Persons with hemophilia or related clotting disorders who have received human-derived clotting factor concentrates.

4. Men and women who have engaged in sex in exchange for money or drugs in the preceding 5 years.

5. Persons who have had sex in the preceding 12 months with any person described in items 1–4 above or with a person known or suspected to have HIV infection.

6. Persons who have been exposed in the preceding 12 months to known or suspected HIV-infected blood through percutaneous inoculation or through contact with an open wound, nonintact skin, or mucous membrane.

7. Inmates of correctional systems. (This exclusion is to address issues such as difficulties with informed consent and increased prevalence of HIV in this population.)
Specific Exclusionary Criteria for Pediatric Donors

1. Children meeting any of the exclusionary criteria listed above for adults should not be accepted as donors.

2. Children born to mothers with HIV infection or mothers who meet the behavioral or laboratory exclusionary criteria for adult donors (regardless of their HIV status) should not be accepted as donors unless HIV infection can be definitively excluded in the child as follows:

   Children >18 months of age who are born to mothers with or at risk for HIV infection, who have not been breast fed within the last 12 months, and whose HIV antibody tests, physical examination, and review of medical records do not indicate evidence of HIV infection can be accepted as donors.

3. Children ≤18 months of age who are born to mothers with or at risk for HIV infection or who have been breast fed within the past 12 months should not be accepted as donors regardless of their HIV test results.

Laboratory and Other Medical Exclusionary Criteria

1. Persons who cannot be tested for HIV infection because of refusal, inadequate blood samples (e.g., hemodilution that could result in false-negative tests), or any other reasons.

2. Persons with a repeatedly reactive screening assay for HIV-1 or HIV-2 antibody regardless of the results of supplemental assays.

3. Persons whose history, physical examination, medical records, or autopsy reports reveal other evidence of HIV infection or high-risk behavior, such as a diagnosis of AIDS, unexplained weight loss, night sweats, blue or purple spots on the skin or mucous membranes typical of Kaposi’s sarcoma, unexplained lymphadenopathy lasting >1 month, unexplained temperature >100.5 F (38.6 C) for >10 days, unexplained persistent cough and shortness of breath, opportunistic infections, unexplained persistent diarrhea, male-to-male sexual contact, sexually transmitted diseases, or needle tracks or other signs of parenteral drug abuse.

Inactivation of HIV in Organs/Tissues

Definitive recommendations cannot yet be made regarding inactivation of HIV in organs and tissues because of lack of information about potentially effective inactivation measures. Research should continue in this area. Efforts to evaluate the effect of certain processing techniques on tissue sterility and quality should be expanded to include virologic studies for HIV. Thus, until more is known, it is prudent to process bone and bone fragments and carefully evacuate all marrow components from whole bone whenever feasible.

Quarantine

For semen donations and, when possible, for tissue donations from living donors, the collection should be placed in frozen quarantine and the donor retested for
antibodies to HIV-1 and HIV-2 after 6 months (75). The quarantined material should be released only if the follow-up test results have been obtained and are negative.

Record Keeping for Tracking of Recipients and Tissues
1. Each establishment involved in the acquisition, processing, distribution, or storage of organs or tissues should have a graft identification system that allows the tracking of organs and tissues from the donor source to the recipient institution and vice versa. Furthermore, each establishment involved in the acquisition of organs or tissues from a single donor should have mechanisms in place to facilitate the communication between establishments for the purposes of tracking organs and tissues to recipients who should be notified if HIV transmission from donor source material is confirmed. Procurement, processing, distribution, and storage centers should keep accurate records of the distribution of each organ/tissue according to the donor identification number, tissue type and identifying number, and identifying information for the receiving center, along with dates of procurement and distribution. Records should be kept a minimum of 10 years after expiration of tissue.

2. The transplantation center, hospital, physician, or dentist should keep accurate records of all organs/tissues received and the disposition of each. These records must be separate from patients’ medical records (e.g., in a log book) so that this information is easily obtainable should tracking be necessary. Recorded information should include the organ/tissue type; donor identification number; name of procurement or distribution center supplying the organ/tissue; recipient-identifying information; name of recipient’s physician or dentist; and dates of a) receipt by the center and b) either transplantation to the recipient or further distribution.

3. The donor identification number and organ or tissue type should be recorded in the recipient’s transplant/medical/dental record.

Testing and Reporting of Recipients
1. Health-care providers for transplant recipients and the recipients themselves should be aware of the small but potential risk of infections, including HIV, from transplanted organs and tissues. The recipient’s informed consent to the transplant should include acknowledgment of the risks, including transmission of HIV and other infections.

2. Until the risk for HIV transmission from screened donors has been clarified, recipients of solid organs should be routinely advised to be tested for HIV immediately before transplantation and at 3 months following the transplant. Testing of recipients should be done with consent of the recipient and should not be mandatory. Recipients of tissues other than solid organs do not require routine testing for HIV following receipt of the tissue from appropriately screened donors. Results of HIV testing of organ recipients should be collected and analyzed by the Scientific Registry for Transplant Recipients. (If data indicate no benefit from recipient testing, then this recommendation for recipient testing may be omitted in a revision of these guidelines.)
3. If a transplant recipient is found to be infected with HIV, the transplant center or health-care provider should, consistent with state law, immediately notify the state health department and the organization from which the tissue was obtained. HIV infection in a solid-organ recipient should also be reported to the Scientific Registry for Transplant Recipients.

Recall of Stored Tissue and Tracking of Recipients of Organs/Tissue from HIV-Infected Donors

1. Upon being notified that an organ/tissue recipient is infected with HIV, the organ/tissue collection center, in collaboration with the state or local health department and with assistance from CDC, is responsible for determining as soon as possible whether the donor was HIV-infected. This is done by determining the HIV-infection status of other recipients of organs/tissues (particularly those recipients of organs and fresh-frozen bone) and by laboratory testing of stored donor material. Experimental diagnostic laboratory assays such as PCR may be useful in these situations and should be used when they become available.

2. If evidence suggests HIV infection in the donor either from testing of stored donor specimens or by finding HIV infection in other recipients, all other recipients of that donor's tissue or organs should be notified through their transplanting physician and informed of the likelihood of HIV exposure and advised to undergo HIV testing.

3. HIV-infected recipients should be counseled about their need for medical evaluation and about prevention of HIV transmission to others. They should also be advised to inform their sex or needle-sharing partners of their potential risk and need for HIV counseling and testing. HIV-infected women should be informed of the risk of transmission of HIV to their children born after the transplant and be advised to have these children evaluated and to avoid breast-feeding. Pregnant women should receive pregnancy counseling about HIV.

4. All stored organs/tissues from a donor found to be HIV-infected should be retrieved and quarantined immediately and either used only for research purposes or destroyed, except when the transplantation of an indispensable organ/tissue is necessary to save the patient's life.

[61 FR 19745, May 2, 1996]
§ 488.1 Definitions.
488.28 Providers or suppliers, other than SNFs and NFs, with deficiencies.

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488.454 Duration of remedies.
488.456 Termination of provider agreement.

Authority: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395(hh)).
Source: 53 FR 22859, June 17, 1988, unless otherwise noted.

Subpart A—General Provisions

§ 488.1 Definitions.
As used in this part—
Accredited provider or supplier means a provider or supplier that has voluntarily applied for and has been accredited by a national accreditation program meeting the requirements of and approved by HCFA in accordance with §488.5 or §488.6.

Act means the Social Security Act.

AOA stands for the American Osteopathic Association.

Certification is a recommendation made by the State survey agency on the compliance of providers and suppliers with the conditions of participation, requirements (for SNFs and NFs), and conditions of coverage.

Conditions for coverage means the requirements suppliers must meet to participate in the Medicare program.

Conditions for participation means the requirements providers other than skilled nursing facilities must meet to participate in the Medicare program and includes conditions of certification for rural health clinics.

Full review means a survey of a hospital for compliance with all conditions of participation for hospitals.

JCAHO stands for the Joint Commission on Accreditation of Healthcare Organizations.

Medicare condition means any condition of participation or for coverage, including any long term care requirements.

Provider of services or provider means a hospital, critical access hospital, skilled nursing facility, home health agency, hospice, comprehensive outpatient rehabilitation facility, or provider of outpatient physical therapy or speech pathology services.

Rate of disparity means the percentage of all sample validation surveys for which a State survey agency finds noncompliance with one or more Medicare conditions and no comparable condition level deficiency was cited by the accreditation organization, where it is reasonable to conclude that the deficiencies were present at the time of the accreditation organization's most recent surveys of providers or suppliers of the same type.

Example: Assume that during a validation review period State survey agencies perform validation surveys at 200 facilities of the same type (for example, ambulatory surgical centers, home health agencies) accredited by the same accreditation organization. The State survey agencies find 60 of the facilities out of compliance with one or more Medicare conditions, and it is reasonable to conclude that these deficiencies were present at the time of the most recent survey by an accreditation organization. The accreditation organization, however, has found deficiencies comparable to the condition level deficiencies at only 22 of the 60 facilities. These validation results would yield ((60-22)/200) a rate of disparity of 19 percent.

Reasonable assurance means that an accreditation organization has demonstrated to HCFA's satisfaction that its requirements, taken as a whole, are at least as stringent as those established by HCFA, taken as a whole.

State includes the District of Columbia, the Commonwealth of Puerto Rico, the Virgin Islands, Guam, and American Samoa.

State survey agency means the State health agency or other appropriate State or local agency used by HCFA to perform survey and review functions for Medicare.

Substantial allegation of noncompliance means a complaint from any of a variety of sources (including complaints submitted in person, by telephone, through written correspondence, or in newspaper or magazine articles) that, if substantiated, would affect the health and safety of patients and raises doubts as to a provider's or supplier's noncompliance with any Medicare condition.

Supplier means any of the following: Independent laboratory; portable X-ray services; physical therapist in independent practice; ESRD facility; rural health clinic; Federally qualified health center; or chiropractor.

Validation review period means the one year period during which HCFA conducts a review of the validation surveys and evaluates the results of the most recent surveys performed by the accreditation organization.

§ 488.2 Statutory basis.

This part is based on the indicated provisions of the following sections of the Act:

1128—Exclusion of entities from participation in Medicare.
1128A—Civil money penalties.
1814—Conditions for, and limitations on, payment for Part A services.
1819—Requirements for SNFs.
1861(f)—Requirements for psychiatric hospitals.
1861(z)—Institutional planning standards that hospitals and SNFs must meet.
1861(ee)—Discharge planning guidelines for hospitals.
1861(ss)(2)—Accreditation of religious non-medical health care institutions.
1864—Use of State survey agencies.
1865—Effect of accreditation.
1880—Requirements for hospitals and SNFs of the Indian Health Service.
1883—Requirements for hospitals that provide SNF care.
1902—Requirements for participation in the Medicaid program.
1913—Medicaid requirements for hospitals that provide NF care.
1919—Medicaid requirements for NFs.

[60 FR 50443, Sept. 29, 1995, as amended at 64 FR 67052, Nov. 30, 1999]

§ 488.3 Conditions of participation; conditions for coverage; and long-term care requirements.

(a) Basic rules. In order to be approved for participation in or coverage under the Medicare program, a prospective provider or supplier must:

(1) Meet the applicable statutory definition in section 1139(b), 1819, 1832(a)(2)(F), 1861, 1881, or 1919 of the Act; and

(2) Be in compliance with the applicable conditions or long-term care requirements prescribed in subpart N, Q or U of part 405, part 416, subpart C of part 418, part 482, part 483, part 484, part 485, subpart A of part 491, or part 494 of this chapter.

(b) Special Conditions. (1) The Secretary, after consultation with the JCAHO or AOA, may issue conditions of participation for hospitals higher or more precise than those of either those accrediting bodies.

(2) The Secretary may, at a State’s request, approve health and safety requirements for providers and suppliers in that State, which are higher than those otherwise applied in the Medicare program.

(3) If a State or political subdivision imposes higher requirements on institutions as a condition for the purchase of health services under a State Medicaid Plan approved under Title XIX of the Act, (or if Guam, Puerto Rico, or the Virgin Islands does so under a State plan for Old Age Assistance under Title I of the Act, or for Aid to the Aged, Blind, and Disabled under the original Title XVI of the Act), the Secretary is required to impose similar requirements as a condition for payment under Medicare in that State or political subdivision.

[53 FR 22859, June 17, 1988, as amended at 58 FR 61838, Nov. 23, 1993]

§ 488.4 Application and reapplication procedures for accreditation organizations.

(a) A national accreditation organization applying for approval of deeming authority for Medicare requirements under § 488.5 or 488.6 of this subpart must furnish to HCFA the information and materials specified in paragraphs (a)(1) through (10) of this section.

A national accreditation organization reapplying for approval must furnish to HCFA whatever information and materials from paragraphs (a)(1) through (10) of this section that HCFA requests. The materials and information are—

(1) The types of providers and suppliers for which the organization is requesting approval;

(2) A detailed comparison of the organization’s accreditation requirements and standards with the applicable Medicare requirements (for example, a crosswalk);

(3) A detailed description of the organization’s survey process, including—

(i) Frequency of the surveys performed;

(ii) Copies of the organization’s survey forms, guidelines and instructions to surveyors;

(iii) Accreditation survey review process and the accreditation status decision-making process;

(iv) Procedures used to notify accredited facilities of deficiencies and the
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Procedures used to monitor the correction of deficiencies in accredited facilities; and

(v) Whether surveys are announced or unannounced;

(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 provider and supplier accredited;

(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) Policies and procedures with respect to an individual's participation in the survey or accreditation decision process of any facility with which the individual is professionally or financially affiliated;

(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) The organization's procedures for responding to and for the investigation of complaints against accredited facilities, including policies and procedures regarding coordination of these activities with appropriate licensing bodies and ombudsman programs;

(7) The organization's policies and procedures with respect to the withholding or removal of accreditation status for facilities that fail to meet the accreditation organization's standards or requirements, and other actions taken by the organization in response to noncompliance with its standards and requirements;

(8) A description of all types (for example, full, partial, type of facility, etc.) and categories (provisional, conditional, temporary, etc.) of accreditation offered by the organization, the duration of each type and category of accreditation and a statement specifying the types and categories of accreditation for which approval of deeming authority is sought;

(9) A list of all currently accredited facilities, the type and category of accreditation currently held by each facility, and the expiration date of each facility's current accreditation; and

(10) A list of all full and partial accreditation surveys scheduled to be performed by the organization.

(b) The accreditation organization must also submit the following supporting documentation—

(1) A written presentation that demonstrates the organization's ability to furnish HCFA with electronic data in ASCII comparable code;

(2) A resource analysis that demonstrates that the organization's staffing, funding and other resources are adequate to perform the required surveys and related activities; and

(3) A statement acknowledging that as a condition for approval of deeming authority, the organization will agree to—

(i) Notify HCFA in writing of any facility that has had its accreditation revoked, withdrawn, or revised, or that has had any other remedial or adverse action taken against it by the accreditation organization within 30 days of any such action taken;

(ii) Notify all accredited facilities within 10 days of HCFA's withdrawal of the organization's approval of deeming authority;

(iii) Notify HCFA in writing at least 30 days in advance of the effective date of any proposed changes in accreditation requirements;

(iv) Within 30 days of a change in HCFA requirements, submit to HCFA an acknowledgement of HCFA's notification of the change as well as a revised crosswalk reflecting the new requirements and inform HCFA about how the organization plans to alter its requirements to conform to HCFA's new requirements;

(v) Permit its surveyors to serve as witnesses if HCFA takes an adverse action based on accreditation findings;

(vi) [Reserved]

(vii) Notify HCFA in writing within ten days of a deficiency identified in any accreditation entity where the deficiency poses an immediate jeopardy to the entity's patients or residents or a hazard to the general public; and

(viii) Conform accreditation requirements to changes in Medicare requirements.
§ 488.5 Effect of JCAHO or AOA accreditation of hospitals.

(a) Deemed to meet. Institutions accredited as hospitals by the JCAHO or AOA are deemed to meet all of the Medicare conditions of participation for hospitals, except—

(1) The requirement for utilization review as specified in section 1861(e)(6) of the Act and in §482.30 of this chapter;

(2) The additional special staffing and medical records requirements that are considered necessary for the provision of active treatment in psychiatric hospitals (section 1861(f) of the Act) and implementing regulations; and

(3) Any requirements under section 1861(e) of the Act and implementing regulations that HCFA, after consulting with JCAHO or AOA, identifies as being higher or more precise than the requirements for accreditation (section 1865(a)(4) of the Act).

(b) Deemed status for providers and suppliers that participate in the Medicaid program. Eligibility for Medicaid participation can be established through Medicare deemed status for providers and suppliers that are not required under Medicaid regulations to comply with any requirements other than Medicare participation requirements for that provider or supplier type.

(c) Release and use of hospital accreditation surveys.

(1) A hospital deemed to meet program requirements must authorize its accreditation organization to release to HCFA and the State survey agency a copy of its most current accreditation survey together with any other information related to the survey that HCFA may require (including corrective action plans).

(2) HCFA may use a validation survey, an accreditation survey or other information related to the survey to determine that a hospital does not meet the Medicare conditions of participation.

(3) HCFA may disclose the survey and information related to the survey to the extent that the accreditation survey and related survey information
are related to an enforcement action taken by HCFA.

§ 488.6 Other national accreditation programs for hospitals and other providers and suppliers.

(a) In accordance with the requirements of this subpart, a national accreditation program for hospitals; psychiatric hospitals; SNFs; HHAs; ASCs; RHCs; CORFs; hospices; religious nonmedical health care institutions; screening mammography services; critical access hospitals; or clinic, rehabilitation agency, or public health agency providers of outpatient physical therapy, occupational therapy or speech pathology services may provide reasonable assurance to HCFA that it requires the providers or suppliers it accredits to meet requirements that are at least as stringent as the Medicare conditions when taken as a whole. In such a case, HCFA may deem the providers or suppliers the program accredits to be in compliance with the appropriate Medicare conditions. These providers and suppliers are subject to validation surveys under § 488.7 of this subpart. HCFA will publish notices in the FEDERAL REGISTER in accordance with § 488.8(b) identifying the programs and deeming authority of any national accreditation program and the providers or suppliers it accredits. The notice will describe how the accreditation organization's accreditation program provides reasonable assurance that entities accredited by the organization meet Medicare requirements. (See § 488.5 for requirements concerning hospitals accredited by JCAHO or AOA.)

(b) Eligibility for Medicaid participation can be established through Medicare deemed status for providers and suppliers that are not required under Medicaid regulations to comply with any requirements other than Medicare participation requirements for that provider or supplier type.

(c) (1) A provider or supplier deemed to meet program requirements under paragraph (a) of this section must authorize its accreditation organization to release to HCFA and the State survey agency a copy of its most current accreditation survey, together with any information related to the survey that HCFA may require (including corrective action plans).

(2) HCFA may determine that a provider or supplier does not meet the Medicare conditions on the basis of its own investigation of the accreditation survey or any other information related to the survey.

(3) Upon written request, HCFA may disclose the survey and information related to the survey—

(i) Of any HHA; or

(ii) Of any other provider or supplier specified at paragraph (a) of this section if the accreditation survey and related survey information relate to an enforcement action taken by HCFA.

§ 488.7 Validation survey.

(a) Basis for survey. HCFA may require a survey of an accredited provider or supplier to validate its organization's accreditation process. These surveys will be conducted on a representative sample basis, or in response to substantial allegations of noncompliance.

(1) When conducted on a representative sample basis, the survey is comprehensive and addresses all Medicare conditions or is focused on a specific condition or conditions.

(2) When conducted in response to a substantial allegation, the State survey agency surveys for any condition that HCFA determines is related to the allegations.

(3) If the State survey agency substantiates a deficiency and HCFA determines that the provider or supplier is out of compliance with any Medicare condition, the State survey agency conducts a full Medicare survey.

(b) Effect of selection for survey. A provider or supplier selected for a validation survey must—

(1) Authorize the validation survey to take place; and

(2) Authorize the State survey agency to monitor the correction of any deficiencies found through the validation survey.

(c) Refusal to cooperate with survey. If a provider or supplier selected for a validation survey fails to comply with
§ 488.8 Federal review of accreditation organizations.

(a) Review and approval of national accreditation organization. HCFA's review and evaluation of a national accreditation organization will be conducted in accordance with, but will not necessarily be limited to, the following general criteria—

1. The equivalency of an accreditation organization's accreditation requirements of an entity to the comparable HCFA requirements for the entity;
2. The organization's survey process to determine—
   i. The composition of the survey team, surveyor qualifications, and the ability of the organization to provide continuing surveyor training;
   ii. The comparability of survey procedures to those of State survey agencies, including survey frequency, and the ability to investigate and respond appropriately to complaints against accredited facilities;
   iii. The organization's procedures for monitoring providers or suppliers found by the organization to be out of compliance with program requirements. These monitoring procedures are to be used only when the organization identifies noncompliance. If noncompliance is identified through validation surveys, the State survey agency monitors corrections as specified at § 488.7(b)(3);
   iv. The ability of the organization to provide HCFA with electronic data in ASCII comparable code and reports necessary for effective validation and assessment of the organization survey process;
   v. The adequacy of staff and other resources;
   vi. The organization's ability to provide adequate funding for performing required surveys; and
   vii. The organization's policies with respect to whether surveys are announced or unannounced; and
3. The accreditation organization's agreement to provide HCFA with a copy of the most current accreditation survey together with any other information related to the survey as HCFA may require (including corrective action plans).

(b) Notice and comment. (1) HCFA will publish a proposed notice in the Federal Register whenever it contemplates approving an accreditation organization.

(2) The organization's survey process to determine—

(i) The composition of the survey team, surveyor qualifications, and the ability of the organization to provide continuing surveyor training;

(ii) The comparability of survey procedures to those of State survey agencies, including survey frequency, and the ability to investigate and respond appropriately to complaints against accredited facilities;

(iii) The organization's procedures for monitoring providers or suppliers found by the organization to be out of compliance with program requirements. These monitoring procedures are to be used only when the organization identifies noncompliance. If noncompliance is identified through validation surveys, the State survey agency monitors corrections as specified at § 488.7(b)(3);

(iv) The ability of the organization to provide HCFA with electronic data in ASCII comparable code and reports necessary for effective validation and assessment of the organization survey process;

(v) The adequacy of staff and other resources;

(vi) The organization's ability to provide adequate funding for performing required surveys; and

(vii) The organization's policies with respect to whether surveys are announced or unannounced; and

(3) The accreditation organization's agreement to provide HCFA with a copy of the most current accreditation survey together with any other information related to the survey as HCFA may require (including corrective action plans).
organization's application for deeming authority. The proposed notice will specify the basis for granting approval of deeming authority and the types of providers and suppliers accredited by the organization for which deeming authority would be approved. The proposed notice will also describe how the accreditation organization's accreditation program provides reasonable assurance that entities accredited by the organization meet Medicare requirements. The proposed notice will also provide opportunity for public comment.

(2) HCFA will publish a final notice in the Federal Register whenever it grants deeming authority to a national accreditation organization. Publication of the final notice will follow publication of the proposed notice by at least six months. The final notice will specify the effective date of the approval of deeming authority and the term of approval (which will not exceed six years).

(c) Effects of approval of an accreditation organization. HCFA will deem providers and suppliers accredited by an approved accreditation organization to meet the Medicare conditions for which the approval of deeming authority has specifically been granted. The deeming authority will take effect 90 days following the publication of the final notice.

(d) Continuing Federal oversight of equivalency of an accreditation organization and removal of deeming authority. This paragraph establishes specific criteria and procedures for continuing oversight and for removing the approval of deeming authority of a national accreditation organization.

(1) Comparability review. HCFA will compare the equivalency of an accreditation organization's accreditation requirements to the comparable HCFA requirements if—

(i) HCFA imposes new requirements or changes its survey process; and

(ii) An accreditation organization proposes to adopt new requirements or change its survey process. An accreditation organization must provide written notification to HCFA at least 30 days in advance of the effective date of any proposed changes in its accreditation requirements or survey process; and

(iii) An accreditation organization's approval has been in effect for the maximum term specified by HCFA in the final notice.

(2) Validation review. Following the end of a validation review period, HCFA will identify any accreditation programs for which—

(i) Validation survey results indicate a rate of disparity between certifications of the accreditation organization and certification of the State agency of 20 percent or more; or

(ii) Validation survey results, irrespective of the rate of disparity, indicate widespread or systematic problems in an organization's accreditation process that provide evidence that there is no longer reasonable assurance that accredited entities meet Medicare requirements.

(3) Reapplication procedures. (i) Every six years, or sooner as determined by HCFA, an approved accreditation organization must reapply for continued approval of deeming authority. HCFA will notify the organization of the materials the organization must submit as part of the reapplication procedure.

(ii) An accreditation organization that is not meeting the requirements of this subpart, as determined through a comparability review, must furnish HCFA, upon request and at any time, with the reapplication materials HCFA requests. HCFA will establish a deadline by which the materials are to be submitted.

(e) Notice. If a comparability or validation review reveals documentation that an accreditation organization is not meeting the requirements of this subpart, as determined through a comparability review, HCFA will provide written notice to the organization indicating that its deeming authority approval may be in jeopardy and that a deeming authority review is being initiated. The notice provides the following information—

(1) A statement of the requirements, instances, rates or patterns of discrepancies that were found as well as other related documentation;

(2) An explanation of HCFA's deeming authority review on which the final determination is based;
§ 488.8

(3) A description of the process available if the accreditation organization wishes an opportunity to explain or justify the findings made during the comparability or validation review;

(4) A description of the possible actions that may be imposed by HCFA based on the findings from the validation review; and

(5) The reapplication materials the organization must submit and the deadline for their submission.

(f) Deeming authority review. (1) HCFA will conduct a review of an accreditation organization's accreditation program if the comparability or validation review produces findings as described at paragraph (d)(1) or (2), respectively, of this section. HCFA will review as appropriate either or both—

(i) The requirements of the accreditation organization; or

(ii) The criteria described in paragraph (a)(1) of this section to reevaluate whether the accreditation organization continues to meet all these criteria.

(2) If HCFA determines, following the deeming authority review, that the accreditation organization has failed to adopt requirements comparable to HCFA's or submit new requirements timely, the accreditation organization may be given a conditional approval of its deeming authority for a probationary period of up to 180 days to adopt comparable requirements.

(3) If HCFA determines, following the deeming authority review, that the rate of disparity identified during the validation review meets either of the criteria set forth in paragraph (d)(2) of this section HCFA—

(i) May give the accreditation organization conditional approval of its deeming authority during a probationary period of up to one year (whether or not there are also noncomparable requirements) that will be effective 30 days following the date of that determination;

(ii) Will require the accreditation organization to release to HCFA upon its request any facility-specific data that is required by HCFA for continued monitoring;

(iii) Will require the accreditation organization to provide HCFA with a survey schedule for the purpose of intermittent onsite monitoring by HCFA staff, State surveyors, or both; and

(iv) Will publish in the Medicare Annual Report to Congress the name of any accreditation organization given a probationary period by HCFA.

(4) Within 60 days after the end of any probationary period, HCFA will make a final determination as to whether or not an accreditation program continues to meet the criteria described at paragraph (a)(1) of this section and will issue an appropriate notice (including reasons for the determination) to the accreditation organization and affected providers or suppliers. This determination will be based on any of the following—

(i) The evaluation of the most current validation survey and review findings. The evaluation must indicate an acceptable rate of disparity of less than 20 percent between the certifications of the accreditation organization and the certifications of the State agency as described at paragraph (d)(2)(i) of this section in order for the accreditation organization to retain its approval;

(ii) The evaluation of facility-specific data, as necessary, as well as other related information;

(iii) The evaluation of an accreditation organization's surveyors in terms of qualifications, ongoing training, composition of survey team, etc.;

(iv) The evaluation of survey procedures; or

(v) The accreditation requirements.

(5) If the accreditation program has not made improvements acceptable to HCFA during the probationary period, HCFA may remove recognition of deemed authority effective 30 days from the date that it provides written notice to the organization that its deeming authority will be removed.

(6) The existence of any validation review, deeming authority review, probationary period, or any other action by HCFA, does not affect or limit the conducting of any validation survey.

(7) HCFA will publish a notice in the Federal Register containing a justification of the basis for removing the deeming authority from an accreditation organization. The notice will provide the reasons the accreditation organization's accreditation program no longer meets Medicare requirements.
§ 488.10 State survey agency review: Statutory provisions.

(a) Section 1864(a) of the Act requires the Secretary to enter into an agreement with any State that is able and willing to do so, under which appropriate State or local survey agencies will determine whether:

(1) Providers or prospective providers meet the Medicare conditions of participation or requirements (for SNFs and NFs);

(2) Suppliers meet the conditions for coverage; and

(3) Rural health clinics meet the conditions of certification.

(b) Section 1865(a) of the Act provides that if an institution is accredited as a hospital by the JCAHO, it will be deemed to meet the conditions of participation:

(1) Except those specified in §488.5;

(2) Provided that such hospital, if it is included within a validation survey, authorizes the JCAHO to release to HCFA (on a confidential basis) upon request a copy of the most current JCAHO accreditation survey.

(c) Section 1865(b) of the Act provides that an accredited hospital which is found after a validation survey to have significant deficiencies related to the health and safety of patients will no longer be deemed to meet the conditions of participation.

(d) Section 1865(a) of the Act also provides that if HCFA finds that accreditation of a hospital; psychiatric hospital; SNF; HHA; hospice; ASC; RHC; CORF; laboratory; screening mammography service; critical access hospital; or clinic, rehabilitation agency, or public health agency provider of outpatient physical therapy, occupational therapy, or speech pathology services by any national accreditation organization provides reasonable assurance that any or all Medicare conditions are met, HCFA may treat the
§ 488.11 State survey agency functions.

State and local agencies that have agreements under section 1864(a) of the Act perform the following functions:

(a) Survey and make recommendations regarding the issues listed in § 488.10.

(b) Conduct validation surveys of accredited facilities as provided in § 488.7.

(c) Perform other surveys and carry out other appropriate activities and certify their findings to HCFA.

(d) Make recommendations regarding the effective dates of provider agreements and supplier approvals in accordance with § 489.13 of this chapter.

§ 488.12 Effect of survey agency certification.

Certifications by the State survey agency represent recommendations to HCFA.

(a) On the basis of these recommendations, HCFA will determine whether:

(1) A provider or supplier is eligible to participate in or be covered under the Medicare program; or

(2) An accredited hospital is deemed to meet the Medicare conditions of participation or is subject to full review by the State survey agency.

(b) Notice of HCFA’s determination will be sent to the provider or supplier.

§ 488.14 Effect of PRO review.

When a PRO is conducting review activities under section 1154 of the Act and part 466 of this chapter, its activities are in lieu of the utilization review and evaluation activities required of health care institutions under sections 1861(e)(6), and 1861(k) of the Act.

§ 488.18 Documentation of findings.

(a) The findings of the State agency with respect to each of the conditions of participation, requirements (for SNFs and NFs), or conditions for coverage must be adequately documented.

When the State agency certifies to the Secretary that a provider or supplier is not in compliance with the conditions or requirements (for SNFs and NFs), and therefore not eligible to participate in the program, such documentation includes, in addition to the description of the specific deficiencies which resulted in the agency’s recommendation, any provider or supplier response.

(b) If a provider or supplier is certified by the State agency as in compliance with the conditions or participation requirements (for SNFs and NFs) or as meeting the requirements for special certification (see § 488.54), with deficiencies not adversely affecting the health and safety of patients, the following information will be incorporated into the finding:

(1) A statement of the deficiencies that were found.

(2) A description of further action that is required to remove the deficiencies.

(3) A time-phased plan of correction developed by the provider and supplier and concurred with by the State agency.

(4) A scheduled time for a resurvey of the institution or agency to be conducted by the State agency within 90 days following the completion of the survey.

(c) If, on the basis of the State certification, the Secretary determines that the provider or supplier is eligible to participate, the information described in paragraph (b) of this section will be incorporated into a notice of eligibility to the provider or supplier.

(d) If the State agency receives information to the effect that a hospital or a critical access hospital (as defined in section 1861(mm)(1) of the Act) has violated § 489.24 of this chapter, the State agency is to report the information to HCFA promptly.


Effective Date Note: At 59 FR 32120, June 22, 1994, in § 488.18, paragraph (d) was added, and will not become effective until the information collection requirements are
§ 488.20 Periodic review of compliance and approval.

(a) Determinations by HCFA to the effect that a provider or supplier is in compliance with the conditions of participation, or requirements (for SNFs and NFs), or the conditions for coverage are made as often as HCFA deems necessary and may be more or less than a 12-month period, except for SNFs, NFs and HHAs. (See § 488.308 for special rules for SNFs and NFs.)

(b) The responsibilities of State survey agencies in the review and certification of compliance are as follows:

(1) Resurvey providers or suppliers as frequently as necessary to ascertain compliance and confirm the correction of deficiencies;

(2) Review reports prepared by a Professional Standards Review Organization (authorized under Part B Title XI of the Act) or a State inspection of care team (authorized under Title XIX of the Act) regarding the quality of a facility’s care;

(3) Evaluate reports that may pertain to the health and safety of patients; and

(4) Take appropriate actions that may be necessary to achieve compliance or certify noncompliance to HCFA.

(c) A State survey agency certification to HCFA that a provider or supplier is no longer in compliance with the conditions of participation or coverage will supersede the State survey agency’s previous certification.

§ 488.24 Certification of noncompliance.

(a) Special rules for certification of noncompliance for SNFs and NFs are set forth in § 488.330.

(b) The State agency will certify that a provider or supplier is not or is no longer in compliance with the conditions of participation or conditions for coverage where the deficiencies are of such character as to substantially limit the provider’s or supplier’s capacity to furnish adequate care or which adversely affect the health and safety of patients; or

(c) If HCFA determines that an institution or agency does not qualify for participation or coverage because it is not in compliance with the conditions of participation or conditions for coverage, or if a provider’s agreement is terminated for that reason, the institution or agency has the right to request that the determination be reviewed. (Appeals procedures are set forth in Part 498 of this chapter.)

§ 488.26 Determining compliance.

(a) Additional rules for certification of compliance for SNFs and NFs are set forth in § 488.330.

(b) The decision as to whether there is compliance with a particular requirement, condition of participation, or condition for coverage depends upon the manner and degree to which the provider or supplier satisfies the various standards within each condition. Evaluation of a provider’s or supplier’s performance against these standards enables the State survey agency to document the nature and extent of deficiencies, if any, with respect to a particular function, and to assess the need for improvement in relation to the prescribed conditions.

(c) The State survey agency must adhere to the following principles in determining compliance with participation requirements:

(1) The survey process is the means to assess compliance with Federal health, safety and quality standards;
(2) The survey process uses resident outcomes as the primary means to establish the compliance status of facilities. Specifically, surveyors will directly observe the actual provision of care and services to residents, and the effects of that care, to assess whether the care provided meets the needs of individual residents;

(3) Surveyors are professionals who use their judgment, in concert with Federal forms and procedures, to determine compliance;

(4) Federal procedures are used by all surveyors to ensure uniform and consistent application and interpretation of Federal requirements;

(5) Federal forms are used by all surveyors to ensure proper recording of findings and to document the basis for the findings.

(d) The State survey agency must use the survey methods, procedures, and forms that are prescribed by HCFA.

(e) The State survey agency must ensure that a facility’s actual provision of care and services to residents and the effects of that care on residents are assessed in a systematic manner.

[59 FR 56237, Nov. 10, 1994]

§ 488.28 Providers or suppliers, other than SNFs and NFs, with deficiencies.

(a) If a provider or supplier is found to be deficient with respect to one or more of the standards in the conditions of participation or conditions for coverage, it may participate in or be covered under the Health Insurance for the Aged and Disabled Program only if the facility has submitted an acceptable plan of correction for achieving compliance within a reasonable period of time acceptable to the Secretary.

(b) The existing deficiencies noted either individually or in combination neither jeopardize the health and safety of patients nor are of such character as to seriously limit the provider’s capacity to render adequate care.

(c)(1) If it is determined during a survey that a provider or supplier is not in compliance with one or more of the standards, it is granted a reasonable time to achieve compliance.

(2) The amount of time depends upon the—

(i) Nature of the deficiency; and

(ii) State survey agency’s judgment as to the capabilities of the facility to provide adequate and safe care.

(d) Ordinarily a provider or supplier is expected to take the steps needed to achieve compliance within 60 days of being notified of the deficiencies but the State survey agency may recommend that additional time be granted by the Secretary in individual situations, if in its judgment, it is not reasonable to expect compliance within 60 days, for example, a facility must obtain the approval of its governing body, or engage in competitive bidding.

[59 FR 56237, Nov. 10, 1994]

Subpart B—Special Requirements

§ 488.52 [Reserved]

§ 488.54 Temporary waivers applicable to hospitals.

(a) General provisions. If a hospital is found to be out of compliance with one or more conditions of participation for hospitals, as specified in part 482 of this chapter, a temporary waiver may be granted by HCFA. HCFA may extend a temporary waiver only if such a waiver would not jeopardize or adversely affect the health and safety of patients. The waiver may be issued for any one year period or less under certain circumstances. The waiver may be withdrawn earlier if HCFA determines this action is necessary to protect the health and safety of patients. A waiver may be granted only if:

(1) The hospital is located in a rural area. This includes all areas not delineated as “urban” by the Bureau of the Census, based on the most recent census;

(2) The hospital has 50 or fewer inpatient hospital beds;

(3) The character and seriousness of the deficiencies do not adversely affect the health and safety of patients; and

(4) The hospital has made and continues to make a good faith effort to comply with personnel requirements consistent with any waiver.

(b) Minimum compliance requirements. Each case will have to be decided on its individual merits, and while the degree and extent of compliance will vary, the institution must, as a minimum, meet
all of the statutory conditions in section 1861(e)(1)–(8), in addition to meeting such other requirements as the Secretary finds necessary under section 1861(e)(9). (For further information relating to the exception in section 1861(e)(5) of the Act, see paragraph (c) of this section.)

(c) Temporary waiver of 24-hour nursing requirement of 24-hour registered nurse requirement. HCFA may waive the requirement contained in section 1861(e)(5) that a hospital must provide 24-hour nursing service furnished or supervised by a registered nurse. Such a waiver may be granted when the following criteria are met:

(1) The hospital’s failure to comply fully with the 24-hour nursing requirement is attributable to a temporary shortage of qualified nursing personnel in the area in which the hospital is located.

(2) A registered nurse is present on the premises to furnish or supervise the nursing services during at least the daytime shift, 7 days a week.

(3) The hospital has in charge, on all tours of duty not covered by a registered nurse, a licensed practical (vocational) nurse.

(4) The hospital complies with all requirements specified in paragraph (a) of this section.

(d) Temporary waiver for technical personnel. HCFA may waive technical personnel requirements, issued under section 1861(e)(9) of the Act, contained in the Conditions of Participation; Hospitals (part 482 of this chapter). Such a waiver must take into account the availability of technical personnel and the educational opportunities for technical personnel in the area in which the hospital is located.

(1) Such facility is located in a rural area and the supply of skilled nursing facility services in such area is not sufficient to meet the needs of individual patients therein,

(2) Such facility has at least one fulltime registered nurse who is regularly on duty at such facility 40 hours a week, and

(3) Such facility (i) has only patients whose attending physicians have indicated (through physicians’ orders or admission notes) that each such patient does not require the services of a registered nurse for a 48-hour period, or (ii) has made arrangements for a registered nurse or a physician to spend such time at the facility as is determined necessary by the patient’s attending physician to provide necessary services on days when the regular fulltime registered nurse is not on duty.

(4) Such facility has made and continues to make a good faith effort to comply with the more than 40-hour registered nurse requirement, but such compliance is impeded by the unavailability of registered nurses in the area.

(b) Waiver of medical director requirement. To the extent that §§488.75(i) of this chapter requires any skilled nursing facility to engage the services of a medical director either part-time or full-time, the Secretary may waive such requirement for such periods as he deems appropriate if, based upon documented findings of the State agency, he determines that:

(1) Such facility is located in an area where the supply of physicians is not sufficient to permit compliance with this requirement without seriously reducing the availability of physician services within the area, and

(2) Such facility has made and continues to make a good faith effort to
§ 488.60 Special procedures for approving end stage renal disease facilities.

(a) Considerations for approval. An ESRD facility which wishes to be approved for coverage, or which wishes any expansion of dialysis services to be approved for coverage in accordance with subpart U of part 405, must secure the Secretary’s determination thereunder. In addition to the certification by the State agency referred to in § 488.12 of this part, data furnished by network organizations and recommendations of the Public Health Service, concerning the contribution of a facility to the furnishing of end-stage renal disease services in its network and concerning the facility’s compliance with professional norms and standards (see subpart U of part 405), shall be considered by the Secretary in determining whether to approve a facility for coverage or for any expansion of services under the End-Stage Renal Disease Program. The facility will also be required to submit data pertaining to its qualifications for approval or for any expansion of services, for consideration in the Secretary’s determination.

(b) Determining compliance with minimal utilization rates: Time limitations—(1) Unconditional status. A facility which meets minimal utilization requirements will be assigned this status as long as it continues to meet these requirements.

(2) Conditional status. A conditional status may be granted to a facility for not more than four consecutive calendar years and will not be renewable (see § 405.2122(b) of this chapter). Its status may be examined each calendar year to ascertain its compliance with Subpart U.

(3) Exception status. Under unusual circumstances (see § 405.2122(b) of this chapter) the Secretary may grant a time-limited exception to a facility which is not in compliance with the minimal utilization rate(s) for either unconditional status or conditional status. This exception status may be granted, and may be renewed on an annual basis, under circumstances where rigid application of minimal utilization rate requirements would adversely affect the achievement of ESRD program objectives.

(c) New applicant. A facility which has not previously participated in the ESRD program must submit a plan detailing how it expects to meet the conditional minimal utilization rate status by the end of the second calendar year of its operation under the program and meet the unconditional minimal utilization rate status by the end of the fourth calendar year of its operation under the program.

(d) Notification. The Secretary will notify each facility and its network coordinating council of its initial and its subsequent minimal utilization rate classification.

(e) Failure to meet minimal utilization rate. A facility failing to meet standards for unconditional status or conditional status, or if applicable, for exception status, will be so notified at the time of such classification.

(f) Interim regulations participant. A facility previously participating under the interim regulations will not be approved under the program established by subpart U until it has demonstrated that it meets all the applicable requirements of this subpart, including the appropriate minimal utilization rate. It may continue under the interim program only for a period not to exceed 1 year from the effective date of these amendments (see § 405.2100(c) of this chapter). During this period it may demonstrate its ability to meet the appropriate minimal utilization rate. Failure to qualify under this subpart will automatically terminate coverage of such facility’s services under the ESRD program at the end of such year.

§ 488.64 Remote facility variances for utilization review requirements.

(a) As used in this section:

1. An “available” individual is one who:
   (i) Possesses the necessary professional qualifications;
   (ii) Is not precluded from participating by reason of financial interest in any such facility or direct responsibility for the care of the patients being reviewed or, in the case of a skilled nursing facility, employment by the facility; and
   (iii) Is not precluded from effective participation by the distance between the facility and his residence, office, or other place of work. An individual whose residence, office, or other place of work is more than approximately one hour’s travel time from the facility shall be considered precluded from effective participation.

2. “Adjacent facility” means a health care facility located within a 50-mile radius of the facility which requests a variance.

(b) The Secretary may grant a requesting facility a variance from the time frames set forth in §§ 405.1137(d) and 482.30 as applicable, within which reviews of all cases must be commenced and completed, upon a showing satisfactory to the Secretary that the requesting facility has been unable to meet one or more of the requirements of § 405.1137 of this chapter or § 482.30 of this chapter, as applicable, by reason of insufficient medical and other professional personnel available to conduct the utilization review required by § 405.1137 of this chapter or § 482.30 of this chapter, as applicable.

(c) The request for variance shall document the requesting facility’s inability to meet the requirements for which a variance is requested and the facility’s good faith efforts to comply with the requirements contained in § 405.1137 of this chapter or § 482.30 of this chapter, as applicable.

(d) The request shall include an assurance by the requesting facility that it will continue its good faith efforts to meet the requirements contained in § 405.1137 of this chapter or § 482.30 of this chapter, as applicable.

(e) A revised utilization review plan for the requesting facility shall be submitted concurrently with the request for a variance. The revised plan shall specify the methods and procedures which the requesting facility will use, if a variance is granted, to assure:

1. That effective and timely control will be maintained over the utilization of services; and

2. That reviews will be conducted so as to improve the quality of care provided to patients.

(f) The request for a variance shall include:

1. The name, location, and type (e.g., hospital, skilled nursing facility) of the facility for which the variance is requested;

2. The total number of patient admissions and average daily patient census at the facility within the previous six months;

3. The total number of title XVIII and title XIX patient admissions and the average daily patient census of title XVIII and title XIX patients in the facility within the previous six months;

4. As relevant to the request, the names of all physicians on the active staff of the facility and the names of all other professional personnel on the staff of the facility, or both;

5. The name, location, and type of each adjacent facility (e.g., hospital, skilled nursing facility);

6. The distance and average travel time between the facility and each adjacent facility;

7. As relevant to the request, the location of practice of available physicians and the estimated number of other available professional personnel, or both (see paragraph (a)(1)(iii) of this section);

8. Documentation by the facility of its attempt to obtain the services of available physicians or other professional personnel, or both; and

9. A statement of whether a PRO exists in the area where the facility is located.

(g) The Secretary shall promptly notify the facility of the action taken on the request. Where a variance is in effect, the validation of utilization review pursuant to § 405.1137 of this chapter or § 482.30 shall be made with reference to the revised utilization review plan.
§ 488.68 State Agency responsibilities for OASIS collection and data base requirements.

As part of State agency survey responsibilities, the State agency or other entity designated by HCFA has overall responsibility for fulfilling the following requirements for operating the OASIS system:

(a) Establish and maintain an OASIS database. The State agency or other entity designated by HCFA must—

(1) Use a standard system developed or approved by HCFA to collect, store, and analyze data;

(2) Conduct basic system management activities including hardware and software maintenance, system backup, and monitoring the status of the database; and

(3) Obtain HCFA approval before modifying any parts of the HCFA standard system including, but not limited to, standard HCFA-approved—

(i) OASIS data items;

(ii) Record formats and validation edits; and

(iii) Agency encoding and transmission methods.

(b) Analyze and edit OASIS data. The State agency or other entity designated by HCFA must—

(1) Upon receipt of data from an HHA, edit the data as specified by HCFA and ensure that the HHA resolves errors within the limits specified by HCFA;

(2) At least monthly, make available for retrieval by HCFA all edited OASIS records received during that period, according to formats specified by HCFA, and correct and retransmit previously rejected data as needed; and

(3) Analyze data and generate reports as specified by HCFA.

(c) Ensure accuracy of OASIS data. The State agency must audit the accuracy of the OASIS data through the survey process.

(d) Restrict access to OASIS data. The State agency or other entity designated by HCFA must do the following:

(1) Ensure that access to data is restricted except for the transmission of data and reports to—

(i) HCFA;

(ii) The State agency component that conducts surveys for purposes related to this function; and

(iii) Other entities if authorized by HCFA.

(2) Ensure that patient identifiable OASIS data is released only to the extent that it is permitted under the Privacy Act of 1974.

(e) Provide training and technical support for HHAs. The State agency or other entity designated by HCFA must—

(1) Instruct each HHA on the administration of the data set, privacy/confidentiality of the data set, and integration of the OASIS data set into the facility’s own record keeping system;

(2) Instruct each HHA on the use of software to encode and transmit OASIS data to the State;

(3) Specify to a facility the method of transmission of data to the State, and instruct the facility on this method.

(4) Monitor each HHA’s ability to transmit OASIS data.

(5) Provide ongoing technical assistance and general support to HHAs in implementing the OASIS reporting requirements specified in the conditions of participation for home health agencies; and

(6) Carry out any other functions as designated by HCFA necessary to maintain OASIS data on the standard State system.

[64 FR 3763, Jan. 25, 1999]
§ 488.100 Long term care survey forms, Part A.

PART A — ADMINISTRATIVE AND PROCEDURAL REQUIREMENTS

MEDICARE / MEDICAID SKILLED NURSING FACILITY AND INTERMEDIATE CARE FACILITY SURVEY REPORT

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<th>PROVIDER NUMBER</th>
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| VENDOR NUMBER   |                                                  |
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| SURVEY DATE      |                                                  |
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Form HCFA-255 (2-06)
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<th>EXPLANATORY STATEMENT</th>
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<td>C. Compliance with Other Laws</td>
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<td>F510 The facility is in compliance with applicable Federal, State and local laws and regulations relating to fire and safety, sanitation, communicable and reportable diseases, postmortem procedures and other relevant health and safety requirements.</td>
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</tr>
<tr>
<td>CODE</td>
<td>COMPLIANCE WITH STATE AND LOCAL LAWS/ GOVERNING BODY AND MANAGEMENT</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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</tr>
<tr>
<td>F512</td>
<td>Construction, maintenance and equipment.</td>
<td></td>
<td></td>
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<td>Exception: Not applicable to SNFs.</td>
</tr>
<tr>
<td>F513</td>
<td>Current reports from all responsible governmental agencies are retained at the facility.</td>
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<tr>
<td>F514</td>
<td>Governing Body and Management (Condition of Participation)</td>
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</tr>
<tr>
<td>F515</td>
<td>SNF (405.1121) (a) (Standard)</td>
<td>MET</td>
<td>NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F516</td>
<td>SNF (405.1121)(c) (Standard)</td>
<td>MET</td>
<td>NOT MET</td>
<td></td>
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</tr>
<tr>
<td>F517</td>
<td>1. Written bylaws address the operation of the facility.</td>
<td></td>
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<tr>
<td>F518</td>
<td>2. Written bylaws and policies address effective resident care.</td>
<td></td>
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</tr>
<tr>
<td>F519</td>
<td>3. Bylaws are reviewed and revised as necessary.</td>
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</tr>
</tbody>
</table>

The facility is in compliance with applicable regulations pertaining to:

- Buying, dispensing, safeguarding, administering, and disposing of medications and controlled substances.
  
  Exception: Not applicable to SNFs.

A. Disclosure

- Full disclosure of ownership has been made in accordance with requirements at 42 CFR 420.206.
### NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>GOVERNING BODY AND MANAGEMENT</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F520</td>
<td>ICF (442.301) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F521</td>
<td>SNF (405.1121(d)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td>The facility has policies which ensure that the facility cooperates in an effective program for regular independent medical evaluations and audit of residents in the facility to the extent required by the programs in which the facility participates.</td>
</tr>
<tr>
<td>F522</td>
<td>SNF (405.1121(d)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F523</td>
<td>ICF (442.303) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
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</tr>
<tr>
<td>F524</td>
<td>The facility has a licensed administrator who has authority for the overall operation of the facility. (Administrator's license or registration number)</td>
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</tr>
<tr>
<td>F525</td>
<td>ICF (442.304) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F526</td>
<td>1. The administrator or another professional staff member is the resident care director (RSD).</td>
<td></td>
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</tr>
<tr>
<td>F527</td>
<td>2. The RSD coordinates and monitors each resident's care.</td>
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<tr>
<td>NAME OF FACILITY</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>CODE</th>
<th>GOVERNING BODY AND MANAGEMENT</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F528</td>
<td>SNF (405.1121(l)) (Standard)</td>
<td>MET</td>
<td></td>
<td></td>
<td>F. Institutional Planning</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>1. The facility has an overall plan and budget prepared by a committee of representatives from the governing body, administrative staff, and the organized medical staff (if any).</td>
</tr>
<tr>
<td></td>
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<td>2. The overall plan and budget is reviewed and updated at least annually.</td>
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<td>3. The plan includes a capital expenditures plan, if necessary.</td>
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<td></td>
<td>G. Personnel Policies and Procedures</td>
</tr>
<tr>
<td></td>
<td>SNF (405.1121(l)) (Standard)</td>
<td>MET</td>
<td></td>
<td></td>
<td>1. The facility has written policies and procedures that support sound resident care and personnel practices and address, at least:</td>
</tr>
<tr>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>a. Control of communicable disease;</td>
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<td></td>
<td></td>
<td>b. The review of employee incidents and accidents to identify health and safety hazards; and</td>
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<td></td>
<td></td>
<td>c. The existence of a safe and sanitary environment.</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>2. Personnel records are current, available to each employee, and contain sufficient information to support placement in the position to which assigned.</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>3. Referral or provision for periodic health examinations to ensure freedom from communicable disease.</td>
</tr>
</tbody>
</table>

Form: HCFA-505 (2-96)
### NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>GOVERNING BODY AND MANAGEMENT</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><strong>H. Outside Resources/Consultant Agreements</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>F538 SNF (405.1121) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
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<td></td>
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<tr>
<td></td>
<td>F539 ICF (442.317) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
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<tr>
<td></td>
<td>F540</td>
<td></td>
<td></td>
<td></td>
<td>The facility has written agreements with qualified persons to render a service (if it does not employ a qualified professional person to do so). The agreements:</td>
</tr>
<tr>
<td></td>
<td>F541</td>
<td></td>
<td></td>
<td></td>
<td>1. Address the responsibilities, functions, objectives, and terms (including financial arrangements and charges);</td>
</tr>
<tr>
<td></td>
<td>F542</td>
<td></td>
<td></td>
<td></td>
<td>2. Are signed by an authorized representative of the facility and the outside resource; and</td>
</tr>
<tr>
<td></td>
<td>F543</td>
<td></td>
<td></td>
<td></td>
<td>3. Specify that the facility retains ultimate responsibility for the services rendered.</td>
</tr>
<tr>
<td></td>
<td><strong>I. Notification of Change in Resident Status</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td>F544 SNF (405.1121) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
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<tr>
<td></td>
<td>F545</td>
<td></td>
<td></td>
<td></td>
<td>The facility has policies and procedures to notify physicians and other responsible persons in the event of an accident involving the resident, or resident's physical, mental or emotional status, or resident changes, billings or related administrative matter.</td>
</tr>
<tr>
<td>CODE</td>
<td>GOVERNING BODY AND MANAGEMENT</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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<tr>
<td>F546</td>
<td>J. Resident Rights (Standard)</td>
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<tr>
<td></td>
<td>SNF (405.1121(k))</td>
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<td></td>
<td>Indicators 1 thru 12 apply to SNFs.</td>
</tr>
<tr>
<td>F547</td>
<td>ICF (442.311) (Standard)</td>
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<td></td>
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<td></td>
<td></td>
<td>1. Information</td>
</tr>
<tr>
<td></td>
<td>a. The facility informs each resident, before or at the time of admission, of his rights and responsibilities.</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>b. The facility informs each resident, before or at the time of admission, of all rules governing resident conduct.</td>
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<tr>
<td></td>
<td>c. The facility informs each resident of amendments to their policies on residents' rights and responsibilities and rules governing conduct.</td>
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<tr>
<td></td>
<td>d. Each resident acknowledges in writing receipt of residents' rights information and any amendment to it.</td>
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<tr>
<td></td>
<td>e. The resident must be informed in writing of all services and charges for services.</td>
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<tr>
<td></td>
<td>f. The resident must be informed in writing of all changes in services and charges before or at the time of admission and on a continuing basis.</td>
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<td></td>
<td>g. The resident must be informed of services not covered by Medicare or Medicaid in the basic rate.</td>
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</tr>
<tr>
<td>CODE</td>
<td>GOVERNING BODY AND MANAGEMENT</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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</tr>
<tr>
<td></td>
<td>2. Medical Condition and Treatment</td>
<td></td>
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<tr>
<td>F555</td>
<td>a. Each resident is informed by a physician of his health and medical condition unless the physician decides that informing the resident is medically contraindicated.</td>
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</tr>
<tr>
<td>F556</td>
<td>b. Each resident is given an opportunity to participate in planning his total care and medical treatment.</td>
<td></td>
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<tr>
<td>F557</td>
<td>c. Each resident is given an opportunity to refuse treatment.</td>
<td></td>
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<tr>
<td>F558</td>
<td>d. Each resident gives informed, written consent before participating in experimental research.</td>
<td></td>
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<tr>
<td>F559</td>
<td>e. If the physician decides that informing the resident of his health and medical condition is medically contraindicated, the physician has documented this decision in the resident's medical record.</td>
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<tr>
<td></td>
<td>3. Transfer and Discharge</td>
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<tr>
<td></td>
<td>Each resident is transferred or discharged only for:</td>
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<tr>
<td>F560</td>
<td>a. Medical reasons.</td>
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<tr>
<td>F561</td>
<td>b. His/her welfare or that of other residents.</td>
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<tr>
<td>F562</td>
<td>c. Nonpayment except as prohibited by the Medicare or Medicaid program.</td>
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<tr>
<td></td>
<td>4. Exercising Rights</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F563</td>
<td>a. Each resident is encouraged and assisted to exercise his/her rights as a resident of the facility and as a citizen.</td>
<td></td>
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</tr>
<tr>
<td>F564</td>
<td>b. Each resident is allowed to submit complaints and recommendations concerning the policies and services of the facility to staff or to outside representatives of the resident's choice or both.</td>
<td></td>
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</tr>
<tr>
<td>F565</td>
<td>c. Such complaints are submitted free from restraint, coercion, discrimination, or reprisal.</td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>CODE</td>
<td>GOVERNING BODY AND MANAGEMENT</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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<tr>
<td></td>
<td>5. Financial Affairs</td>
<td></td>
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</tr>
<tr>
<td>F566</td>
<td>a. Residents are allowed to manage their own personal financial affairs.</td>
<td></td>
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<tr>
<td></td>
<td>b. The facility establishes and maintains a system that assures full and complete accounting of residents' personal funds. An accounting report is made to residents in skilled nursing facilities at least on a quarterly basis.</td>
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<tr>
<td></td>
<td>c. The facility does not commingle resident funds with any other funds other than resident funds.</td>
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<tr>
<td></td>
<td>d. If a resident requests assistance from the facility in managing his personal financial affairs, resident's delegation is in writing.</td>
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</tr>
<tr>
<td></td>
<td>e. The facility system of accounting includes written receipts for:</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td>1. All personal possessions and funds received by or deposited with the facility.</td>
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</tr>
<tr>
<td></td>
<td>2. All disbursement made to or for the resident.</td>
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<tr>
<td></td>
<td>f. The financial record must be available to the resident and his/her family.</td>
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<tr>
<td></td>
<td>6. Freedom from Abuse and Restraints</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>a. Each resident is free from mental and physical abuse</td>
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</tr>
<tr>
<td></td>
<td>b. Chemical and physical restraints are only used when authorized by a physician in writing for a specified period of time or in emergencies.</td>
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<tr>
<td></td>
<td>c. If used in emergencies, they are necessary to protect the resident from injury to himself or others.</td>
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</tr>
</tbody>
</table>
### NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>GOVERNING BODY AND MANAGEMENT</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F576</td>
<td>d. The use is authorized by a professional staff member identified in the written policies and procedures of the facility.</td>
<td></td>
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</tr>
<tr>
<td>F577</td>
<td>e. The use is reported promptly to the resident's physician by the staff member.</td>
<td></td>
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</tr>
<tr>
<td>F578</td>
<td>a. Each resident is treated with respect, consideration and full recognition of his/her dignity and individuality.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F579</td>
<td>b. Each resident is given privacy during treatment and care of personal needs.</td>
<td></td>
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</tr>
<tr>
<td>F580</td>
<td>c. Each resident's records, including information in an automated data bank, are treated confidentially.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F581</td>
<td>d. Each resident must give written consent before the facility releases information from his/her record to someone not otherwise authorized to receive it.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F582</td>
<td>e. Married residents are given privacy during visits by their spouses.</td>
<td></td>
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</tr>
<tr>
<td>F583</td>
<td>f. Married residents are permitted to share a room.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F584</td>
<td>8. Work</td>
<td>No resident may be required to perform services for the facility.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F585</td>
<td>9. Freedom of Association and Correspondence</td>
<td>a. Each resident is allowed to communicate, associate and meet privately with individuals of his choice unless this intrudes upon the rights of another resident.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F586</td>
<td>b. Each resident is allowed to send and receive personal mail unopened.</td>
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<td></td>
</tr>
<tr>
<td>CODE</td>
<td>GOVERNING BODY AND MANAGEMENT</td>
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<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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</tr>
<tr>
<td>F587</td>
<td>10. Activities</td>
<td></td>
<td></td>
<td></td>
<td>Each resident is allowed to participate in social, religious, and community group activities.</td>
</tr>
<tr>
<td>F588</td>
<td>11. Personal Possessions</td>
<td></td>
<td></td>
<td></td>
<td>Each resident is allowed to retain and use his personal possessions and clothing as space permits.</td>
</tr>
<tr>
<td>F589</td>
<td>12. Written Policies and Procedures: Delegation of Rights and Responsibilities</td>
<td></td>
<td></td>
<td></td>
<td>ICF (442.312) (Standard)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>MET</td>
<td></td>
<td>NOT MET</td>
<td>a. The facility has written policies and procedures that provide that all the rights and responsibilities of a resident pass to the resident’s guardian, next of kin or sponsoring agency or agencies if the resident is adjudicated incompetent under State law or is determined by his physician to be incapable of understanding his rights and responsibilities.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>b. Physician determinations of incapability and the specific reasons thereof are recorded by the physician in the resident’s record.</td>
</tr>
<tr>
<td>F592</td>
<td>K. Resident Care Policies</td>
<td></td>
<td></td>
<td></td>
<td>SNF (405.1121)(g) (Standard)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>MET</td>
<td></td>
<td>NOT MET</td>
<td>1. The facility has written policies to govern the continuing skilled nursing care and related medical or other services provided.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2. Those policies reflect awareness of and provision for meeting the total medical and psychosocial needs of residents including admission, transfer, discharge planning, and the range of services available to residents; and</td>
</tr>
<tr>
<td>CODE</td>
<td>GOVERNING BODY AND MANAGEMENT</td>
<td>YES</td>
<td>NO</td>
<td>WA</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
<tr>
<td>------</td>
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</tr>
<tr>
<td>F595</td>
<td>3. The protection of residents' personal and property rights.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F596</td>
<td>4. The policies are developed by a group of professional personnel, including the Medical Director or the organized medical staff, and are periodically reviewed and revised (if necessary).</td>
<td></td>
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</tr>
<tr>
<td>F597</td>
<td>5. These policies are available to admitting physicians, sponsoring agencies, residents, and the public.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F598</td>
<td>6. The Medical Director or a registered nurse is designated as responsible for the execution of the policies.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L. Public Availability</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>F599</td>
<td>ICF (442.305) (Standard)</td>
<td>MET</td>
<td>NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F600</td>
<td>1. The facility has written policies and procedures governing all the services it provides.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F601</td>
<td>2. The policies and procedures are available to the staff and residents, members of the family, the public, and legal representatives of residents.</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>M. Admissions</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F602</td>
<td>ICF (442.306) (Standard)</td>
<td>MET</td>
<td>NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F603</td>
<td>The facility has written policies and procedures that ensure that it admits as residents only those residents whose needs can be met by:</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F604</td>
<td>1. the facility itself.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F605</td>
<td>2. the facility in cooperation with community resources.</td>
<td></td>
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</tr>
<tr>
<td>F606</td>
<td>3. the facility in cooperation with other providers of care affiliated with or under contract to the facility.</td>
<td></td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

Form HCFA-625 (2.66)
### Name of Facility

<table>
<thead>
<tr>
<th>CODE</th>
<th>Governing Body and Management</th>
<th>Yes</th>
<th>No</th>
<th>N/A</th>
<th>Explanatory Statement</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N. Transfers</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F606</td>
<td>ICF (442.307) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F607</td>
<td>1. The facility has written policies and procedures to ensure that residents are transferred promptly to a hospital, SNF, or other appropriate facility when a change is necessary.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F608</td>
<td>2. Except in emergencies, the facility consults the resident, his next of kin, the attending physician, and the responsible agency, if any, at least five days before discharge.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F609</td>
<td>3. The facility uses casework services and other means to ensure that adequate arrangements are made to meet resident's needs through other resources.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

|      | O. Restraints                  |     |    |     |                       |
| F610 | ICF (442.308) (Standard)       | ☐ MET | ☐ NOT MET | |                       |
| F611 | The facility has written policies and procedures that: | | | |                       |
| F612 | 1. Define the uses of chemical and physical restraints. | | | |                       |
| F613 | 2. Identify the professional personnel who may authorize the use of restraints in emergencies under 442.311(f). | | | |                       |
| F614 | 3. Describe procedures for monitoring and controlling the use of these restraints. | | | |                       |

<p>|      | P. Complaints                  |     |    |     |                       |
| F615 | ICF (442.309) (Standard)       | ☐ MET | ☐ NOT MET | |                       |
| F616 | The facility has written policies and procedures that: | | | |                       |
| F617 | 1. Describe the procedures the facility uses to receive complaints and recommendations from residents. | | | |                       |
| F618 | 2. Ensure that the facility responds to complaints and recommendations. | | | |                       |</p>
<table>
<thead>
<tr>
<th>CODE</th>
<th>GOVERNING BODY AND MANAGEMENT</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F617</td>
<td>SNF (405.1121(h)) (Standard)</td>
<td></td>
<td></td>
<td></td>
<td>1. The facility conducts an orientation program for all new employees that includes a review of all its policies.</td>
</tr>
<tr>
<td>F618</td>
<td>ICF (442.314) (Standard)</td>
<td></td>
<td></td>
<td></td>
<td>2. The facility plans and conducts an inservice staff development program for all personnel to assist them in developing and improving their skills.</td>
</tr>
<tr>
<td>F619</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>3. The facility maintains a record of the orientation and staff development programs it conducts.</td>
</tr>
<tr>
<td>F620</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>4. The record includes the content of the program and the names of participants.</td>
</tr>
<tr>
<td>F621</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>5. Inservice training includes at least prevention and control of infections, fire prevention and safety, confidentiality of resident information, and preservation of resident dignity including protection of resident's privacy and personal and property rights.</td>
</tr>
<tr>
<td>CODE</td>
<td>MEDICAL DIRECTION</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
<tr>
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<tr>
<td>F624</td>
<td>Medical Direction (Condition of Participation)</td>
<td></td>
<td></td>
<td></td>
<td>The facility has a written agreement with a licensed physician to serve as Medical Director on a part-time or full-time basis as is appropriate to the needs of the residents and the facility. (See 405.1911(b) regarding waiver of this requirement.)</td>
</tr>
<tr>
<td></td>
<td>SNF (405.1122)</td>
<td></td>
<td></td>
<td></td>
<td>MET</td>
</tr>
</tbody>
</table>

A. Coordination of Medical Care

<table>
<thead>
<tr>
<th>F625</th>
<th>SNF (405.1122)(a) (Standard)</th>
<th>MET</th>
<th>NOT MET</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1. Medical direction and coordination of medical care in the facility are provided by a Medical Director.</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>2. The Medical Director is responsible for development of policies approved by the governing body.</td>
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<tr>
<td></td>
<td>3. Coordination of medical care includes liaison with attending physicians to ensure their writing orders promptly upon admission of a resident, and periodic evaluation of the adequacy and appropriateness of health professional and supportive staff and services.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

B. Responsibilities to the Facility

<table>
<thead>
<tr>
<th>F629</th>
<th>SNF (405.1122)(b) (Standard)</th>
<th>MET</th>
<th>NOT MET</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1. The Medical Director is responsible for surveillance of the health status of the facility's employees.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2. Incidents and accidents that occur on the premises are reviewed by the Medical Director to identify hazards to health and safety.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>PHYSICIAN SERVICES</td>
<td>(Condition of Participation)</td>
<td></td>
</tr>
<tr>
<td>------</td>
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<td></td>
</tr>
<tr>
<td>F33</td>
<td>SNF (485.1123)</td>
<td>NOT MET</td>
<td></td>
</tr>
</tbody>
</table>

The facility must be a facility that is under the supervision of a physician. To the extent such medical care is required and is not provided by the facility, the facility must make arrangements with a physician to furnish necessary medical care in case of emergency.

<table>
<thead>
<tr>
<th>CODE</th>
<th>PHYSICIAN SUPERVISION</th>
<th>Standard</th>
<th>(Standard)</th>
<th>MET</th>
<th>NOT MET</th>
</tr>
</thead>
<tbody>
<tr>
<td>F44</td>
<td>ICF (485.1124)</td>
<td>Standard</td>
<td>Standard</td>
<td>NOT MET</td>
<td>NOT MET</td>
</tr>
<tr>
<td>F45</td>
<td>SNF (485.1125)</td>
<td>Standard</td>
<td>Standard</td>
<td>NOT MET</td>
<td>NOT MET</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>CODE</th>
<th>EMERGENCY SERVICES</th>
<th>Standard</th>
<th>(Standard)</th>
<th>MET</th>
<th>NOT MET</th>
</tr>
</thead>
<tbody>
<tr>
<td>F63</td>
<td>SNF (485.1126)</td>
<td>Standard</td>
<td>Standard</td>
<td>NOT MET</td>
<td>NOT MET</td>
</tr>
<tr>
<td>NAME OF FACILITY</td>
<td></td>
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<td></td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>CODE</th>
<th>NURSING SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F638</td>
<td>Nursing Services (Condition of Participation)</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>F639</td>
<td>ICF (442.342) (Standard)</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

A. Director of Nursing Services

<table>
<thead>
<tr>
<th>F640</th>
<th>SNF (405.1124(a)) (Standard)</th>
<th>☐</th>
<th>☐</th>
<th>☐</th>
<th>☐</th>
</tr>
</thead>
<tbody>
<tr>
<td>F641</td>
<td>1. The director of nursing services is a qualified registered nurse employed full-time.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F642</td>
<td>2. The director of nursing services has, in writing, administrative authority, responsibility, and accountability for the functions, activities, and training of the nursing services staff, and serves only one facility in this capacity.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F643</td>
<td>3. If the director of nursing services has other institutional responsibilities, a qualified registered nurse serves as assistant so that there is the equivalent of a full-time director of nursing services on duty.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>NURSING SERVICES</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
<tr>
<td>------</td>
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</tr>
<tr>
<td>F644</td>
<td>ICF (442.339)</td>
<td>MET</td>
<td></td>
<td></td>
<td>1. The facility has a full-time registered nurse, or a licensed practical or vocational nurse to supervise the health services 7 days a week on the day shift.</td>
</tr>
<tr>
<td>F645</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2. The nurse has a current State license.</td>
</tr>
<tr>
<td>F646</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>3. If the supervisor of health services is a licensed practical or vocational nurse, the facility has a formal contract with a registered nurse to serve as a consultant no less than 4 hours a week.</td>
</tr>
<tr>
<td>F647</td>
<td></td>
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<td></td>
<td>4. To qualify to serve as a health services supervisor, a licensed practical or vocational nurse must:</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>a. Have graduated from a State-approved school of practical nursing, or</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>b. Have education or other training that the State authority responsible for licensing practical nurses considered equal to graduation from a State-approved school of practical nursing, or</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>c. Have passed the Public Health Service examination for waived licensed practical or vocational nurses.</td>
</tr>
<tr>
<td>F650</td>
<td></td>
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<td></td>
<td>5. If the nurse in charge is licensed by the State in a category other than registered nurse or licensed practical or vocational nurse:</td>
</tr>
<tr>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>a. The individual has completed a training program to get the license that includes at least the same number of classroom and practice hours in all nursing subjects as in the program of a State-approved school of practical or vocational nursing, and</td>
</tr>
</tbody>
</table>
### NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>NURSING SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>NIA</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F652</td>
<td>b. The State agency responsible for licensing the individual submits a report to the Medicaid agency comparing State-licensed practical nurse or vocational nurse course requirements with those for the program completed by the individual.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F653</td>
<td>SNF (405.112)(c)(2) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td>C. Twenty-four Hour Nursing Service</td>
</tr>
<tr>
<td>F654</td>
<td>ICF (442.338) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### F655 1. 24-Hour Nursing
- Nursing policies and procedures address the total nursing needs of the residents.
- The policies are designed to ensure that each resident receives:
  - Treatment.
- Medications as prescribed.
- Diet as prescribed.
- Rehabilitative nursing care as needed.
- Proper care to prevent decubitus ulcers and deformities.
- Proper care to ensure that residents are clean, well-groomed and comfortable.
- Protection from accident and injury.
- Protection from infection.
- Encouragement, assistance, and training in self-care and group activities.

Form HHC-515 (4-96)
<table>
<thead>
<tr>
<th>CODE</th>
<th>NURSING SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F665</td>
<td>2. Weekly time schedules are maintained and indicate the number and classifications of nursing personnel including relief personnel, who worked on each unit for each tour of duty.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F666</td>
<td>D. Rehabilitative Nursing Care</td>
<td>☐ ME T</td>
<td>☐ NOT ME T</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F667</td>
<td>SNF (405.1124(g)) (Standard)</td>
<td>☐ ME T</td>
<td>☐ NOT ME T</td>
<td></td>
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</tr>
<tr>
<td>F668</td>
<td>Nursing personnel are trained in rehabilitative nursing.</td>
<td></td>
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</tr>
<tr>
<td>F669</td>
<td>E. Supervision of Resident Nutrition</td>
<td>☐ ME T</td>
<td>☐ NOT ME T</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F670</td>
<td>SNF (405.1124(f)) (Standard)</td>
<td>☐ ME T</td>
<td>☐ NOT ME T</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F671</td>
<td>A procedure is established to inform dietician service of physicians' diet orders and of residents' dietetic problems.</td>
<td></td>
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</tr>
<tr>
<td>F672</td>
<td>F. Administration of Drugs</td>
<td>☐ ME T</td>
<td>☐ NOT ME T</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F673</td>
<td>SNF (405.1124(g)) (Standard)</td>
<td>☐ ME T</td>
<td>☐ NOT ME T</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F674</td>
<td>G. Conformance with Physicians' Drug Orders</td>
<td>☐ ME T</td>
<td>☐ NOT ME T</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F675</td>
<td>Procedures are established by the Pharmaceutical Services Committee (see 405.1127(d)) to ensure that drugs are checked against physicians' orders.</td>
<td></td>
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</tr>
<tr>
<td>F676</td>
<td>SNF (405.1124(h)) (Standard)</td>
<td>☐ ME T</td>
<td>☐ NOT ME T</td>
<td></td>
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</tr>
<tr>
<td>F677</td>
<td>Indicators 1 thru 4 apply to SNFs.</td>
<td></td>
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</tr>
<tr>
<td>F678</td>
<td>ICF (442.335) (Standard)</td>
<td>☐ ME T</td>
<td>☐ NOT ME T</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F679</td>
<td>1. Drugs not specifically limited as to time or number of doses when ordered are controlled by automatic stop orders or other methods in accordance with written policies.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>NURSING SERVICES</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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<td>------</td>
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<tr>
<td>F675</td>
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<td></td>
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<td>2. The attending physician is notified of an automatic stop order prior to the last dose so that the physician may decide if the administration of the drug or biological is to be continued or altered.</td>
</tr>
<tr>
<td>F676</td>
<td>ICF (442.334)</td>
<td></td>
<td></td>
<td></td>
<td>(Standard) MET NOT MET</td>
</tr>
<tr>
<td>F677</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>3. Physicians' verbal orders for drugs are given only to a licensed nurse, pharmacist, or physician and are immediately recorded and signed by the person receiving the order. (Verbal orders for Schedule II drugs are permitted only in the case of a bona fide emergency situation.)</td>
</tr>
<tr>
<td>F678</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>4. Such orders are countersigned by the attending physician within a reasonable time.</td>
</tr>
<tr>
<td></td>
<td>H. Storage of Drugs and Biologicals</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F679</td>
<td>SNF (405.1124(e))</td>
<td></td>
<td></td>
<td></td>
<td>(Standard) MET NOT MET</td>
</tr>
<tr>
<td>F680</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1. Procedures for storing and disposing of drugs and biologicals are established by the pharmaceutical services committee.</td>
</tr>
<tr>
<td>F681</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2. In accordance with State and Federal laws, all drugs and biologicals are stored in locked compartments under proper temperature controls.</td>
</tr>
<tr>
<td>F682</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>3. Only authorized personnel have access to the keys.</td>
</tr>
<tr>
<td>F683</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>4. Separately locked, permanently affixed compartments are provided for storage of controlled drugs listed in Schedule II of the Comprehensive Drug Abuse Prevention &amp; Control Act of 1970 and other drugs subject to abuse, except under single unit dosage distribution systems in which the quantity stored is minimal and a missing dose can be readily detected.</td>
</tr>
<tr>
<td>F684</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>5. An emergency medication kit approved by the pharmaceutical services committee is kept readily available.</td>
</tr>
</tbody>
</table>
### Diabetic Services (Condition of Participation)

<table>
<thead>
<tr>
<th>CODE</th>
<th>DIETETIC SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F685</td>
<td>SNF (405.1125)</td>
<td></td>
<td></td>
<td></td>
<td>The facility provides a hygienic dietetic service that meets the daily nutritional needs of patients, ensures that special dietary needs are met, and provides palatable and attractive meals. A facility that has a contract with an outside food management company may be found to be in compliance with this condition provided the facility and/or company meets the standards listed herein.</td>
</tr>
</tbody>
</table>

#### A. Staffing

<p>| F686  | SNF (405.1125(a)) (Standard) |     |    |     | 1. Overall supervisory responsibility for the dietetic service is assigned to a full-time qualified dietetic service supervisor. |
|-------|------------------------------|-----|----|-----| 2. If the dietetic service supervisor is not a qualified dietician, the dietetic service supervisor functions with frequent, regularly scheduled consultation from a person so qualified. (§405.1101(a).) |
| F687  | SNF (405.1125(a)) (Standard) |     |    |     | 3. In addition, the facility employs sufficient supportive personnel competent to carry out the functions of the dietetic service. |
| F688  | SNF (405.1125(a)) (Standard) |     |    |     | 4. If consultant dietetic services are used, the consultant's visits are at appropriate times, and of sufficient duration and frequency to provide continuing liaison with medical and nursing staffs, advice to the administrator, resident counseling, guidance to the supervisor and staff of the dietetic service, approval of all menus, and participation in the development or revisions of dietetic policies and procedures. (See §405.1121(i).) |</p>
<table>
<thead>
<tr>
<th>NAME OF FACILITY</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>CODE</th>
<th>DIETETIC SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
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</thead>
<tbody>
<tr>
<td>F691</td>
<td>ICF (442.332) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
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</tr>
<tr>
<td>F692</td>
<td>1. The facility has a staff member trained or experienced in food management or nutrition who is responsible for:</td>
<td></td>
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</tr>
<tr>
<td>F693</td>
<td>a. Planning meals that meet the nutritional needs of each resident.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F694</td>
<td>b. Following the orders of the resident’s physician.</td>
<td></td>
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</tr>
<tr>
<td>F695</td>
<td>c. To the extent medically possible, following the recommended dietary allowances of the Food and Nutrition Board of the National Research Council, National Academy of Sciences (Recommended Dietary Allowances, 8th Ed., 1974).</td>
<td></td>
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</tr>
<tr>
<td>F696</td>
<td>d. Supervising the meal preparation and service to ensure that the menu plan is followed.</td>
<td></td>
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</tr>
<tr>
<td>F697</td>
<td>2. For residents who required medically prescribed special diets, the facility:</td>
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</tr>
<tr>
<td>F698</td>
<td>a. Has menus for those residents planned by a professionally qualified dietitian or reviewed and approved by the attending physician; and</td>
<td></td>
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<tr>
<td>F699</td>
<td>b. Supervises the preparation and serving of meals to ensure that the resident accepts the special diet.</td>
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<tr>
<td>F700</td>
<td>3. The facility keeps for 30 days a record of each menu as served.</td>
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</tr>
<tr>
<td>CODE</td>
<td>DIETETIC SERVICES/ SPECIALIZED REHABILITATION SERVICES</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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<tr>
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</tr>
<tr>
<td>F699</td>
<td>C. Hygiene of Staff</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td></td>
<td>SNF (405.1125(j)) (Standard)</td>
<td>MET</td>
<td>NOT</td>
<td>MET</td>
<td></td>
</tr>
<tr>
<td>F700</td>
<td>In the event food service employees are assigned duties outside the dietetic service, these duties do not interfere with the sanitation, safety, or the time required for dietetic work assignments. (See §405.1121(g).)</td>
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<tr>
<td>F701</td>
<td>D Sanitary Conditions</td>
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<tr>
<td></td>
<td>SNF (405.1125(g)) (Standard)</td>
<td>MET</td>
<td>NOT</td>
<td>MET</td>
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<tr>
<td>F702</td>
<td>Written reports of inspections by State and local health authorities are on file at the facility, with notation made of action taken by the facility to comply with any recommendations.</td>
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<tr>
<td>F703</td>
<td>Specialized Rehabilitation Services (Condition of Participation)</td>
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<tr>
<td></td>
<td>SNF (405.1126)</td>
<td>MET</td>
<td>NOT</td>
<td>MET</td>
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<tr>
<td></td>
<td>The facility provides, or arranges for, under written agreement, specialized rehabilitative services by qualified personnel (i.e., physical therapy, speech pathology and audiology, and occupational therapy) as needed by residents to improve and maintain functioning. Safe and adequate space and equipment are available, commensurate with the services offered. If the facility does not offer such services directly, it does not admit nor retain residents in need of this care unless provision is made for such services under arrangement with qualified outside resources under which the facility assumes professional responsibility for the services rendered. (See §405.1121(i).)</td>
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<tr>
<td>CODE</td>
<td>SPECIALIZED REHABILITATION SERVICES</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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<tr>
<td>F704</td>
<td>Staffing and Organization</td>
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<tr>
<td></td>
<td>SNF (405.1126(a)) (Standard)</td>
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<td></td>
<td></td>
<td>Indicators 1 thru 3 apply to SNFs</td>
</tr>
<tr>
<td>F705</td>
<td>ICF (442.343) (Standard)</td>
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<tr>
<td>F706</td>
<td>1. Specialized rehabilitative services are provided, in accordance with accepted professional practices, by qualified therapists or by qualified assistants or other supportive personnel under the supervision of qualified therapists.</td>
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<tr>
<td>F707</td>
<td>2. Other rehabilitative services also may be provided, but must be in a facility where all rehabilitative services are provided through an organized rehabilitative service under the supervision of a physician qualified in physical medicine who determines the goals and limitations of these services and assigns duties appropriate to the training and experience of those providing such services.</td>
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<td></td>
<td>Exception: Does not apply to ICFs.</td>
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<tr>
<td>F708</td>
<td>3. Written administrative and resident care policies and procedures are developed for rehabilitative services by appropriate therapists and representatives of the medical, administrative, and nursing staffs.</td>
<td></td>
<td></td>
<td></td>
<td>Exception: Does not apply to ICF's See General Requirements 442.305</td>
</tr>
</tbody>
</table>
NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>SPECIALIZED REHABILITATION SERVICES/ PHARMACEUTICAL SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F709</td>
<td>SNF (405.1126(d)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td>The physician's order, the plan of rehabilitative care, services rendered, evaluations of progress, and other pertinent information are recorded in the patient's medical record, and are dated and signed by the physician ordering the service and the person who provided the service.</td>
</tr>
<tr>
<td>F710</td>
<td>SNF (405.1126(d)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td>If the facility provides outpatient physical therapy services, it meets the applicable health and safety regulations pertaining to such services as are included in Subpart Q of this part. (See §405.1719, 405.1720, 405.1722(a) and (b)(1)(3)(9), (4), (10), (7), and (8); and 405.1725.)</td>
</tr>
<tr>
<td>F711</td>
<td>SNF (405.1127)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td>The facility has appropriate methods and procedures for the dispensing and administering of drugs and biologicals. The facility is responsible for providing such drugs and biologicals for its residents, to the extent that they are covered under the programs, and for ensuring that pharmaceutical services are provided in accordance with accepted professional principles.</td>
</tr>
<tr>
<td>CODE</td>
<td>PHARMACEUTICAL SERVICES</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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</tr>
<tr>
<td>A. Supervision of Services</td>
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<tr>
<td>F712</td>
<td>SNF (405.1127(a)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
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</tr>
<tr>
<td>F713</td>
<td>1. The pharmaceutical services are under the general supervision of a qualified pharmacist.</td>
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<tr>
<td>F714</td>
<td>2. The pharmacist is responsible to the administrative staff for developing coordinating, and supervising all pharmaceutical services.</td>
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</tr>
<tr>
<td>F715</td>
<td>3. The pharmacist (if not a full-time employee) devotes a sufficient number of hours, based upon the needs of the facility, during regularly scheduled visits to carry out these responsibilities.</td>
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<tr>
<td>F716</td>
<td>ICF (442.333) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
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</tr>
<tr>
<td>F717</td>
<td>1. The facility employs a licensed pharmacist, or</td>
<td></td>
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<tr>
<td>F718</td>
<td>2. The facility has formal arrangements with a licensed pharmacist to advise the facility on ordering, storage, administration, disposal and recordkeeping of drugs and biologicals.</td>
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<tr>
<td>B. Control and Accountability</td>
<td></td>
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<tr>
<td>F719</td>
<td>SNF (405.1127(b)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
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</tr>
<tr>
<td>F720</td>
<td>1. The pharmaceutical service has procedures for control and accountability of all drugs and biologicals throughout the facility.</td>
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</tr>
<tr>
<td>F721</td>
<td>2. Only approved drugs and biologicals are used in the facility.</td>
<td></td>
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<tr>
<td>F722</td>
<td>3. Records of receipt and disposition of all controlled drugs are maintained in sufficient detail to enable an accurate reconciliation.</td>
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<tr>
<td>CODE</td>
<td>EXPLANATORY STATEMENT</td>
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<td></td>
</tr>
<tr>
<td>C. Pharmaceutical Services Committee</td>
<td></td>
<td></td>
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<tr>
<td>F723</td>
<td>SNF (405.1127(d)) (Standard)</td>
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<td></td>
<td>□ MET □ NOT MET</td>
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<tr>
<td>F724</td>
<td>1. A pharmaceutical services committee or its equivalent develops written policies and procedures for safe and effective drug therapy, distribution, control and use.</td>
<td></td>
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<tr>
<td>F725</td>
<td>2. The committee is comprised of at least the pharmacist, the director of nursing services, the administrator, and one physician.</td>
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<tr>
<td>F726</td>
<td>3. The committee oversees pharmaceutical services in the facility, makes recommendations for improvement, and monitors the service to ensure its accuracy and adequacy.</td>
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</tr>
<tr>
<td>Laboratory and Radiologic Services (Condition of Participation)</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>F727</td>
<td>SNF (405.1128)</td>
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<td></td>
<td>□ MET □ NOT MET</td>
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<tr>
<td></td>
<td>The facility has provision for promptly obtaining required laboratory, X-ray, and other diagnostic services.</td>
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</tr>
<tr>
<td>A. Provision for Services</td>
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</tr>
<tr>
<td>F728</td>
<td>SNF (405.1128(a)(1)) (Standard)</td>
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<tr>
<td></td>
<td>□ MET □ NOT MET</td>
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<tr>
<td>F729</td>
<td>1. If the facility provides its own laboratory and X-ray services, these meet the applicable conditions established for certification of hospitals that are contained in 405.1028 and 405.1029, respectively.</td>
<td></td>
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</table>
### NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>LABORATORY AND RADIOLOGIC SERVICES/DENTAL SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F730</td>
<td>2. If the facility itself does not provide such services, arrangements are made for obtaining these services from a physician's office, a participating hospital or skilled nursing facility, or a portable X-ray supplier or independent laboratory which is approved to provide these services under the program.</td>
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<tr>
<td>F731</td>
<td>3. The facility assists the resident, if necessary, in arranging for transportation to and from the source of service.</td>
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<tr>
<td>B. Blood and Blood Products</td>
<td></td>
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<tr>
<td>F732</td>
<td>SNF (405.1128(d)) (Standard)</td>
<td>□ MET</td>
<td>□ NOT MET</td>
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</tr>
<tr>
<td>F733</td>
<td>1. Blood handling and storage facilities are safe, adequate, and properly supervised.</td>
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</tr>
<tr>
<td>F734</td>
<td>2. If the facility provides for maintaining and transfusing blood and blood products, it meets the conditions established for certification of hospitals that are contained in §405.1028(b).</td>
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</tr>
<tr>
<td>F735</td>
<td>3. If the facility does not provide its own facility but does provide transfusion services alone, it meets at least the requirements of §405.1028(b)(1), (3), (4), (6), and (9).</td>
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<tr>
<td>Dental Services (Condition of Participation)</td>
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</tr>
<tr>
<td>F736</td>
<td>SNF (405.1129)</td>
<td>□ MET</td>
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</tbody>
</table>

The facility has satisfactory arrangements to assist residents to obtain routine and emergency dental care (See §405.1121(j). The basic Hospital Insurance Program does not cover the services of a dentist in a skilled nursing facility in connection with the care, treatment, filling, removal, or replacement of teeth or structures supporting the teeth; and only certain oral surgery is included in the Supplemental Medical Insurance Program.)
NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>DENTAL SERVICES/SOCIAL SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
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<tbody>
<tr>
<td>F737</td>
<td>A. Advisory Dentist</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>SNF (405.1129(a))</td>
<td></td>
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<tr>
<td></td>
<td>(Standard)</td>
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<td>□ NOT MET</td>
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<tr>
<td>F738</td>
<td>□ MET</td>
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<td></td>
<td>A dentist recommends oral hygiene policies and practices for the care of residents. (§405.1121(h)).</td>
</tr>
<tr>
<td>F739</td>
<td>B. Arrangements of Outside Services</td>
<td></td>
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<tr>
<td></td>
<td>SNF (405.1129(b))</td>
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<tr>
<td></td>
<td>□ NOT MET</td>
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<tr>
<td>F740</td>
<td>1. The facility has a cooperative agreement with a dentist, and</td>
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<tr>
<td>F741</td>
<td>2. Maintains a list of dentists in the community for residents who do not have a private dentist.</td>
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</tr>
<tr>
<td>F742</td>
<td>3. The facility assists the resident, if necessary, in arranging for transportation to and from the dentist’s office.</td>
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</tr>
<tr>
<td></td>
<td>Social Services (Condition of Participation)</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>F743</td>
<td>SNF (405.1130)</td>
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<tr>
<td></td>
<td>□ MET</td>
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<tr>
<td></td>
<td>□ NOT MET</td>
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</tr>
</tbody>
</table>

The facility has satisfactory arrangements for identifying the medically related social and emotional needs of the resident. It is not mandatory that the skilled nursing facility itself provide social services in order to participate in the program. If the facility does not provide social services, it has written procedures for referring residents in need of social services to appropriate social agencies. If social services are offered by the facility, they are provided under a clearly defined plan, by qualified persons, to assist each resident to adjust to the social and emotional aspects of the resident’s illness, treatment, and stay in the facility.
### NAME OF FACILITY

### CODE SOCIAL SERVICES YES NO WA EXPLANATORY STATEMENT

<table>
<thead>
<tr>
<th>CODE</th>
<th>SOCIAL SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>WA</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F744</td>
<td>SNF (405.1130(a)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F745</td>
<td>Services are provided to meet the social and emotional needs of residents by qualified staff of the facility, or by referral, based on established procedures, to appropriate social agencies.</td>
<td></td>
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</tr>
<tr>
<td>F746</td>
<td>ICF (442.544(b))</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td>The facility either provides these services itself or arranges for them with qualified outside resources.</td>
</tr>
</tbody>
</table>

### B. Staffing

<table>
<thead>
<tr>
<th>CODE</th>
<th>SOCIAL SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>WA</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F747</td>
<td>SNF (405.1130(b)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
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</tr>
<tr>
<td>F748</td>
<td>1. If the facility offers social services, a member of the staff of the facility is designated as responsible for social services.</td>
<td></td>
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</tr>
<tr>
<td>F749</td>
<td>2. If the designated person is not a qualified social worker, the facility has a written agreement with a qualified social worker or recognized social agency for consultation and assistance on a regularly scheduled basis. (See §405.1101(d).)</td>
<td></td>
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</tr>
<tr>
<td>F750</td>
<td>3. The social service also has sufficient supportive personnel to meet resident needs.</td>
<td></td>
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</tr>
<tr>
<td>F751</td>
<td>4. Facilities are adequate for social service personnel, easily accessible to residents and medical and other staff, and ensure privacy for interviews.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>SOCIAL SERVICES/ACTIVITIES</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
<tr>
<td>------</td>
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</tr>
<tr>
<td>F752</td>
<td>ICF (442.344(c))</td>
<td>☐ MET ☐ NOT MET</td>
<td></td>
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</tr>
<tr>
<td>F753</td>
<td>The facility designates one staff member, qualified by training or experience, to be responsible for:</td>
<td></td>
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<td></td>
</tr>
<tr>
<td></td>
<td>a. Arranging for social services; and</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F754</td>
<td>b. Integrating social services with other elements of the plan of care.</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>C. Records and Confidentiality</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F755</td>
<td>SNF (405.1130(c)) (Standard)</td>
<td>☐ MET ☐ NOT MET</td>
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<td></td>
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</tr>
<tr>
<td>F756</td>
<td>Records of pertinent social data about personal and family problems medically related to the resident’s illness and care, and of action taken to meet the resident’s needs, are maintained in the resident’s medical records.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F757</td>
<td>If social services are provided by an outside resource, a record is maintained of each referral to such resource.</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>Activities (Condition of Participation)</td>
<td></td>
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</tr>
<tr>
<td>F758</td>
<td>SNF (405.1131)</td>
<td>☐ MET ☐ NOT MET</td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td>The facility provides for an activities program, appropriate to the needs and interests of each resident, to encourage self care, resumption of normal activities, and maintenance of an optimal level of psychosocial functioning.</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>CODE</td>
<td>ACTIVITIES/MEDICAL RECORDS</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
<tr>
<td>------</td>
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</tr>
<tr>
<td></td>
<td>A. Staffing</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F759</td>
<td>SNF (405.1131(a)) Standard</td>
<td></td>
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</tr>
<tr>
<td>F760</td>
<td>A member of the facility's staff is designated as responsible for the activities program.</td>
<td></td>
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</tr>
<tr>
<td>F761</td>
<td>If not a qualified activities coordinator, this staff member functions with frequent, regularly scheduled consultation from a person so qualified. (See §405.1101(a).)</td>
<td></td>
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<tr>
<td>F762</td>
<td>ICF (442.345(b))</td>
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</tr>
<tr>
<td></td>
<td>The facility designates one staff member, qualified by training or experience in directing group activity, to be responsible for activity service.</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>Medical Records (Condition of Participation)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F763</td>
<td>SNF (405.1132)</td>
<td></td>
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<tr>
<td></td>
<td>The facility maintains clinical (medical) records on all residents in accordance with accepted professional standards and practices. The medical record service has sufficient staff, facilities, and equipment to provide medical records that are completely and accurately documented, readily accessible, and systematically organized to facilitate retrieving and compiling information.</td>
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<tr>
<td>F764</td>
<td>ICF (442.318(a))</td>
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<tr>
<td></td>
<td>The facility maintains an organized resident record system that contains a record for each resident.</td>
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<tr>
<td>NAME OF FACILITY</td>
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</tr>
<tr>
<td>CODE</td>
<td>MEDICAL RECORDS</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
<tr>
<td>A. Staffing</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F765</td>
<td>SNF (405.1132(a)) (Standard)]</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
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</tr>
<tr>
<td>F766</td>
<td>1. Overall supervisory responsibility for the medical record service is assigned to a full-time employee of the facility.</td>
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<td></td>
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</tr>
<tr>
<td>F767</td>
<td>2. The facility also employs sufficient supportive personnel competent to carry out the functions of the medical record service.</td>
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</tr>
<tr>
<td>F768</td>
<td>3. If the medical record supervisor is not a qualified medical record practitioner, this person functions with consultation from a person qualified. (See §405.1101(b).)</td>
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</tr>
<tr>
<td>B. Protection of Medical Record Information</td>
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</tr>
<tr>
<td>F769</td>
<td>SNF (405.1132(b)) (Standard)]</td>
<td>☐ MET</td>
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<tr>
<td>F770</td>
<td>ICF (442.318(d))</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
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</tr>
<tr>
<td>F771</td>
<td>The facility safeguards medical record information against loss, destruction, or unauthorized use.</td>
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<tr>
<td>C. Physician Documentation</td>
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<tr>
<td>F772</td>
<td>SNF (405.1132(d)) (Standard)]</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
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</tr>
<tr>
<td>F773</td>
<td>1. Only physicians enter or authenticate in medical records opinions that require medical judgment (in accordance with medical staff bylaws, rules, and regulations, if applicable).</td>
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<tr>
<td>F774</td>
<td>2. All physicians sign their entries into the medical record.</td>
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</tr>
<tr>
<td>CODE</td>
<td>MEDICAL RECORDS</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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<tr>
<td>D. Completion of Records and Centralization of Reports</td>
<td></td>
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</tr>
<tr>
<td>F775</td>
<td>SNF (405.1132(w)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
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</tr>
<tr>
<td>F776</td>
<td>1. Current medical records and those of discharged residents are completed promptly.</td>
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<tr>
<td>F777</td>
<td>2. All clinical information pertaining to a resident’s stay is centralized in the resident’s medical record.</td>
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<tr>
<td>E. Retention and Preservation</td>
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<tr>
<td>F778</td>
<td>SNF (405.1132(j)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td>Medical records are retained for a period of time not less than that determined by the respective State statute, the statute of limitations in the State, or 5 years from the date of discharge in the absence of a State statute, or, in the case of a minor, 3 years after the resident becomes of age under State law.</td>
</tr>
<tr>
<td>F779</td>
<td>ICF (440.318(e))</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td>The facility must keep a resident’s record for at least 3 years after the resident is discharged.</td>
</tr>
<tr>
<td>F. Location and Facilities</td>
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<tr>
<td>F780</td>
<td>SNF (405.1132(h)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td>The facility maintains adequate facilities and equipment, conveniently located to provide efficient processing of medical records (reviewing, indexing, filing, and prompt retrieval).</td>
</tr>
<tr>
<td>CODE</td>
<td>TRANSFER AGREEMENT</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
<tr>
<td>------</td>
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<tr>
<td>F78</td>
<td>SNF (405,1133)</td>
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<tr>
<td>F782</td>
<td>ICF (442.316)</td>
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<tr>
<td></td>
<td>(Standard)</td>
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<tr>
<td>F783</td>
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<td></td>
<td>The facility has in effect a transfer agreement with one or more hospitals approved for participation under the programs, which provides the basis for effective working arrangements under which inpatient hospital care or other hospital services are available promptly to the facility's residents when needed. (A facility that has been unable to establish a transfer agreement with the hospital(s) in the community or service area after documented attempts to do so is considered to have such an agreement in effect.) Exception: A facility that has been unable to establish a written agreement after documented attempts to do so, is considered to have such an agreement.</td>
</tr>
<tr>
<td>F784</td>
<td>SNF (405,1133(a))</td>
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<tr>
<td></td>
<td>(Standard)</td>
<td></td>
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<tr>
<td>F785</td>
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<td></td>
<td>A hospital and a skilled nursing facility shall be considered to have a transfer agreement in effect if, by reason of a written agreement between them or (in case of two institutions are under common control) by reason of a written undertaking by the person or body which controls them, there is reasonable assurance that:</td>
</tr>
<tr>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>1. Transfer of patients will be effected between the hospital and the skilled nursing facility, ensuring timely admission, whenever such transfer is medically appropriate as determined by the attending physician.</td>
</tr>
<tr>
<td>CODE</td>
<td>TRANSFER AGREEMENT/PHYSICAL ENVIRONMENT</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
<tr>
<td>------</td>
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</tr>
<tr>
<td>F786</td>
<td>2. There will be interchange of medical and other information necessary or useful in the care and treatment of individuals transferred between institutions, or in determining whether such individuals can be adequately cared for otherwise than in either of such institutions.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F787</td>
<td>3. Security and accountability for residents' personal effects are provided on transfer.</td>
<td></td>
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</tr>
<tr>
<td>F788</td>
<td><strong>Physical Environment (Condition of Participation)</strong></td>
<td></td>
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<td></td>
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</tr>
<tr>
<td></td>
<td>SNF (405.1134) (Standard)</td>
<td>MET</td>
<td>NOT MET</td>
<td></td>
<td>The facility is constructed, equipped, and maintained to protect the health and safety of residents, personnel, and the public.</td>
</tr>
<tr>
<td></td>
<td>ICF (442.321) (Standard)</td>
<td>MET</td>
<td>NOT MET</td>
<td></td>
<td>(See appropriate HCFA Fire Safety survey form.)</td>
</tr>
<tr>
<td></td>
<td><strong>A. Life Safety from Fire</strong></td>
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<tr>
<td></td>
<td>SNF (405.1134(a)) (Standard)</td>
<td>MET</td>
<td>NOT MET</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>ICF (442.321) (Standard)</td>
<td>MET</td>
<td>NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>B. Maintenance of Equipment, Building, and Grounds</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td>SNF (405.1134(i)) (Standard)</td>
<td>MET</td>
<td>NOT MET</td>
<td></td>
<td>The facility establishes a written preventative maintenance program to ensure that all equipment is operative.</td>
</tr>
</tbody>
</table>
### Infection Control (Condition of Participation)

<table>
<thead>
<tr>
<th>CODE</th>
<th>INFECTION CONTROL</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F791</td>
<td>SNF (405.1135)</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>The facility establishes an infection control committee of representative professional staff with responsibility for overall infection control in the facility. All necessary housekeeping and maintenance services are provided to maintain a sanitary and comfortable environment and to help prevent the development and transmission of infection.</td>
</tr>
</tbody>
</table>

#### A. Infection Control Committee

<table>
<thead>
<tr>
<th>CODE</th>
<th>INFECTION CONTROL</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F792</td>
<td>SNF (405.1135(a)) (Standard)</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
<tr>
<td>F793</td>
<td>1. The infection control committee is composed of members of the medical and nursing staffs, administration, and the dietary, pharmacy, housekeeping, maintenance, and other services.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
<tr>
<td>F794</td>
<td>2. The committee establishes policies and procedures for investigating, controlling, and preventing infection in the facility.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
<tr>
<td>F795</td>
<td>3. The committee monitors staff performance to ensure that the policies and procedures are executed.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
</tbody>
</table>

#### B. Aseptic and Isolation Techniques

<table>
<thead>
<tr>
<th>CODE</th>
<th>INFECTION CONTROL</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F796</td>
<td>SNF (405.1135(b)) (Standard)</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
<tr>
<td>F797</td>
<td>1. The facility has written procedures for aseptic and isolation techniques.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
<tr>
<td>F798</td>
<td>2. These procedures are reviewed and revised for effectiveness and improvement as necessary.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>INFECTION CONTROL</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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</tr>
<tr>
<td>F799</td>
<td><strong>C. Housekeeping</strong></td>
<td>MET</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F800</td>
<td>SNF (405.1135(c)) (Standard)</td>
<td>MET</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1. The facility employs sufficient housekeeping personnel.</td>
<td></td>
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</tr>
<tr>
<td>F801</td>
<td>2. Provides all necessary equipment to maintain a safe, clean and orderly interior.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F802</td>
<td>3. A full-time employee is designated responsible for the services and for supervision and training of personnel.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F803</td>
<td>4. If a facility has a contract with an outside resource for housekeeping services, the facility and/or outside resource meets the requirements of the standards.</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>F804</td>
<td><strong>D. Pest Control</strong></td>
<td>MET</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>SNF (405.1135(p)) (Standard)</td>
<td>MET</td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td>The facility has an ongoing pest control program.</td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>
### NAME OF FACILITY

#### DISASTER PREPAREDNESS

<table>
<thead>
<tr>
<th>CODE</th>
<th>DISASTER PREPAREDNESS</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F805</td>
<td>SNF (405.1136)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Disaster Preparedness (Condition of Participation)</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>The facility has a written plan, periodically rehearsed, with procedures to be followed in the event of an internal or external disaster and for the care of casualties (residents and personnel) arising from such disasters.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### A. Plan

<table>
<thead>
<tr>
<th>CODE</th>
<th>DISASTER PREPAREDNESS</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F806</td>
<td>ICF (442.313) (Standard)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1. The facility has a written plan for staff and residents to follow in case of emergencies such as fire or explosion.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F807</td>
<td>2. The facility rehearses the plan regularly.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F808</td>
<td>3. The facility has written procedures for the staff to follow in case of an emergency involving an individual resident.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F809</td>
<td>4. These procedures include: a. Caring for the resident.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F810</td>
<td>b. Notifying the attending physician and other individuals responsible for the resident.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F811</td>
<td>c. Arranging for transportation, hospitalization, and other appropriate services.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F812</td>
<td>SNF (405.1136(a)) (Standard)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F813</td>
<td>1. The facility has an acceptable written plan in operation, with procedures to be followed in the event of fire, explosion, or other disaster.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F814</td>
<td>2. The plan is developed and maintained with the assistance of qualified fire, safety, and other appropriate experts.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F815</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>DISASTER PREPAREDNESS/UTILIZATION REVIEW</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
<tr>
<td>------</td>
<td>----------------------------------------</td>
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</tr>
<tr>
<td>FB16</td>
<td>3. Includes procedures for prompt transfer of casualties and records.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>FB17</td>
<td>4. Instructions regarding the location and use of alarm systems and signals and of fire-fighting equipment.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>FB18</td>
<td>5. Information regarding methods of containing fire.</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>FB20</td>
<td>7. Specifications of evacuation routes and procedures. (See §405.1134(a).)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

B. Orientation and training

FB21 SNF (405.1136(b)) (standard) ☐ MET ☐ NOT MET

The disaster program includes orientation and ongoing training and drills for all personnel in all procedures so that each employee promptly and correctly carries out a specific role in case of a disaster (See §405.1121(b)).

Utilization Review (Condition of Participation)

FB23 SNF (405.1137) ☐ MET ☐ NOT MET

The facility carries out utilization review of the services provided in the facility to residents who are entitled to benefits under the program(s). Utilization review assures the maintenance of high quality care and appropriate and efficient utilization of facility services. There are two elements to utilization review: medical care evaluation studies and review of extended duration cases.
<table>
<thead>
<tr>
<th>CODE</th>
<th>UTILIZATION REVIEW</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A. Plan</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F824</td>
<td>SNF (405.1137(a)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F825</td>
<td>1. The facility has a currently applicable written description of its utilization review plan.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F826</td>
<td>2. Such description includes:</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td>a. The organization and composition of the committee or group which will be responsible for the utilization review function.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F827</td>
<td>b. Methods of criteria (including norms where available) to be used to define periods of continuous extended duration and to assign or select subsequent dates for continued stay review.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F828</td>
<td>c. Methods for selection and conduct of medical care evaluation studies.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>B. Organization and Composition of Utilization Review Committees</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F829</td>
<td>SNF (405.1137(b)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F830</td>
<td>1. The utilization review (UR) function is conducted by:</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>a. A staff committee of the skilled nursing facility which is composed of two or more physicians, with participation of other professional personnel; or,</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>CODE</td>
<td>UTILIZATION REVIEW</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
<tr>
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<tr>
<td>F831</td>
<td>b. A group outside the facility which is similarly composed and which is established by the local medical or osteopathic society and some or all of the hospitals and skilled nursing facilities in the locality; or (indicate name of the outside group and briefly describe the organization.)</td>
<td></td>
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<tr>
<td>F832</td>
<td>c. A group established and organized in a manner approved by the Secretary that is capable of performing such function.</td>
<td></td>
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</tr>
<tr>
<td>F833</td>
<td>2. The medical care evaluation studies, educational duties of the review program, and the review of admissions and long-stay cases are performed by:</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>a. the same committee or group:</td>
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<td></td>
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<tr>
<td>F834</td>
<td>b. or more committees or groups.</td>
<td></td>
<td></td>
<td></td>
<td>Briefly explain who performs these functions.</td>
</tr>
<tr>
<td></td>
<td>C. Medical Care Evaluation Studies</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F856</td>
<td>1. Medical care evaluation studies are performed to promote the most effective and efficient use of available health facilities and services consistent with resident needs and professionally recognized standards of health care.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F857</td>
<td>2. Studies emphasize identification and analysis of patterns of resident care and suggest, where appropriate, possible changes for maintaining consistently high quality care and effective and efficient use of services.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>UTILIZATION REVIEW</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
<tr>
<td>------</td>
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</tr>
<tr>
<td>F838</td>
<td>3. Each medical care evaluation study identifies and analyzes factors related to the care rendered in the facility and where indicated, results in recommendations for change beneficial to residents, staff, the facility, and the community.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F839</td>
<td>4. Studies, on a sample or other basis, include, but need not be limited to, admissions, durations of stay, ancillary services furnished (including drugs and biologicals), and professional services performed on premises.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F840</td>
<td>At least one study was completed during the last year. Type of study last completed:</td>
<td></td>
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</tr>
<tr>
<td>F841</td>
<td>D. Extended Stay Review</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td></td>
<td>SNF (405.1137(d)) (Standard)</td>
<td>□ MET</td>
<td>□ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F842</td>
<td>1. Periodic review is made of each current inpatient skilled nursing facility beneficiary case of continuous extended duration, and the length of which is defined in the utilization review plan to determine whether further inpatient stay is necessary.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F843</td>
<td>2. The review is based on the attending physician's reasons for and plan for continued stay and any other documentation the committee or group deems appropriate.</td>
<td></td>
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<tr>
<td>F844</td>
<td>3. Cases are screened by:</td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td>a. A qualified non-physician representative of the committee.</td>
<td></td>
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<tr>
<td>F845</td>
<td>b. The group.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F846</td>
<td>c. The reviewer uses criteria established by the physician members of the committee.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>EXPLANATORY STATEMENT</td>
<td></td>
<td></td>
<td></td>
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<td>------</td>
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<tr>
<td>F847</td>
<td>In instances when non-physician members are utilized, cases are referred to a physician member for further review when it appears that the resident no longer requires further inpatient care.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F848</td>
<td>Non-physician representatives used to screen extended stay review cases have experience in such screening or appropriate training in the application of the screening criteria used, or both.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F849</td>
<td>Before the expiration of each new period, the case must be reviewed again in like manner with such reviews being repeated as long as the stay continues beyond the scheduled review dates and notice has not been given pursuant to paragraph (e) of this section.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F850</td>
<td>E. Further Stay Not Medically Necessary</td>
<td></td>
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</tr>
<tr>
<td>F851</td>
<td>A final determination of the committee or group that the continued stay is not medically necessary is made by at least two physician members of the committee or group, except that the final determination may be made by one physician where the attending physician, when given an opportunity to express his views, does not do so, or does not contest the finding that the continued stay is not medically necessary.</td>
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<td></td>
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</tr>
<tr>
<td>F852</td>
<td>2. If the committee or group, or its nonphysician representative where a physician member concurs, has reason to believe from the review of an extended duration case or a case reviewed as part of a medical care evaluation study that further stay is no longer medically necessary, the committee or group shall notify the individual’s attending physician and afford him an opportunity to present his views before it makes a final determination.</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>
### NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>UTILIZATION REVIEW</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F853</td>
<td>3. If the final determination of the committee or group is that further stay is no longer medically necessary, written notification of the finding is given to the facility, the attending physician, and the individual (or where appropriate, his next of kin) no later than 2 days after such final determination is made and, in no event in the case of an extended duration case, later than 3 working days after the end of the extended duration period specified pursuant to paragraph (d) of this section.</td>
<td></td>
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</tr>
</tbody>
</table>

### F. Administrative Responsibilities

<table>
<thead>
<tr>
<th>F854</th>
<th>SNF (405.1137(j)) (Standard)</th>
<th>MET</th>
<th>NOT MET</th>
</tr>
</thead>
<tbody>
<tr>
<td>F855</td>
<td>The administrative staff of the facility is kept directly and fully informed of committee activities to facilitate support and assistance. (Explain)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### G. Utilization Review Records

<table>
<thead>
<tr>
<th>F856</th>
<th>SNF (405.1137(g)) (Standard)</th>
<th>MET</th>
<th>NOT MET</th>
</tr>
</thead>
<tbody>
<tr>
<td>F857</td>
<td>1. Written records of committee activities are maintained.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F858</td>
<td>2. Appropriate reports, signed by the committee chairman, are made regularly to the medical staff, administrative staff, governing body, and sponsors (if any).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F859</td>
<td>3. Minutes of each committee meeting is maintained and include at least: a. Name of committee. b. Date and duration of meeting. c. Names of committee members present and absent.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>UTILIZATION REVIEW</td>
<td>YES</td>
<td>NO</td>
</tr>
<tr>
<td>------</td>
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<td>-----</td>
<td>----</td>
</tr>
<tr>
<td>F862</td>
<td>4. Description of activities presently in progress to satisfy the requirements for medical care evaluation studies, including the subject, reason for study, dates of commencement and expected completion, summary of studies completed since the last meeting, conclusions and follow-up on implementation of recommendations made from previous studies.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F863</td>
<td>5. Summary of extended duration cases reviewed including the number of cases, identification number, admission and review dates, and decision reached, including the basis for each determination and action taken for each case not approved for extended care.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F854</td>
<td>H. Discharge Planning</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F864</td>
<td>SNF (405.1137(h)) (Standard) ☐ MET ☐ NOT MET</td>
<td></td>
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<tr>
<td></td>
<td>The facility maintains a centralized, coordinated program to ensure that each resident has a planned program of continuing care which meets his postdischarge needs.</td>
<td></td>
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</tr>
<tr>
<td>F865</td>
<td>1. The facility has in operation an organized discharge planning program.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F866</td>
<td>The utilization review committee, in its evaluation of the current status of each extended duration case, has available to it the results of such discharge planning and information on alternative available community resources to which the resident may be referred.</td>
<td></td>
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</tr>
<tr>
<td>F867</td>
<td>2. The facility maintains written discharge planning procedures which describe:</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>a. How the discharge coordinator will function, and his authority and relationships with the facility's staff.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F868</td>
<td>b. The maximum time period after which reevaluation of each resident's discharge plan is made.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
§ 488.105 Long term care survey forms, Part B.

DEPARTMENT OF HEALTH AND HUMAN SERVICES
HEALTH CARE FINANCING ADMINISTRATION

PART B

MEDICARE / MEDICAID SKILLED NURSING FACILITY AND INTERMEDIATE CARE FACILITY SURVEY REPORT

<table>
<thead>
<tr>
<th>PROVIDER NUMBER</th>
<th>FACILITY NAME AND ADDRESS (City, State, Zip)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
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</table>

<table>
<thead>
<tr>
<th>VENDOR NUMBER</th>
</tr>
</thead>
<tbody>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>SURVEY DATE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>SURVEYORS' NAMES</th>
<th>TITLES</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>SURVEY TEAM COMPOSITION</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1 Indicate the Number of Surveyors According to Discipline:</td>
</tr>
<tr>
<td>A. Administrator</td>
</tr>
<tr>
<td>B. Nurse</td>
</tr>
<tr>
<td>C. Dietitian</td>
</tr>
<tr>
<td>D. Pharmacist</td>
</tr>
<tr>
<td>E. Records Administrator</td>
</tr>
<tr>
<td>F. Social Worker</td>
</tr>
<tr>
<td>G. Qualified Mental Health Professional</td>
</tr>
</tbody>
</table>

Note: More than one discipline may be marked for surveyors qualified in multiple disciplines.

F2 Indicate the Total Number of Surveyors Onsite: ______________

Form HCFA-619 (2-90) (CONTINUED ON REVERSE)
<table>
<thead>
<tr>
<th>PROVIDER NO.</th>
<th>F3 MEDICARE</th>
<th>F4 MEDICAID</th>
<th>F5 OTHER</th>
<th>F6 TOTAL RESIDENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>BATHING</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F7</td>
<td></td>
<td></td>
<td></td>
<td>Number of residents requiring assistance in bathing more than one part of body—or does not bathe self.</td>
</tr>
<tr>
<td>F8</td>
<td></td>
<td></td>
<td></td>
<td>Number of residents requiring assistance in bathing only a single part (as back or disabled extremity) or bathes self completely.</td>
</tr>
<tr>
<td>F9</td>
<td>TOTAL*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>DRESSING</strong></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F10</td>
<td></td>
<td></td>
<td></td>
<td>Number of residents totally dressed by another person.</td>
</tr>
<tr>
<td>F11</td>
<td></td>
<td></td>
<td></td>
<td>Number of residents needing assistance to dress self or remain partly dressed. (Exclude those residents totally dressed.)</td>
</tr>
<tr>
<td>F12</td>
<td></td>
<td></td>
<td></td>
<td>Number of residents able to get clothes from closets and drawers, puts on clothes, outer garments, brassieres, fasteners. Act of tying shoes is excluded.</td>
</tr>
<tr>
<td>F13</td>
<td>TOTAL*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>TOILETING</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F14</td>
<td></td>
<td></td>
<td></td>
<td>Number of residents not toileted. (Use protective padding, catheter.)</td>
</tr>
<tr>
<td>F15</td>
<td></td>
<td></td>
<td></td>
<td>Number of residents who must use a bedpan or commode and/or receive assistance in getting to and using a toilet.</td>
</tr>
<tr>
<td>F16</td>
<td></td>
<td></td>
<td></td>
<td>Number of residents able to get to toilet—gets on and off toilet—cleans self—arranges clothes.</td>
</tr>
<tr>
<td>F17</td>
<td>TOTAL*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>TRANSFERRING</strong></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F18</td>
<td></td>
<td></td>
<td></td>
<td>Number of residents needing assistance in all transfers (moving in or out of bed and/or chair, toilet, tub, transfers).</td>
</tr>
<tr>
<td>F19</td>
<td></td>
<td></td>
<td></td>
<td>Number of residents needing assistance in transferring to toilet and tub only.</td>
</tr>
<tr>
<td>F20</td>
<td></td>
<td></td>
<td></td>
<td>Number of residents able to complete all transfers independently (may or may not be using mechanical support).</td>
</tr>
<tr>
<td>F21</td>
<td>TOTAL*</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**CONTINENT**

- F22: Number of residents with indwelling or external catheters.
- F23: Number of residents with partial or total incontinence in urination or defecation—partial or total control by suppositories or enemas, regulated use of urinals and/or bedpans.
- F24: Number of residents with incontinence in urination and defecation entirely self-controlled.
- F25: TOTAL*

**FEEDING**

- F26: Number of residents who receive enteral/parenteral feedings.
- F27: Number of residents who receive NG tube feedings.
- F28: Number of residents who require assistance in act of eating.
- F29: Number of residents who get food from plate or its equivalent into mouth—pre-cutting of meat and preparation of food, buttering bread, opening cans, removing plate covers, etc. are excluded from evaluation.
- F30: TOTAL*
### Name of Facility

<table>
<thead>
<tr>
<th>Code</th>
<th>Governing Body</th>
<th>Yes</th>
<th>No</th>
<th>N/A</th>
<th>Explanatory Statement</th>
</tr>
</thead>
<tbody>
<tr>
<td>F50</td>
<td>SNF (405.1121)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Resident Rights

<table>
<thead>
<tr>
<th>Code</th>
<th>Governing Body (Condition of Participation)</th>
<th>Yes</th>
<th>No</th>
<th>N/A</th>
<th>Explanatory Statement</th>
</tr>
</thead>
<tbody>
<tr>
<td>F51</td>
<td>SNF (405.1121)(K) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td>Indicators A thru K apply to this standard for SNF's.</td>
</tr>
<tr>
<td>F52</td>
<td>ICF (442.311) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td>Indicators A thru K apply to this standard for ICF's.</td>
</tr>
</tbody>
</table>

### A. Information

1. The facility informs each resident, before or at the time of admission, of his/her rights and responsibilities.

2. The facility informs each resident, before or at the time of admission, of all rules governing resident conduct.

3. The facility informs each resident of amendments to their policies on residents' rights and responsibilities and rules governing conduct.

4. Each resident acknowledges in writing receipt of residents' rights information and any amendment to it.

5. The resident must be informed in writing of all services and charges for services.

6. The resident must be informed in writing of all changes in services and charges before or at the time of admission and on a continuing basis.

7. The resident must be informed of services not covered by Medicare or Medicaid and not covered in the basic rate.
## NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>GOVERNING BODY</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>B. Medical Condition and Treatment</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F60</td>
<td>1. Each resident is informed by a physician of his/her health and medical condition unless the physician decides that informing the resident is medically contraindicated.</td>
<td></td>
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</tr>
<tr>
<td>F61</td>
<td>2. Each resident is given an opportunity to participate in planning his/her total care and medical treatment.</td>
<td></td>
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</tr>
<tr>
<td>F62</td>
<td>3. Each resident is given an opportunity to refuse treatment.</td>
<td></td>
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</tr>
<tr>
<td>F63</td>
<td>4. Each resident gives informed, written consent before participating in experimental research.</td>
<td></td>
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</tr>
<tr>
<td>F64</td>
<td>5. If the physician decides that informing the resident of his/her health and medical condition is medically contraindicated, the physician has documented this decision in the resident’s medical record.</td>
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<tr>
<td>C. Transfer and Discharge</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F65</td>
<td>Each resident is transferred or discharged only for:</td>
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</tr>
<tr>
<td>F66</td>
<td>1. Medical reasons.</td>
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</tr>
<tr>
<td>F67</td>
<td>2. Higher welfare or that of other residents.</td>
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</tr>
<tr>
<td>F68</td>
<td>3. Resident’s care is transfered to the Medicare or Medicaid programs.</td>
<td></td>
<td></td>
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<tr>
<td>D. Exercising Rights</td>
<td></td>
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</tr>
<tr>
<td>F69</td>
<td>1. Each resident is encouraged and assisted to exercise his/her rights as a resident of the facility and as a citizen.</td>
<td></td>
<td></td>
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<tr>
<td>F70</td>
<td>2. Each resident is allowed to submit complaints and recommendations concerning the policies and services of the facility to staff or to outside representatives of the resident’s choice or both.</td>
<td></td>
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</tbody>
</table>
### NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>GOVERNING BODY</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F71</td>
<td>3. Such complaints are submitted free from restraint, coercion, discrimination, or reprisal.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td><strong>E. Financial Affairs</strong></td>
<td></td>
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</tr>
<tr>
<td>F72</td>
<td>1. Residents are allowed to manage their own personal financial affairs.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F73</td>
<td>2. The facility establishes and maintains a system that assures full and complete accounting of residents' personal funds. An accounting report is made to each resident in a skilled nursing facility at least on a quarterly basis.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F74</td>
<td>3. The facility does not commingle resident funds with any other funds.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F75</td>
<td>4. If a resident requests assistance from the facility in managing his/her personal financial affairs, resident's delegation is in writing.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F76</td>
<td>5. The facility system of accounting includes written receipts for: All personal possessions and funds received by or deposited with the facility.</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>F77</td>
<td>All disbursements made to or for the resident.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F78</td>
<td>6. The financial record must be available to the resident and his/her family.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td><strong>F. Freedom from Abuse and Restraints</strong></td>
<td></td>
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</tr>
<tr>
<td>F79</td>
<td>1. Each resident is free from mental and physical abuse.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F80</td>
<td>2. Chemical and physical restraints are only used when authorized by a physician in writing for a specified period of time or in emergencies.</td>
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<tr>
<td>CODE</td>
<td>GOVERNING BODY</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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<tr>
<td>F81</td>
<td>3. If used in emergencies, they are necessary to protect the resident from injury to himself/herself or others.</td>
<td></td>
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</tr>
<tr>
<td>F82</td>
<td>4. The emergency use is authorized by a professional staff member identified in the written policies and procedures of the facility.</td>
<td></td>
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<tr>
<td>F83</td>
<td>5. The emergency use is reported promptly to the resident's physician by the staff member.</td>
<td></td>
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<tr>
<td>G</td>
<td>Privacy</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F84</td>
<td>1. Each resident is treated with respect, consideration and full recognition of his/her dignity and individuality.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F85</td>
<td>2. Each resident is given privacy during treatment and care of personal needs.</td>
<td></td>
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</tr>
<tr>
<td>F86</td>
<td>3. Each resident's records, including information in an automated data bank, are treated confidentially.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F87</td>
<td>4. Each resident must give written consent before the facility releases information from his/her record to someone not otherwise authorized to receive it.</td>
<td></td>
<td></td>
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<tr>
<td>F88</td>
<td>5. Married residents are given privacy during visits by their spouses.</td>
<td></td>
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</tr>
<tr>
<td>F89</td>
<td>6. Married residents are permitted to share a room.</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>H</td>
<td>Work</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F90</td>
<td>No resident may be required to perform services for the facility.</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>CODE</td>
<td>GOVERNING BODY</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
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<tr>
<td>F91</td>
<td>Freedom of Association and Correspondence</td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td>1. Each resident is allowed to communicate, associate and meet privately with individuals of his/her choice unless this infringes upon the rights of another resident.</td>
<td></td>
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</tr>
<tr>
<td>F92</td>
<td>2. Each resident is allowed to send and receive personal mail unopened.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F93</td>
<td>J. Activities</td>
<td></td>
<td></td>
<td></td>
<td>Each resident is allowed to participate in social, religious, and community group activities.</td>
</tr>
<tr>
<td>F94</td>
<td>K. Personal Possessions</td>
<td></td>
<td></td>
<td></td>
<td>Each resident is allowed to retain and use his/her personal possessions and clothing as space permits.</td>
</tr>
<tr>
<td>F95</td>
<td>L. Delegation of Rights and Responsibilities</td>
<td></td>
<td></td>
<td></td>
<td>ICF (442.312) (Standard)</td>
</tr>
<tr>
<td></td>
<td>1. All the rights and responsibilities of a resident pass to the resident's guardian, next of kin or sponsoring agency or agencies if the resident is adjudicated incompetent under State law or is determined by his/her physician to be incapable of understanding his/her rights and responsibilities.</td>
<td></td>
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</tr>
<tr>
<td>F97</td>
<td>2. Physician determinations of incapability and the specific reasons thereof are recorded by the physician in the resident's record.</td>
<td></td>
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</tr>
</tbody>
</table>
### NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>GOVERNING,body</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
</table>

#### F98
- **SNF (405.112)(b)** (Standard):  
  - MET: 
  - NOT MET: 

#### F99
- **ICF (442.314)** (Standard):  
  - MET: 
  - NOT MET: 

#### F100
1. Facility staff are knowledgeable about the problems and needs of the aged, ill, and disabled.

#### F101
2. Facility staff practices proper techniques in providing care to the aged, ill, and disabled.

#### F102
3. Facility staff practice proper technique for prevention and control of infection, fire prevention and safety, accident prevention, confidentiality of resident information, and preservation of resident dignity, including protection of privacy and personal and property rights.

#### STATUS CHANGE NOTIFICATIONS

#### F103
- **SNF (405.112)(b)** (Standard):  
  - MET: 
  - NOT MET: 

#### F104
- **ICF (442.307)** (Standard):  
  - Met: 
  - Not Met: 

#### F105
1. The facility notifies the resident's attending physician and other responsible persons in the event of an accident involving the resident, or other significant change in the resident's physical, mental, or emotional status, or resident charges, billings, and related administrative matters.

#### F106
2. Except in a medical emergency, a resident is not transferred or discharged, nor is treatment altered radically, without consultation with the resident or, if the resident is incompetent, without prior notification of next of kin or sponsor.
<table>
<thead>
<tr>
<th>CODE</th>
<th>PHYSICIANS' SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>P107</td>
<td>SNF (405.1123)</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td></td>
<td>MET</td>
<td>NOT</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>A. Medical Findings and Orders at Time of Admission</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>P108</td>
<td>SNF (405.1123a) (Standard)</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>MET</td>
<td>NOT</td>
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</tr>
<tr>
<td></td>
<td>1. There is made available to the facility prior to or at the time of admission, resident information which includes current medical findings, diagnoses, and orders from a physician for immediate care of the resident.</td>
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<tr>
<td>P109</td>
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<tr>
<td></td>
<td>2. Information about the rehabilitation potential of the resident and a summary of prior treatment are made available to the facility at the time of admission or within 48 hours thereafter.</td>
<td></td>
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</tbody>
</table>
### NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>PHYSICIAN SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F11</td>
<td>Resident Supervision by Physician</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F12</td>
<td>SNF (405.1120) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F12</td>
<td>ICF (442.346) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td>Applies to ICFs</td>
</tr>
<tr>
<td></td>
<td>Indicators B and C apply to</td>
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<tr>
<td></td>
<td>each standard for ICFs.</td>
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</tr>
<tr>
<td>F13</td>
<td>1. Every resident must be under the supervision of a</td>
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<tr>
<td></td>
<td>physician.</td>
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<tr>
<td>F14</td>
<td>2. A physician prescribes a planned regimen of care based</td>
<td></td>
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<tr>
<td></td>
<td>on a medical evaluation of each resident’s immediate and</td>
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<td></td>
<td>long-term care needs.</td>
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<tr>
<td></td>
<td>Exception: Not required for ICF residents</td>
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<tr>
<td>F15</td>
<td>3. A physician is available to provide care in the absence of</td>
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<tr>
<td></td>
<td>any resident’s attending physician.</td>
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<tr>
<td>F16</td>
<td>4. Medical evaluation is done within 48 hours of admission</td>
<td></td>
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<tr>
<td></td>
<td>unless done within 5 days prior to admission.</td>
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<tr>
<td></td>
<td>Exception: Not required for ICF residents.</td>
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</tr>
<tr>
<td>F17</td>
<td>5. Each resident is seen by their attending physician at least</td>
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<td>once every 30 days for the first 60 days after admission.</td>
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<td></td>
<td>Exception: ICF residents must be seen every 60 days unless</td>
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<td>otherwise justified and documented by the attending physician.</td>
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<td>F18</td>
<td>6. Each resident’s total program of care including medications and treatments is reviewed during a visit by the attending physician at least once every 30 days for the first 90 days and revised as necessary.</td>
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<td></td>
<td>Exception: Only medications must be reviewed quarterly for ICF residents.</td>
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<td>F119</td>
<td>7. Progress notes are written and signed by the physician at the time of each visit, and all orders are signed by the physician.</td>
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<tr>
<td>F120</td>
<td>8. Alternate physician visit schedules that exceed a 30-day schedule adopted after the 90th day following admission are justified by the attending physician in the medical record. These visits cannot exceed 60 days or apply to residents who require specialized rehabilitation schedules.</td>
<td></td>
<td></td>
<td></td>
<td>EXCEPTION: Not required for ICF residents.</td>
</tr>
<tr>
<td>F121</td>
<td>C. Emergency Services</td>
<td></td>
<td></td>
<td></td>
<td>SNF (405.1123(c)) (Standard)</td>
</tr>
<tr>
<td>F122</td>
<td>Emergency services from a physician are available and provided to each resident who requires emergency care.</td>
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<tr>
<td>F123</td>
<td>NURSING SERVICES (CONDITION OF PARTICIPATION)</td>
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<td></td>
<td></td>
<td>SNF (405.1124)</td>
</tr>
<tr>
<td>F124</td>
<td>SRF (405.1124(c)) (Standard)</td>
<td>MET</td>
<td>NOT MET</td>
<td>Indicators A and B apply to this standard for SNFs</td>
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</tr>
<tr>
<td>F125</td>
<td>ICF (440.238)</td>
<td>MET</td>
<td>NOT MET</td>
<td>Indicators A thru E apply to this standard for ICFs except where noted.</td>
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</tr>
<tr>
<td></td>
<td>A. The facility provides nursing services which are sufficient to meet nursing needs of all residents all hours of each day.</td>
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<tr>
<td>F126</td>
<td>1. Each resident receives all treatments, medications and diets as prescribed. Deviations are reported and appropriate action is taken.</td>
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<td>EXPLANATORY STATEMENT</td>
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<tr>
<td>F127</td>
<td>2. Each resident receives daily personal hygiene as needed to assure cleanliness, good skin care, good grooming, and oral hygiene taking into account individual preferences. Residents are encouraged to engage in self care activity.</td>
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<tr>
<td>F128</td>
<td>3. Each resident receives care necessary to prevent skin breakdown.</td>
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<td>F129</td>
<td>4. Each resident with a decubitus receives care necessary to promote the healing of the decubitus including proper dressing.</td>
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<td>F130</td>
<td>5. When residents require restraints the application is ordered by the physician, applied properly, and released at least every 2 hours.</td>
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<tr>
<td>F131</td>
<td>6. Each resident with incontinence is provided with care necessary to encourage continence including frequent toileting and opportunities for rehabilitative training.</td>
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<td>F132</td>
<td>7. Each resident with a urinary catheter receives proper routine care including periodic evaluation.</td>
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<td>F133</td>
<td>8. Each resident receives proper care for the following needs:</td>
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<td></td>
<td>Injections</td>
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<td></td>
<td>Parenteral Fluids</td>
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<td></td>
<td>Colostomy/Stoma</td>
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<td>Tracheostomy Care</td>
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<td>Suctioning</td>
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<td></td>
<td>Tube Feeding</td>
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<tr>
<td>F134</td>
<td>9. Infection Control Techniques are properly carried out in the provision of care to each resident.</td>
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<tr>
<td>CODE</td>
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<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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<td>F135</td>
<td>10. Proper nursing and sanitary procedures and techniques are used when medications are given to residents.</td>
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<td>F136</td>
<td>11. Adequate resident care supplies are available for providing treatments.</td>
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<tr>
<td>F137</td>
<td><strong>B. Twenty-Four Hour Nursing Service</strong></td>
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<tr>
<td></td>
<td>1. Nursing personnel, including registered nurses, licensed practical (vocational) nurses, nurse aides, orderlies, and ward clerks, are assigned duties consistent with their education and experience, and based on the characteristics of the resident load.</td>
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<td><strong>EXCEPTION:</strong> Not required for ICFs.</td>
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<td>F138</td>
<td>2. Weekly time schedules are maintained and indicate the number and classifications of nursing personnel including relief personnel, who worked on each unit for each tour of duty.</td>
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<td>(If a distinct part certification, show the staffing for the IP and, if appropriate, any nonparticipating remainder and explain any sharing of nursing personnel.)</td>
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<td><strong>Exception:</strong> Not required for Freestanding ICFs.</td>
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<tr>
<td>F139</td>
<td>3. There is a sufficient number of nursing staff available to meet the total needs of all residents.</td>
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<tr>
<td>F140</td>
<td>4. There is a registered nurse on the day tour of duty 7 days a week.</td>
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<td><strong>Exception:</strong> Not required for ICF residents.</td>
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<tr>
<td>CODE</td>
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<td>EXPLANATORY STATEMENT</td>
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<tr>
<td>F1A1</td>
<td>C. Charge Nurse</td>
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<td>NOT MET</td>
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<tr>
<td>F142</td>
<td>1. A registered nurse or a qualified licensed practical (or vocational) nurse is designated as charge nurse by the director of nursing for each tour of duty.</td>
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<td>Exception: Not required for ICFs.</td>
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<tr>
<td>F143</td>
<td>2. The director of nursing services does not serve as charge nurse in a facility with an average daily total occupancy of 60 or more residents.</td>
<td></td>
<td>Exception: Not required for ICFs.</td>
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<tr>
<td>F144</td>
<td>3. The ICF must have a registered nurse, or a licensed practical or vocational nurse full-time, 7 days a week, on the day shift.</td>
<td></td>
<td>Exception: Not required for SNFs.</td>
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</table>
List the number of full-time equivalents of RN's, LPN's, Aides/Orderlies assigned to nursing duty from the last 3 complete weeks. (Note only actual staff on duty.)

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## Staffing Pattern Worksheets Day of Survey (Optional)

### Entire Facility Staffing Pattern (Day of Survey)

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### Unit Staffing Pattern Worksheet (Day of Survey)

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**NAME OF FACILITY**

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<td>N/A</td>
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<td>F167</td>
<td>SNF (405.1124d)</td>
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<td></td>
<td>Each resident's needs are addressed in a written plan of care which demonstrates that the plans of all services are integrated, consonant with the physician's plan of medical care, and implemented shortly after admission.</td>
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<tr>
<td>F168</td>
<td>ICF (442.341)</td>
<td></td>
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<td></td>
<td>Each professional service identifies needs, goals, plans, and evaluates the effectiveness of interventions, plus institutes changes in the plan of care in a timely manner.</td>
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<tr>
<td>F169</td>
<td>SNF (405.1124w)</td>
<td></td>
<td></td>
<td></td>
<td>Rehabilitative Nursing Services are performed daily, and recorded for those residents who require such service.</td>
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<tr>
<td>F170</td>
<td>ICF (442.342)</td>
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<td></td>
<td></td>
<td>Each resident receives rehabilitative nursing care to promote maximum physical functioning to prevent immobility, deformities, and contractures.</td>
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<tr>
<td>F171</td>
<td>SNF (405.1124w)</td>
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<td></td>
<td>There is an ongoing evaluation of each resident's rehabilitative nursing needs. This may include:</td>
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<tr>
<td>F172</td>
<td>ICF (442.342)</td>
<td></td>
<td></td>
<td></td>
<td>(a) Range of motion, ambulation, turning and positioning and other activities;</td>
<td></td>
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</tr>
<tr>
<td>F173</td>
<td>SNF (405.1124w)</td>
<td></td>
<td></td>
<td></td>
<td>(b) Assistance and instruction in the activities of daily living such as feeding, dressing, grooming, oral hygiene and toilet activities;</td>
<td></td>
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</tr>
<tr>
<td>F174</td>
<td>ICF (442.342)</td>
<td></td>
<td></td>
<td></td>
<td>(c) Remotivation therapy and/or reality orientation when appropriate.</td>
<td></td>
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</tr>
<tr>
<td>F175</td>
<td>SNF (405.1124w)</td>
<td></td>
<td></td>
<td></td>
<td>3. These activities are coordinated with other resident care services.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>NURSING SERVICES</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
<td></td>
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</tr>
<tr>
<td>118</td>
<td>1. Each resident is provided with the amount of food and fluid on a daily basis necessary to maintain their appropriate minimum average weight. Between meal feedings are offered and the amount consumed is observed. Daily food and fluid intake is observed and encouraged.</td>
<td></td>
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</tr>
<tr>
<td>118</td>
<td>2. Each resident needing assistance in eating or drinking is provided prompt assistance. Specific self-help devices are available when necessary.</td>
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</tr>
<tr>
<td>118</td>
<td>3. Deviations from normal food and fluid intake are recorded and reported to the charge nurse and the attending physician.</td>
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</table>
### NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>NURSING SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F183</td>
<td>Administration of Drugs</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>SNF (405.1124(g)) (Standard)</td>
<td>□ MET</td>
<td>□ NOT MET</td>
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<tr>
<td>F184</td>
<td>ICF (442.337) (Standard)</td>
<td>□ MET</td>
<td>□ NOT MET</td>
<td></td>
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</tr>
<tr>
<td>F185</td>
<td>The resident is identified prior to administration of a drug.</td>
<td></td>
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</tr>
<tr>
<td>F186</td>
<td>Drugs and biologicals are administered as soon as possible after doses are prepared.</td>
<td></td>
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</tr>
<tr>
<td>F187</td>
<td>Administered by same person who prepared the doses for administration except under single unit dose package distribution systems.</td>
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<tr>
<td>F188</td>
<td>Exception: ICF residents may self-administer medication only with their physician’s permission.</td>
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</table>

#### Conformance with Physician Drug Orders

<table>
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<th>CODE</th>
<th>NURSING SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
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</thead>
<tbody>
<tr>
<td>F189</td>
<td>SNF (405.1124(h)) (Standard)</td>
<td>□ MET</td>
<td>□ NOT MET</td>
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<tr>
<td>F190</td>
<td>ICF (442.334) (Standard)</td>
<td>□ MET</td>
<td>□ NOT MET</td>
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</tr>
<tr>
<td>F191</td>
<td>Drugs are administered in accordance with written orders of the attending physician.</td>
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<tr>
<td>F192</td>
<td>Drug Error Rate %</td>
<td></td>
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<td></td>
<td>(See Form HCFA-522)</td>
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</tr>
<tr>
<td>CODE</td>
<td>DIETETIC SERVICES (CONDITION OF PARTICIPATION)</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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<tr>
<td>F193</td>
<td>SNF (405.1125)</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>A. Menus and Nutritional Adequacy</td>
</tr>
<tr>
<td>F194</td>
<td>ICF (428.332) (Standard)</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>Menus are planned and followed to meet the nutritional needs of each resident in accordance with physicians’ orders and, to the extent medically possible, based on the recommended dietary allowances of the Food and Nutrition Board of the National Research Council, National Academy of Sciences.</td>
</tr>
<tr>
<td>F195</td>
<td>SNF (405.1125(0)) (Standard)</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>B. Therapeutic Diets</td>
</tr>
<tr>
<td>F196</td>
<td></td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>1. Therapeutic diets are prescribed by the attending physician.</td>
</tr>
<tr>
<td>F197</td>
<td>SNF (405.1125(f)) (Standard)</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>2. Therapeutic menus are planned in writing, prepared, and served as ordered with supervision from the dietician and advice from the physician when necessary.</td>
</tr>
<tr>
<td>F198</td>
<td>Number of Regular Diets</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F199</td>
<td>Number of Therapeutic Diets</td>
<td></td>
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<td></td>
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<tr>
<td>F200</td>
<td>Number of Mechanically Altered Diets</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>F201</td>
<td>Number of Tube Feedings</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>CODED TEXT</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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<td>------</td>
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</tr>
<tr>
<td>F206</td>
<td>C. Preparation</td>
<td>□ MET</td>
<td>□ NOT MET</td>
<td></td>
<td>1. Food is prepared by methods that conserve its nutritive value and flavor.</td>
</tr>
<tr>
<td>F205</td>
<td>2. Meals are palatable, served at proper temperatures. They are cut, ground, chopped, pureed or in a form which meets individual resident needs.</td>
<td></td>
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</tr>
<tr>
<td>F207</td>
<td>3. If a resident refuses food served, appropriate substitutes of similar nutritive value are offered.</td>
<td></td>
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</tr>
<tr>
<td>F208</td>
<td>D. Frequency</td>
<td>□ MET</td>
<td>□ NOT MET</td>
<td></td>
<td>1. At least three meals are served daily at regular hours with not more than a 14-hour span between a substantial evening meal and breakfast.</td>
</tr>
<tr>
<td>F209</td>
<td>2. To the extent medically possible, bedtime nourishments are offered to all residents.</td>
<td></td>
<td></td>
<td></td>
<td><em>Exception: Not required for ICF Residents.</em></td>
</tr>
<tr>
<td>F211</td>
<td>E. Staffing</td>
<td>□ MET</td>
<td>□ NOT MET</td>
<td></td>
<td>1. Food service personnel are on duty daily over a period of 12 or more hours.</td>
</tr>
<tr>
<td>CODE</td>
<td>SPECIALIZED REHABILITATIVE SERVICES</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
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</tr>
<tr>
<td>F214</td>
<td>SNF (405.1120) (Condition of Participation)</td>
<td>□ MET</td>
<td>□ NOT MET</td>
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<tr>
<td>F215</td>
<td>SNF-405,1120(b)(3)</td>
<td>□ MET</td>
<td>□ NOT MET</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>ICF-442,343</td>
<td>□ MET</td>
<td>□ NOT MET</td>
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</tr>
</tbody>
</table>

A. Plan of Care

Rehabilitative services are provided under a written plan of care, initiated by the attending physician and developed in consultation with appropriate therapist(s) and the nursing service.

B. Therapy

Therapy is provided according to orders of the attending physician in accordance with accepted professional practices by qualified therapists or qualified assistants.

C. Progress

1. A report of the resident's progress is communicated to the attending physician within 2 weeks of the initiation of specialized rehabilitative services.

   Exception: ICF resident's progress must be reviewed regularly.
<table>
<thead>
<tr>
<th>CODE</th>
<th>SPECIALIZED REHABILITATIVE SERVICES/PHARMACEUTICAL SERVICES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F220</td>
<td>2. The resident's progress is thereafter reviewed regularly, and the plan of rehabilitative care is reevaluated as necessary, but at least every 30 days, by the physician and the therapist. Exceptions: ICF residents' plans must be revised as necessary.</td>
<td>☑</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
<tr>
<td>F221</td>
<td>PHARMACEUTICAL SERVICES (CONDITION OF PARTICIPATION)</td>
<td>☑</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
<tr>
<td></td>
<td>SNF (405.1127)</td>
<td>☑</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
<tr>
<td>F222</td>
<td>A. Supervision</td>
<td>☑</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
<tr>
<td></td>
<td>SNF (405.1127(p)) (Standard)</td>
<td>☑</td>
<td>☐</td>
<td>☐</td>
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<tr>
<td>F223</td>
<td>ICF (442.336) (Standard)</td>
<td>☑</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
<tr>
<td>F224</td>
<td>The pharmacist reviews the drug regimen of each resident at least monthly and reports any irregularities to the medical director and administrator.</td>
<td>☑</td>
<td>☐</td>
<td>☐</td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>PHARMACEUTICAL SERVICES</td>
<td>LABORATORY AND RADIOLOGIC SERVICES/SOCIAL SERVICES</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
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<tr>
<td>P225</td>
<td>SNF (405.1127)(g) (Standard)</td>
<td>☐ MET ☐ NOT MET</td>
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<tr>
<td>P226</td>
<td>ICF (442.333) (Standard)</td>
<td>☐ MET ☐ NOT MET</td>
<td></td>
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</tr>
<tr>
<td>P227</td>
<td>The labeling of drugs and biologicals is based on currently accepted professional principles and includes the appropriate accessory and cautionary instructions as well as an expiration date where applicable.</td>
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</tr>
<tr>
<td>P228</td>
<td>SNF (405.1128)(a)</td>
<td>☐ MET ☐ NOT MET</td>
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<tr>
<td>P229</td>
<td>SNF (405.1128)(a) (Standard)</td>
<td>☐ MET ☐ NOT MET</td>
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</tr>
</tbody>
</table>

**Provision of Services**

1. All services are provided only on the orders of a physician.
2. The attending physician is notified promptly of diagnostic findings.
3. Signed and dated reports of a clinical laboratory, X-ray and other diagnostic services are filed with the resident's medical record.
<table>
<thead>
<tr>
<th>CODE</th>
<th>SOCIAL SERVICES/ACTIVITIES</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F233</td>
<td>SNF (405.1130)</td>
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<tr>
<td>F234</td>
<td>SNF (405.1130)(a) (Standard)</td>
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<tr>
<td>F235</td>
<td>ICF (442.344) (Standard)</td>
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</tbody>
</table>

**SOCIAL SERVICES (CONDITION OF PARTICIPATION)**

A. Plan

The medically related social and emotional needs of the resident are identified.

B. Provision of Services

1. Services are provided to meet the social and emotional needs by the facility or by referral to an appropriate social agency.

2. If financial assistance is indicated, arrangements are made promptly for referral to an appropriate agency.
<table>
<thead>
<tr>
<th>CODE</th>
<th>ACTIVITIES</th>
<th>VERIFICAION</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F241</td>
<td>ICF (442.345) (Standard)</td>
<td>☑ MET   ☐ NOT MET</td>
<td>An ongoing program of meaningful activities is provided based on identified needs and interests of each resident. It is designed to promote opportunities for engaging in normal pursuits, including religious activities of their choice, if any.</td>
</tr>
<tr>
<td>F242</td>
<td>1. An ongoing program of meaningful activities is provided based on identified needs and interests of each resident. It is designed to promote opportunities for engaging in normal pursuits, including religious activities of their choice, if any.</td>
<td></td>
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</tr>
<tr>
<td>F243</td>
<td>2. Unless contraindicated by the attending physicians each resident is encouraged to participate in the activities program.</td>
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<tr>
<td>F244</td>
<td>3. The activities promote the physical, social and mental well-being of the resident.</td>
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<tr>
<td>F245</td>
<td>4. Equipment is maintained in good working order.</td>
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<tr>
<td>F246</td>
<td>5. Supplies and equipment are available.</td>
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</tr>
<tr>
<td>CODE</td>
<td>MEDICAL REASON</td>
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<tr>
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<tr>
<td>F248</td>
<td>SNF (405.1132)(c)(Standard)</td>
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<tr>
<td>F249</td>
<td>ICF (442.31E)(Standard)</td>
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<tr>
<td>F250</td>
<td>1. The medical record contains sufficient information to identify the resident clearly, to justify diagnoses and treatment, and to document results accurately.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>MEDICAL RECORDS</td>
<td>YES</td>
<td>NO</td>
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<tr>
<td>F251</td>
<td>2. The medical record contains the following information:</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>a. Identification information</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F252</td>
<td>b. Admission data including past medical and social history</td>
<td></td>
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</tr>
<tr>
<td>F253</td>
<td>c. Transfer form, discharge summary from any transferring facility</td>
<td></td>
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</tr>
<tr>
<td>F254</td>
<td>d. Report of resident's attending physician</td>
<td></td>
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<tr>
<td>F255</td>
<td>e. Report of physical examinations</td>
<td></td>
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<tr>
<td>F256</td>
<td>f. Reports of physicians' periodic evaluations and progress notes</td>
<td></td>
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</tr>
<tr>
<td>F257</td>
<td>g. Diagnostic reports and therapeutic orders</td>
<td></td>
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</tr>
<tr>
<td>F258</td>
<td>h. Reports of treatments</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F259</td>
<td>i. Medications administered</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F260</td>
<td>j. An overall plan of care setting forth goals to be accomplished through each service's designed activities, therapies and treatments.</td>
<td></td>
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</tr>
<tr>
<td>F261</td>
<td>k. Assessments and goals of each service's plan of care</td>
<td></td>
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</tr>
<tr>
<td>F262</td>
<td>l. Treatments and services rendered</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F263</td>
<td>m. Progress notes</td>
<td></td>
<td></td>
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<tr>
<td>F264</td>
<td>n. All symptoms and other indications of illness or injury including date, time and action taken regarding each problem.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>NAME OF FACILITY</td>
<td>TRANSFER AGREEMENT (CONDITION OF PARTICIPATION)</td>
<td>YES</td>
</tr>
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<td>------</td>
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<td>-----------------------------------------------</td>
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<tr>
<td>F265</td>
<td>SNF (405.1133)</td>
<td>☐ MET ☐ NOT MET</td>
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<tr>
<td>F266</td>
<td>SNF (405.1133(a)) (Standard)</td>
<td>☐ MET ☐ NOT MET</td>
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</tr>
<tr>
<td>F267</td>
<td>ICF (442.316) (Standard)</td>
<td>☐ MET ☐ NOT MET</td>
<td></td>
</tr>
</tbody>
</table>

F268 A. Whenever the attending physician determines that a transfer is medically appropriate between a hospital or a facility providing more specialized care and the nursing facility, admission to the new facility shall be effected in a timely manner.

F269 B. Information necessary for providing care and treatment to transferred individuals is provided.
<table>
<thead>
<tr>
<th>CODE</th>
<th>PHYSICAL ENVIRONMENT</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F270</td>
<td>SNF (405.1134)</td>
<td></td>
<td></td>
<td></td>
<td>PHYSICAL ENVIRONMENT (CONDITION OF PARTICIPATION)</td>
</tr>
<tr>
<td></td>
<td>A. Nursing Unit</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F271</td>
<td>SNF (405.1134(d))</td>
<td></td>
<td></td>
<td></td>
<td>(Standard)</td>
</tr>
<tr>
<td>F272</td>
<td>1. The unit is properly equipped for preparation and storage of drugs and biologicals.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F273</td>
<td>2. Utility and storage rooms are adequate in size.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F274</td>
<td>3. The unit is equipped to register resident calls with a functioning communication system from resident areas including resident rooms and toilet and bathing facilities.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>B. Dining and Activities Area</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F275</td>
<td>SNF (405.1134(g))</td>
<td></td>
<td></td>
<td></td>
<td>(Standard)</td>
</tr>
<tr>
<td>F276</td>
<td>ICF (442.329)</td>
<td></td>
<td></td>
<td></td>
<td>(Standard)</td>
</tr>
<tr>
<td>F277</td>
<td>1. The facility provides one or more clean, orderly and appropriately furnished rooms of adequate size designated for resident dining and resident activities.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F278</td>
<td>2. Dining and activity rooms are well lighted and ventilated.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F279</td>
<td>3. Any multipurpose room used for dining and resident activities has sufficient space to accommodate all activities and prevent their interference with each other.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### NAME OF FACILITY

<table>
<thead>
<tr>
<th>CODE</th>
<th>PHYSICAL ENVIRONMENT</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F280</td>
<td>SNF (405.1134(e)) (Standard)</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

**INDICATORS C AND D APPLY TO THIS STANDARD FOR SNF**

#### C. Resident Rooms

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>F281</td>
<td>ICF (442.325) (Standard)</td>
</tr>
</tbody>
</table>

1. Single resident rooms have at least 100 square feet.
2. Multiple resident rooms have no more than four residents and at least 80 square feet per resident.
3. Each room is equipped with or conveniently located near toilet and bathing facilities.
4. There is capability of maintaining privacy in each.
5. There is adequate storage space for each resident.
6. There is a comfortable and functioning bed and chair plus a functional cabinet and light.
7. The resident call system functions in resident rooms.
8. Each room is designed and equipped for adequate nursing care and the comfort and privacy of the residents.
9. Each room is at or above grade level.
10. Each room has direct access to a corridor and outside exposure.

*Exception: Not required for ICF residents.*
<table>
<thead>
<tr>
<th>CODE</th>
<th>PHYSICAL ENVIRONMENT</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F292</td>
<td>D. Toilet and Bath Facilities</td>
<td></td>
<td></td>
<td></td>
<td>ICF (442.326) (Standard) □ MET □ NOT MET</td>
</tr>
<tr>
<td>F293</td>
<td>1. Facilities are clean, sanitary and free of odors.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F294</td>
<td>2. Facilities have safe and comfortable hot water temperatures.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F295</td>
<td>3. Facilities maintain privacy.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F296</td>
<td>4. Facilities have grab bars and other safeguards against slipping.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F297</td>
<td>5. Facilities have fixtures in good condition.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F298</td>
<td>6. The resident call system functions in toilet and bath facilities.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F299</td>
<td>E. Social Service Area</td>
<td></td>
<td></td>
<td></td>
<td>SNF (405.1130(b)) (Standard) □ MET □ NOT MET</td>
</tr>
<tr>
<td>F300</td>
<td>1. Ensures privacy for social service interviewing.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F301</td>
<td>2. Adequate space for clerical and interviewing functions is provided.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F302</td>
<td>3. Facilities are easily accessible to residents and staff.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>PHYSICAL ENVIRONMENT</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td>EXPLANATORY STATEMENT</td>
</tr>
<tr>
<td>------</td>
<td>-----------------------</td>
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<td>-----</td>
<td>-----------------------</td>
</tr>
<tr>
<td>F303</td>
<td>SNF (405.1126(a)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F304</td>
<td>ICF (442.328(a))</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F305</td>
<td>1. Space is adequate for proper use of equipment by all residents receiving treatments.</td>
<td></td>
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</tr>
<tr>
<td>F306</td>
<td>2. Equipment is safe and in proper working condition.</td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>F307</td>
<td>SNF (405.1134(f)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F308</td>
<td>ICF (442.328(b))</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F309</td>
<td>1. Single rooms with private toilet and handwashing facilities are available for isolating residents.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F310</td>
<td>2. Precautionary signs are used to identify these rooms when in use.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F311</td>
<td>SNF (405.1134(j)) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F312</td>
<td>ICF (442.324) (Standard)</td>
<td>☐ MET</td>
<td>☐ NOT MET</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F313</td>
<td>1. All common resident areas are clean, sanitary and free of odors.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F314</td>
<td>2. Provision is made for adequate and comfortable lighting levels in all areas.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F315</td>
<td>3. There is limitation of sounds at comfort levels.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CODE</td>
<td>PHYSICAL ENVIRONMENT</td>
<td>YES</td>
<td>NO</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>------</td>
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</tr>
<tr>
<td>F316</td>
<td>4. A comfortable room temperature is maintained.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>F317</td>
<td>5. There is adequate ventilation through windows or mechanical means or a combination of both.</td>
<td></td>
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</tr>
<tr>
<td>F318</td>
<td>6. Corridors are equipped with firmly secured handrails on each side.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F319</td>
<td>7. Staff are aware of procedures to ensure water to all essential areas in the event of loss of normal supply.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### I. Maintenance of Building and Equipment

- **SNF (405.1134(h)) (Standard)**
  - [ ] MET
  - [ ] NOT MET

#### J. Dietary Service Area

- **SNF (405.1134(h)) (Standard)**
  - [ ] MET
  - [ ] NOT MET

**Indicators J thru L apply to ICFs.**

- **F320**
- **F321**
- **F322**
- **F323**
- **F324**
- **F325**
- **F326**
- **F327**
- **F328**
<table>
<thead>
<tr>
<th>NAME OF FACILITY</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>CODE</th>
<th>PHYSICAL ENVIRONMENT/INFECTION CONTROL</th>
<th>YES</th>
<th>NO</th>
<th>N/A</th>
<th>EXPLANATORY STATEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>F329</td>
<td>E. HYGIENE OF DENTAL STAFF</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F330</td>
<td>SNF (405.1135(b)) (Standard)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F331</td>
<td>F. DENTAL SANITARY CONDITIONS</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F332</td>
<td>SNF (405.1135(g))</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F333</td>
<td>H. EMERGENCY POWER</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F334</td>
<td>SNF (405.1135(b))</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F335</td>
<td>1. An emergency source of electrical power necessary to protect the health and safety of residents is available in the event the normal electrical supply is interrupted.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F336</td>
<td>2. Emergency power is adequate at least for lighting in all means of egress; equipment to maintain fire detection, alarm, and extinguishing systems; and life safety support systems.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F337</td>
<td>3. Emergency power is provided by an emergency electrical generator located on the premises where life support systems are used.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F338</td>
<td>INFECTION CONTROL (CONDITION OF PARTICIPATION)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F339</td>
<td>SNF (405.1135)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F340</td>
<td>A. Infection Control</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F341</td>
<td>SNF (405.1135(b))</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F342</td>
<td>Aseptic and isolation techniques are followed by all personnel.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## NAME OF FACILITY

### CODE | INFECTION CONTROL/DISASTER PREPAREDNESS | YES | NO | N/A | EXPLANATORY STATEMENT
--- | --- | --- | --- | --- | ---
6. Sanitation
- F341 SNF (405.1135(c)) (Standard) | Met | Not Met | | | 
- F342 The facility maintains a safe, clean, and orderly interior. | | | | | 
C. Linen
- F343 SNF (405.1135(d)) (Standard) | Met | Not Met | | | 
- F344 ICF (442.327) (Standard) | Met | Not Met | | | 
- F345 1. The facility has available at all times a quantity of linen essential for proper care and comfort of residents. | | | | | 
- F346 2. Linens are handled, stored, processed, and transported in such a manner as to prevent the spread of infection. | | | | | 
D. PEST CONTROL
- F347 SNF (405.1135(e)) (Standard) | Met | Not Met | | | 
- F348 ICF (442.315(c)) (Standard) | Met | Not Met | | | 
- F349 The facility is maintained free from insects and rodents. | | | | | 
### DISASTER PREPAREDNESS (CONDITION OF PARTICIPATION)
- F350 SNF (405.1136) | Met | Not Met | | | 
- F351 SNF (405.1136(a)) (Standard) | Met | Not Met | | | 
- F352 ICF (442.313) (Standard) | Met | Not Met | | | 
**INDICATORS A AND B APPLY TO THIS STANDARD FOR ICFs:**

### A. Disaster Plan
- F353 1. Facility staff are aware of plans, procedures to be followed for fire, explosion or other disaster.
<table>
<thead>
<tr>
<th>CODE</th>
<th>DISASTER PREPAREDNESS</th>
</tr>
</thead>
<tbody>
<tr>
<td>F354</td>
<td>2. Facility staff are knowledgeable about evacuation routes.</td>
</tr>
<tr>
<td>F355</td>
<td>3. Facility staff are aware of their specific responsibilities in regard to evaluation and protection of residents.</td>
</tr>
<tr>
<td>F356</td>
<td>4. Facility staff are aware of methods of containing fire.</td>
</tr>
</tbody>
</table>

**B. Drills**

<table>
<thead>
<tr>
<th>CODE</th>
<th>DISASTER PREPAREDNESS</th>
</tr>
</thead>
<tbody>
<tr>
<td>F357</td>
<td>SNF (405.1136(b))(Standard) □ MET □ NOT MET</td>
</tr>
<tr>
<td>F358</td>
<td>1. All employees are trained, as part of their employment orientation in all aspects of preparedness for any disaster.</td>
</tr>
<tr>
<td>F359</td>
<td>2. Facility staff participate in ongoing training and drills in all procedures so that each employee promptly and correctly carries out a specific role in case of a disaster.</td>
</tr>
</tbody>
</table>
§ 488.105
42 CFR Ch. IV (10–1–00 Edition)

SKILLED NURSING FACILITY & INTERMEDIATE CARE FACILITY
SURVEY REPORT — PART B
OFFICIAL DATA EXTRACT
(To be used with 2–88 Revision of Form HCFA-418)

<table>
<thead>
<tr>
<th>PROVIDER NO.</th>
<th>FACILITY NAME</th>
<th>SURVEY DATE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

SURVEY TEAM COMPOSITION

*F1: INDICATE THE NUMBER OF SURVEYORS ACCORDING TO DISCIPLINE:

| A. _____ | ADMINISTRATOR | H. _____ | LIFE SAFETY CODE SPECIALIST |
| B. _____ | NURSE | I. _____ | LABORATORIAN |
| C. _____ | DIETITIAN | J. _____ | SANITARIAN |
| D. _____ | PHARMACIST | K. _____ | THERAPIST |
| E. _____ | RECORDS ADMINISTRATOR | L. _____ | PHYSICIAN |
| F. _____ | SOCIAL WORKER | M. _____ | NATIONAL INSTITUTE OF MENTAL HEALTH |
| G. _____ | QUALIFIED MENTAL RETARDATION PROFESSIONAL | N. _____ | OTHER |

NOTE: MORE THAN ONE DISCIPLINE MAY BE MARKED FOR SURVEYORS QUALIFIED IN MULTIPLE DISCIPLINES.

*F2: INDICATE THE TOTAL NUMBER OF SURVEYORS ONSITE: _____

*F3: DRUG ERROR RATE: _____% (Round % to nearest whole number.)

*SFS Survey Form Indicator (Check one)

Traditional Survey

New LTC Survey

(1) ☐ (2) ☐

NOTE: PLEASE ATTACH COPY OF PAGES 2, 14 AND 15.

*Mandatory

Form HCFA-418E (2–88)

### TOUR NOTES WORKSHEET

**INDEPTH SAMPLE**

<table>
<thead>
<tr>
<th>Facility Type</th>
<th>0-50</th>
<th>61-190</th>
<th>191-290</th>
<th>291+</th>
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</thead>
<tbody>
<tr>
<td>Census</td>
<td>256</td>
<td>156</td>
<td>155</td>
<td>154</td>
</tr>
<tr>
<td>Sample</td>
<td>255</td>
<td>210</td>
<td>155</td>
<td>154</td>
</tr>
</tbody>
</table>

**GROOMING/PERSOAL HYGIENE**

- Positioning

- Assistive Devices

- Ambulation

- Restraints

- Hydration

- Infection Control

- Patient Rights

- Other

---

*[42 CFR Ch. IV (10–1–00 Edition)](http://example.com)*

[ln 664]
## OBSERVATION / INTERVIEW RECORD REVIEW WORKSHEET

<table>
<thead>
<tr>
<th>PROVIDER NUMBER</th>
<th>SURVEY DATE</th>
<th>OBSERVATION/INTERVIEW OF: (RESIDENT IDENTIFIER)</th>
</tr>
</thead>
</table>

### INSTRUCTIONS

1. Observe each resident in sample to identify ADL needs and potential problems. Check appropriate boxes.
2. Interview only residents in sample who are capable and willing.
3. Review each resident's record to ensure assessments, plans, interventions, and evaluations are appropriate and current.
4. Note deficiencies on survey report form after reviewing all residents in sample.

### RESIDENT NEEDS

<table>
<thead>
<tr>
<th>ADLs</th>
<th>GROOMING/HYGIENE</th>
<th>RESIDING</th>
<th>COLOSTOMY/CONTINENT</th>
<th>RESPIRATORY</th>
<th>REHYDRATION NURSING</th>
<th>ACTIVITY NEEDS</th>
<th>PATIENT RIGHTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bathing</td>
<td>Dressing</td>
<td>Grooming</td>
<td>Personal卫生</td>
<td>Oral Care</td>
<td>Not Well Regular</td>
<td>Incontinence</td>
<td>Impaired Communication</td>
</tr>
<tr>
<td>Feeding</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
</tbody>
</table>

### POSITIONING

- Head/Eye Support
- Not Perform
- Perform
- Not Perform

### DRESSING

- Present
- Not Present

### FOOD/PARTIAL FEEDING

- Present
- Not Present

### INFECTIONS

- Present
- Not Present

### BOWEL/BLADDER

- Present
- Not Present

### Weights

- Present
- Not Perform

### RESPIRATORY

- Present
- Not Perform

### PULMONARY

- Present
- Not Perform

### DEMENTIA

- Present
- Not Perform

### OTHER PROBLEMS

- Present
- Not Perform

### INTERVENTIONS

- Present
- Not Perform

### PASSIVE MISTREATMENT

- Present
- Not Perform

### BREVITY

- Present
- Not Perform

### NUTRITIONAL/FEEDING NEEDS

- Present
- Not Perform

### FELD'S HOUND

- Present
- Not Perform

### WEIGHT

- Present
- Not Perform

### SISSON'S PROBLEM

- Present
- Not Perform

### FEEDING

- Present
- Not Perform

### SUGGESTIONS

- Present
- Not Perform

### OTHER

- Present
- Not Perform

### NOTES:

- Present
- Not Perform

---

Form HCFA 124 (3-89)  SEE 40-2/6/82

Health Care Financing Administration, HHS

868.105
# DRUG PASS WORKSHEET

<table>
<thead>
<tr>
<th>IDENTIFIER</th>
<th>FAVOR</th>
<th>PASS</th>
<th>RECORD</th>
</tr>
</thead>
<tbody>
<tr>
<td>RESIDENT'S FULL NAME, ROOM NUMBER, TIME</td>
<td>DRUG PRESCRIPTION NAME, DOSE AND FORM</td>
<td>OBSERVATION OF ADMINISTRATION</td>
<td>DRUG ORDER WRITTEN AS (IF DIFFERS FROM ADRAMS ONLY)</td>
</tr>
</tbody>
</table>

**INSTRUCTIONS**

2. Record Observation of each Opportunity.
3. Compare Observation Notes with Physician Orders.
4. Calculate and Note Error Rate.
5. Note Deficiencies on Survey Report Form.

**DEFICIENCY FORMULA**

\[
\text{Deficiency} = \frac{\text{Significant + Non-significant}}{\text{Doses Given + Doses Ordered But Not Given}} \times 100 \times \text{Error Rate}
\]
§ 488.105 42 CFR Ch. IV (10–1–00 Edition)

DRUG ERROR CALCULATION
(SEE SOM Appendix N Part 2)

How to Calculate a Medication Error Rate—in calculating the percentage of errors, the numerator in the ratio is the total number of errors that you observe, both significant and non-significant. The denominator is all the doses observed being administered plus the doses ordered but not administered. The equation for calculating a medication error rate is as follows:

Medication Error Rate = Number of errors observed \times \frac{100}{	ext{Opportunities for errors}}

Where: Opportunities for errors equals the number of doses administered plus the number of doses ordered but not administered.

Comments

For example, you observed the administration of drugs to 20 patients. There were a total of 47 drugs administered (47 opportunities for errors). At the completion of the reconciliation of your Observations with the physicians' orders, you find that three medication errors were made in administration and one medication was omitted (ordered but not administered). The omitted dose is included in both the numerator and the denominator. Therefore, following the above formula, your equation would be as follows:

\[
\frac{3 + 1}{47 + 1} \times 100 = 8.3\%
\]
### DINING AREA & EATING ASSISTANCE WORKSHEET

<table>
<thead>
<tr>
<th>PROVIDER NUMBER</th>
<th>SURVEY DATE</th>
</tr>
</thead>
</table>

**INSTRUCTIONS**

1. **DINING AREA AND MEALS**
   - a. Size does not restrict movement.
   - b. Accommodates all residents.
   - c. Cleanliness.
   - d. Adequate/comfortable lighting.
   - e. Adequate/comfortable ventilation.

2. **SERVING OF MEALS**
   - a. Number of meals/time span between meal.
   - b. Conformance to physicians order.
   - c. Nutritional adequacy.
   - d. Adequacy of portions.
   - e. Residents eat approximately 75% of meals.
   - f. Puree dishes served individually.
   - g. Food cut, chopped or ground for individual resident needs.
   - h. Acceptable taste.
   - i. Proper temperature.
   - j. Plates covered.

---

*Note: Assistance Provided*
§ 488.110 Procedural Guidelines. 

The purpose for implementing a new SNF/ICF survey process is to assess whether the quality of care, as intended by the law and regulations, as needed by the resident, is actually being provided in nursing homes. Although the onsite review procedures have been changed, facilities must continue to meet all applicable Conditions/Standards in order to participate in Medicare/Medicaid programs. That is, the methods used to evaluate facilities must continue to meet the law and regulations, as needed by the resident.

### 2. SERVING OF MEALS *(continued)*

<table>
<thead>
<tr>
<th>Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>k.</td>
<td>Served promptly.</td>
</tr>
<tr>
<td>l.</td>
<td>Residents ready for meal when served.</td>
</tr>
<tr>
<td>m.</td>
<td>Attractive.</td>
</tr>
<tr>
<td>n.</td>
<td>Utensils available.</td>
</tr>
<tr>
<td>o.</td>
<td>Functional trays for bedfast residents.</td>
</tr>
<tr>
<td>p.</td>
<td>Salt, pepper, sugar, other condiments on resident's trays unless contraindicated.</td>
</tr>
<tr>
<td>q.</td>
<td>Medically able residents eating in dining area.</td>
</tr>
<tr>
<td>r.</td>
<td>Bedtime nourishment offered.</td>
</tr>
</tbody>
</table>

### 3. SUPERVISION OF RESIDENT NUTRITION

<table>
<thead>
<tr>
<th>Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>a.</td>
<td>Prompt assistance.</td>
</tr>
<tr>
<td>b.</td>
<td>Proper assistance (spoon-feeding, supervision or instruction to develop eating skills).</td>
</tr>
<tr>
<td>c.</td>
<td>Courteous and unhurried assistance.</td>
</tr>
<tr>
<td>d.</td>
<td>Self-help devices present (straws, easy grip utensils, special cup, etc.).</td>
</tr>
<tr>
<td>e.</td>
<td>Intake recorded/deviations from normal are reported.</td>
</tr>
</tbody>
</table>
compile information about compliance with law and regulations are changed; the law and regulations themselves are not changed. The new process differs from the traditional process, principally in terms of its emphasis on resident outcomes. In ascertaining whether residents' grooming and personal hygiene needs are met, for example, surveyors will no longer routinely evaluate a facility's written policies and procedures. Instead, surveyors will observe residents in order to make that determination. In addition, surveyors will confirm, through interviews with residents and staff, that such needs are indeed met on a regular basis. In most reviews, then, surveyors will ascertain whether the facility is actually providing the required and needed care and services, rather than whether the facility is capable of providing the care and services.

THE OUTCOME-ORIENTED SURVEY PROCESS—SKILLED NURSING FACILITIES (SNFs) AND INTERMEDIATE CARE FACILITIES (ICFs)

(a) General.
(b) The Survey Tasks.
(c) Task 1—Entrance Conference.
(d) Task 2—Resident Sample—Selection Methodology.
(e) Task 3—Tour of the Facility.
(f) Task 4—Observation/Interview/Medical Record Review (including drug regimen review).
(g) Task 5—Drug Pass Observation.
(h) Task 6—Dining Area and Eating Assistance Observation.
(i) Task 7—Forming the Deficiency Statement (if necessary).
(j) Task 8—Exit Conference.
(k) Plan of Correction.
(l) Followup Surveys.
(m) Role of Surveyor.
(n) Confidentiality and Respect for Resident Privacy.
(o) Team Composition.
(p) Type of Facility—Application of SNF or ICF Regulations.

(a) General. A complete SNF/ICF facility survey consists of three components:
- Life Safety Code requirements;
- Administrative and structural requirements (Part A of the Survey Report, Form HCFA-525); and
- Direct resident care requirements (Part B of the Survey Report, Form HCFA-519), along with the related worksheets (HCFA-520 through 524).

Use this survey process for all surveys of SNFs and ICFs—whether free-standing, distinct parts, or dually certified. Do not use this process for surveys of Intermediate Care Facilities for Mentally Retarded (ICFs/MR), swing-bed hospitals or skilled nursing sections of hospitals that are not separately certified as SNF distinct parts. Do not announce SNF/ICF surveys ahead of time.

(b) The Survey Tasks. Listed below are the survey tasks for easy reference:
- Task 1. Entrance Conference.
- Task 2. Resident Sample—Selection Methodology.
- Task 4. Observation/Interview/Medical Record. Review of Each Individual in the Resident Sample (including drug regimen review).
- Task 5. Drug Pass Observation.
- Task 6. Dining Area and Eating Assistance Observation.
- Task 7. Forming the Deficiency Statement (if necessary).
- Task 8. Exit Conference.

(c) Task 1—Entrance Conference. Perform these activities during the entrance conference in every certification and recertification survey:
- Introduce all members of the team to the facility staff, if possible, even though the whole team may not be present for the entire entrance conference. (All surveyors wear identification tags.)
- Explain the SNF/ICF survey process as resident centered in focus, and outline the basic steps.
- Ask the facility for a list showing the names of residents by room number with each of the following care needs/treatments identified for each resident to whom they apply:
  - Decubitus care
  - Restraints
  - Catheters
  - Injections
  - Parenteral fluids
  - Rehabilitation service
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Colostomy/ileostomy care
Respiratory care
Tracheostomy care
Suctioning
Tube feeding

Use this list for selecting the resident sample.

- Ask the facility to complete page 2 of Form HCFA-519 (Resident Census) as soon as possible, so that the information can further orient you to the facility’s population. In a survey of a SNF with a distinct part ICF, you may collect two sets of census data. However, consolidate the information when submitting it to the regional office. You may modify the Resident Census Form to include the numbers of licensed and certified beds, if necessary.

- Ask the facility to post signs on readily viewed areas (at least one on each floor) announcing that State surveyors are in the facility performing an “inspection,” and are available to meet with residents in private. Also indicate the name and telephone number of the State agency. Hand-printed signs with legible, large letters are acceptable.

- If the facility has a Resident Council, make mutually agreeable arrangements to meet privately with the president and officers and other individuals they might invite.

- Inform the facility that interviews with residents and Resident Councils are conducted privately, unless they independently request otherwise, in order to enhance the development of rapport as well as to allay any resident anxiety. Tell the facility that information is gathered from interviews, the tour, observations, discussions, record review, and facility officials. Point out that the facility will be given an opportunity to respond to all findings.

(d) Task 2—Resident Sample—Selection Methodology. This methodology is aimed at formulating a sample that reflects the actual distribution of care needs/treatments in the facility population.

Primarily performed on a random basis, it also ensures representation in the sample of certain care needs and treatments that are assessed during the survey.

(1) Sample Size. Calculate the size of the sample according to the following guide:

<table>
<thead>
<tr>
<th>Number of residents in facility</th>
<th>Number of residents in sample1</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-60 residents</td>
<td>25% of residents (minimum—10).</td>
</tr>
<tr>
<td>61-120 residents</td>
<td>20% of residents (minimum—15).</td>
</tr>
<tr>
<td>121-200 residents</td>
<td>15% of residents (minimum—24).</td>
</tr>
<tr>
<td>201+ residents</td>
<td>10% of residents (minimum—30).</td>
</tr>
<tr>
<td></td>
<td>Maximum—50.</td>
</tr>
</tbody>
</table>

Note that the calculation is based on the resident census, not beds. After determining the appropriate sample size, select residents for the sample in a random manner. You may, for example, select every fifth resident from the resident census, beginning at a random position on the list. For surveys of dually certified facilities or distinct part SNFs/ICFs, first use the combined SNF/ICF population to calculate the size of the sample, and then select a sample that reflects the proportions of SNF and ICF residents in the facility’s overall population.

(2) Special Care Needs/Treatments. The survey form specifies several care needs/treatments that must always be reviewed when they apply to any facility residents. These include:

- Decubitus Care
- Restraints
- Catheters
- Injections, Parenteral Fluids, Colostomy/Ileostomy, Respiratory Care, Tracheostomy Care, Suctioning, Tube Feeding
- Rehabilitative Services (physical therapy, speech pathology and audiology services, occupational therapy)

Due to the relatively low prevalence of these care needs/treatments, appropriate residents may be either under-represented or entirely omitted from the sample. Therefore, determine during the tour how many residents in the random selection fall into each of these care categories. Then, compare the number of such residents in the random selection with the total number of residents in the facility with each specified care need/treatment (based on either the resident census or other information provided by the facility).

Review no less than 25 percent of the residents in each of these special care needs/treatments categories. For example, if the facility has 10 residents with
decubitus ulcers, but only one of these residents is selected randomly, review two more residents with decubitis ulcers (25% of 10 equals 2.5, so review a total of 3). Or, if the facility has two residents who require tube feeding, neither of whom is in the random selection, review the care of at least one of these residents. This can be accomplished in the following manner:

Conduct in-depth reviews of the randomly selected residents and then perform limited reviews of additional residents as needed to cover the specified care categories. Such reviews are limited to the care and services related to the pertinent care areas only, e.g., catheters, restraints, or colostomy. Utilize those worksheets or portions of worksheets which are appropriate to the limited review. Refer to the Care Guidelines, as a resource document, when appropriate.

Always keep in mind that neither the random selection approach nor the review of residents within the specified care categories precludes investigation of other resident care situations that you believe might pose a serious threat to a resident’s health or safety. Add to the sample, as appropriate.

(e) Task 3—Tour of the Facility. (1) Purpose. Conduct the tour in order to:

• Develop an overall picture of the types and patterns of care delivery present within the facility;
• View the physical environment; and
• Ascertain whether randomly selected residents are communicative and willing to be interviewed.

(2) Protocol. You may tour the entire facility as a team or separately, as long as all areas of the facility are examined by at least one team member. Success of the latter approach, however, is largely dependent on open intra-team communication and the ability of each team member to identify situations for further review by the team. Access to all areas of the facility is required. The tour may be conducted with or without facility staff accompanying you, as you prefer. Facilities, however, vary in staff member availability. Record your notes on the Tour Notes Worksheet, Form HCFA-521.

Allow approximately three hours for the tour. Converse with residents, family members/significant others (if present), and staff, asking open-ended questions in order to confirm observations, obtain additional information, or corroborate information, (e.g., accidents, odors, apparent inappropriate dress, adequacy and appropriateness of activities). Converse sufficiently with residents selected for in-depth review to ascertain whether they are willing to be interviewed and are communicative. Observe staff interactions with other staff members as well as with residents for insight into matters such as resident rights and assignments of staff responsibilities.

Always knock and/or get permission before entering a room or interrupting privacy. If you wish to inspect a resident’s skin, observe a treatment procedure, or observe a resident who is exposed, courteously ask permission from the resident if she/he comprehends, or ask permission from the staff nurse if the resident cannot communicate. Do not do “hands-on” monitoring such as removal of dressings; ask staff to remove a dressing or handle a resident.

(3) Resident Needs. While touring, focus on the residents’ needs—physical, emotional, psychosocial, or spiritual—and whether those needs are being met. Refer to the following list as needed:

—Personal hygiene, grooming, and appropriate dress
—Position
—Assistive and other restorative devices
—Rehabilitation issues
—Functional limitations in ADL
—Functional limitations in gait, balance and coordination
—Hydration and nutritional status
—Resident rights
—Activity for time of day (appropriate or inappropriate)
—Emotional status
—Level of orientation
—Awareness of surroundings
—Behaviors
—Cleanliness of immediate environment (wheelchair, bed, bedside table, etc.)
—Odors
—Adequate clothing and care supplies as well as maintenance and cleanliness of same

(4) Review of the Physical Environment. As you tour each resident’s room and
auxiliary rooms, also examine them in connection with the physical environment requirements. You need not document physical environment on the Tour Notes Worksheet. Instead, you may note any negative findings directly on the Survey Report Form in the remarks section.

(5) Meeting With Resident Council Representatives. If a facility has a Resident Council, one or more surveyors meet with the representatives in a private area. Facility staff members do not attend unless specifically requested by the Council. Explain the purpose of the survey and briefly outline the steps in the survey process, i.e., entrance conference * * * exit conference. Indicate your interest in learning about the strengths of the facility in addition to any complaints or shortcomings. State that this meeting is one part of the information gathering; the findings have not yet been completed nor the conclusions formulated. Explain further, however, that the official survey findings are usually available within three months after the completion of the survey, and give the telephone number of the State agency office.

Use this meeting to ascertain strengths and/or problems, if any, from the consumer’s perspective, as well as to develop additional information about aspects of care and services gleaned during the tour that were possibly substandard.

Conduct the meeting in a manner that allows for comments about any aspect of the facility. (See the section on Interview Procedures.) Use open-ended questions such as:

- “What is best about this home?”
- “What is worst?”
- “What would you like to change?”

In order to get more detail, use questions such as:

- “Can you be more specific?”
- “Can you give me an example?”
- “What can anyone else tell me about this?”

If you wish to obtain information about a topic not raised by the residents, use an approach like the following:

- “Tell me what you think about the food/staff/cleanliness here.”
- “What would make it better?”
- “What don’t you like? What do you like?”

(6) Tour Summation and Focus of Remaining Survey Activity. When the tour is completed, review the resident census data provided by the facility. Determine if the care categories specified in the section on Resident Sample are sufficiently represented in the random selection, make adjustments as needed, and complete the listing of residents on the worksheet labeled “Residents Selected for In-depth Review”, Form HCFA-520.

Transcribe notes of a negative nature onto the SRF in the “Remarks” column under the appropriate rule. Findings from a later segment in the survey or gathered by another surveyor may combine to substantiate a deficiency. You need not check “met” or “not met” at this point in the survey. Discuss significant impressions/conclusions at the completion of each subsequent survey task, and transfer any negative findings onto the Survey Report Form in the Remarks section.

(f) Task 4—Observation/Interview/Medical Record Review (including drug regimen review). Perform the in-depth review of each individual in the resident sample in order to ascertain whether the facility is meeting resident needs. Evaluate specific indicators for each resident, utilizing the front and back of the “Observation/Interview/Record Review (OIRR)” worksheet, Form HCFA-524. You may prefer to perform the record review first, complete resident/staff/family observations and interviews, and finally, return to the record for any final unresolved issues. On the other hand, you may prefer to do the interviews first. Either method is acceptable. Whenever possible, however, complete one resident’s observation/interview/medical record review and document the OIRR before moving onto another resident. If because of the facility layout, it is more efficient to do more than one record review at a time, limit such record review to two or three residents so your familiarity with the particular resident and continuity of the OIRR are not compromised.

(1) Observation. Conduct observations concurrently with interviews of residents, family/significant others, and
discussions with direct care staff of the various disciplines involved. In multi-facility operations, whenever possible, observe staff that is regularly assigned to the facility in order to gain an understanding of the care and services usually provided. Maintain respect for resident privacy. Minimize disruption of the operations of the facility or impositions upon any resident as much as possible. Based upon your observations of the residents' needs, gather information about any of the following areas, as appropriate:

- Bowel and bladder training
- Catheter care
- Restraints
- Injections
- Parenteral fluids
- Tube feeding/gastrostomy
- Colostomy/ileostomy
- Respiratory therapy
- Tracheostomy care
- Suctioning

(2) Interviews. Interview each resident in private unless he/she independently requests that a facility staff member or other individual be present. Conduct the in-depth interview in a nonthreatening and noninvasive fashion so as to decrease anxiety and defensiveness. The open-ended approach described in the section on the Resident Council is also appropriate for the in-depth interview. While prolonged time expenditure is not usually a worthwhile use of resources or the resident's time, do allow time initially to establish rapport.

At each interview:

- Introduce yourself.
- Address the resident by name.
- Explain in simple terms the reason for your visit (e.g., to assure that the care and services are adequate and appropriate for each resident).
- Briefly outline the process—entrance conference, tour, interviews, observations, review of medical records, resident interviews, and exit conference.
- Mention that the selection of a particular resident for an interview is not meant to imply that his/her care is substandard or that the facility provides substandard care. Also mention that most of those interviewed are selected randomly.

- Assure that you will strive for anonymity for the resident and that the interview is used in addition to medical records, observations, discussions, etc., to capture an accurate picture of the treatment and care provided by the facility. Explain that the official findings of the survey are usually available to the public about three months after completion of the survey, but resident names are not given to the public.

- When residents experience difficulty expressing themselves:
  - Avoid pressuring residents to verbalize
  - Accept and respond to all communication
  - Ignore mistakes in word choice
  - Allow time for recollection of words
  - Encourage self-expression through any means available
- When interviewing residents with decreased receptive capacity:
  - Speak slowly and distinctly
  - Speak at conversational voice level
  - Sit within the resident's line of vision
- Listen to all resident information/allegations without judgment. Information gathered subsequently may substantiate or repudiate an allegation.

The length of the interview varies, depending on the condition and wishes of the resident and the amount of information supplied. Expect the average interview, however, to last approximately 15 minutes. Courteously terminate an interview whenever the resident is unable or unwilling to continue, or is too confused or disoriented to continue. Do, however, perform the other activities of this task (observation and record review). If, in spite of your conversing during the tour, you find that less than 40 percent of the residents in your sample are sufficiently alert and willing to be interviewed, try to select replacements so that a complete OIRR is performed for a group this size, if possible. There may be situations, however, where the resident population has a high percentage of confused individuals and this percentage is not achievable. Expect that the information from confused individuals can be, but is not necessarily, less
§ 488.110

reliable than that from more alert individuals.
Include the following areas in the interview of each resident in the sample:
Activities of daily living
Grooming/hygiene
Nutrition/dietary
Restorative/rehabilitation care and services
Activities
Social services
Resident rights
Refer to the Care Guidelines “evaluation factors” as a resource for possible elements to consider when focusing on particular aspects of care and resident needs.
Document information obtained from the interviews/observations on the OIRR Worksheet. Record in the “Notes” section any additional information you may need in connection with substandard care or services. Unless the resident specifically requests that he/she be identified, do not reveal the source of the information gleaned from the interview.
(3) Medical Record Review. The medical record review is a three-part process, which involves first reconciling the observation/interview findings with the record, then reconciling the record against itself, and lastly performing the drug regimen review.
Document your findings on the OIRR Worksheet, as appropriate, and summarize on the Survey Report Form the findings that are indicative of problematic or substandard care. Be alert for repeated similar instances of substandard care developing as the number of completed OIRR Worksheets increases.

NOTE: The problems related to a particular standard or condition could range from identical (e.g., meals not in accordance with dietary plan) to different but related (e.g., nursing services—lapse in care provided to residents with catheters, to residents with contractures, to residents needing assistance for personal hygiene and residents with improperly applied restraints).

(ii) Reconciling the observation/interview findings with the record. Determine if:
• An assessment has been performed.
• A plan with goals has been developed.
• The interventions have been carried out.
• The resident has been evaluated to determine the effectiveness of the interventions.

For example, if a resident has developed a decubitus ulcer while in the facility, record review can validate staff and resident interviews regarding the facility’s attempts at prevention. Use your own judgment; review as much of the record(s) as necessary to evaluate the care planning. Note that facilities need not establish specific areas in the record stating “Assessment,” “Plan,” “Intervention,” or “Evaluation” in order for the documentation to be considered adequate.

(ii) Reconciling the record with itself. Determine:
• If the resident has been properly assessed for all his/her needs.
• That normal and routine nursing practices such as periodic weights, temperatures, blood pressures, etc., are performed as required by the resident’s conditions.

(iii) Performing the drug regimen review. The purpose of the drug regimen review is to determine if the pharmacist has reviewed the drug regimen on a monthly basis. Follow the procedures in Part One of Appendix N, Surveyor Procedures for Pharmaceutical Service Requirements in Long-Term Care Facilities. Fill in the appropriate boxes on the top left hand corner of the reverse side of the OIRR Worksheet, Form HCFA-524. Appendix N lists many irregularities that can occur. Review at least six different indicators on each survey. However, the same six indicators need not be reviewed on every survey.

NOTE: If you detect irregularities and the documentation demonstrates that the pharmacist has notified the attending physician, do not cite a deficiency. Do, however, bring the irregularity to the attention of the medical director or other facility official, and note the official’s name and date of notification on the Survey Report Form.

(g) Task 5—Drug Pass Observation. The purpose of the drug pass observation is to observe the actual preparation and administration of medications to residents. With this approach, there is no doubt that the errors detected, if any, are errors in drug administration, not
documentation. Follow the procedure in Part Two of Appendix N, Surveyor Procedures for Pharmaceutical Service Requirements in Long-Term Care Facilities, and complete the Drug Pass Worksheet, Form HCFA-522. Be as neutral and unobtrusive as possible during the drug pass observation. Whenever possible, select one surveyor, who is a Registered Nurse or a pharmacist, to observe the drug pass of approximately 20 residents. In facilities where fewer than 20 residents are receiving medications, observe as many residents receiving medications as possible. Residents selected for the in-depth review need not be included in the group chosen for the drug pass; however, their whole or partial inclusion is acceptable. In order to get a balanced view of a facility’s practices, observe more than one person administering a drug pass, if feasible. This might involve observing the morning pass one day in Wing A, for example, and the morning pass the next day in Wing B.

Transfer findings noted on the “Drug Pass” worksheet to the SRF under the appropriate rule. If your team concludes that the facility’s medication error rate is 5 percent or more, cite the deficiency under Nursing Services/Administration of Drugs. Report the error rate under F209. If the deficiency is at the standard level, cite it in Nursing Services, rather than Pharmacy.

(h) Task 6—Dining Area and Eating Assistance Observation. The purpose of this task is to ascertain the extent to which the facility meets dietary needs, particularly for those who require eating assistance. This task also yields information about staff interaction with residents, promptness and appropriateness of assistance, adaptive equipment usage and availability, as well as appropriateness of dress and hygiene for meals.

For this task, use the worksheet entitled “Dining Area and Eating Assistance Observation” (Form HCFA-523). Observe two meals; for a balanced view, try to observe meals at different times of the day. For example, try to observe a breakfast and a dinner rather than two breakfasts. Give particular care to performing observations as unobtrusively as possible. Chatting with residents and sitting down nearby may help alleviate resident anxiety over the observation process.

Select a minimum of five residents for each meal observation and include residents who have their meals in their rooms. Residents selected for the in-depth review need not be included in the dining and eating assistance observation; however, their whole or partial inclusion is acceptable. Ascertain the extent to which the facility assesses, plans, and evaluates the nutritional care of residents and eating assistance needs by reviewing the sample of 10 or more residents. If you are unable to determine whether the facility meets the standards from the sample reviewed, expand the sample and focus on the specific area(s) in question, until you can formulate a conclusion about the extent of compliance. As with the other survey tasks, transfer the findings noted on the “Dining & Eating Assistance Observation” worksheet to the Survey Report Form.

(i) Task 7—Forming the Deficiency Statement. (1) General. The Survey Report Form contains information about all of the negative findings of the survey. Be sure to transfer to the Survey Report Form data from the tour, drug pass observation, dining area and eating assistance observation, as well as in-depth review of the sample of residents. Transfer only those findings which could possibly contribute to a determination that the facility is deficient in a certain area.

Meet as a group in a pre-exit conference to discuss the findings and make conclusions about the deficiencies, subject to information provided by facility officials that may further explain the situation. Review the summaries/conclusions from each task and decide whether any further information and/or documentation is necessary to substantiate a deficiency. As the facility for additional information for clarification about particular findings, if necessary. Always consider information provided by the facility. If the facility considers as acceptable, practices which you believe are not acceptable, ask the facility to backup its contention with suitable reference material or sources and submit them for your consideration.
(2) Analysis. Analyze the findings on the Survey Report Form for the degree of severity, frequency of occurrence and impact on delivery of care or quality of life. The threshold at which the frequency of occurrences amounts to a deficiency varies from situation to situation. One occurrence directly related to a life-threatening or fatal outcome can be cited as a deficiency. On the other hand, a few sporadic occurrences may have so slight an impact on delivery of care or quality of life that they do not warrant a deficiency citation. Review carefully all the information gathered. What may appear during observation as a pattern, may or may not be corroborated by records, staff, and residents. For example, six of the 32 residents in the sample are dressed in mismatched, poorly buttoned clothes. A few of the six are wearing slippers without socks. A few others are wearing worn clothes. Six occurrences might well be indicative of a pattern of substandard care. Close scrutiny of records, discussions with staff, and interviews reveal, however, that the six residents are participating in dressing retraining programs. Those residents who are without socks, chose to do so. The worn clothing items were also chosen—they are favorites.

Combinations of substandard care such as poor grooming of a number of residents, lack of ambulation of a number of residents, lack of attention to positioning, poor skin care, etc., can yield a deficiency in nursing services just as 10 out of 10 residents receiving substandard care for decubiti yields a deficiency.

(3) Deficiencies Alleged by Staff or Residents. If staff or residents allege deficiencies, but records, interviews, and observation fail to confirm the situation, it is unlikely that a deficiency exists. Care and services that are indeed confirmed by the survey to be in compliance with the regulatory requirements, but considered deficient by residents or staff, cannot be cited as deficient for certification purposes. On the other hand, if an allegation is of a very serious nature (e.g., resident abuse) and the tools of record review and observation are not effective because the problem is concealed, obtain as much information as possible or necessary to ascertain compliance, and cite accordingly. Residents, family, or former employees may be helpful for information gathering.

(4) Composing the Deficiency Statement. Write the deficiency statement in terms specific enough to allow a reasonably knowledgeable person to understand the aspect(s) of the requirement(s) that is (are) not met. Do not delve into the facility's policies and procedures to determine or speculate on the root cause of a deficiency, or sift through various alternatives in an effort to prescribe an acceptable remedy. Indicate the data prefix tag and regulatory citation, followed by a summary of the deficiency and supporting findings using resident identifiers, not resident names, as in the following example.

F102 SNF 405.1123(b).—Each resident has not had a physician’s visit at least once every 30 days for the first 90 days after admission. Resident ι1602 has not been seen by a physician since she was admitted 50 days ago. Her condition has deteriorated since that time (formulation of decubiti, infections).

When the data prefix tag does not repeat the regulations, also include a short phrase that describes the prefix tag (e.g., F117 decubitus ulcer care). List the data tags in numerical order, whenever possible.

(j) Task 8—Exit Conference. The purpose of the exit conference is to inform the facility of survey findings and to arrange for a plan of correction, if needed. Keep the tone of the exit conference consistent with the character of the survey process—inspection and enforcement. Tactful, business-like, professional presentation of the findings is of paramount importance. Recognize that the facility may wish to respond to various findings. Although deficiency statements continue to depend, in part, on surveyor professional judgment, support your conclusions with resident-specific examples (identifiers other than names) whenever you can do so without compromising confidentiality. Before formally citing deficiencies, discuss any allegations or findings that could not be substantiated during earlier tasks in the process. For example, if information is gathered that suggests a newly hired
Health Care Financing Administration, HHS § 488.110

R.N. is not currently licensed, ask the facility officials to present current licensure information for the nurse in question. Identify residents when the substandard care is readily observed or discerned through record review. Ensure that the facility improves the care provided to all affected residents, not only the identified residents. Make clear to the facility that during a follow-up visit the surveyors may review residents other than those with significant problems from the original sample, in order to see that the facility has corrected the problems overall. Do not disclose the source of information provided during interviews, unless the resident has specifically requested you to inform the facility of his/her comments or complaints. In accordance with your Agency’s policy, present the Statement of Deficiencies, form HCFA-2567, on site or after supervisory review, no later than 10 calendar days following the survey.

(k) Plan of Correction. Explain to the facility that your role is to identify care and services which are not consistent with the regulatory requirements, rather than to ascertain the root causes of deficiencies. Each facility is expected to review its own care delivery. Subsequent to the exit conference, each facility is required to submit a plan of correction that identifies necessary changes in operation that will have an impact on the correction of the cited deficiencies. In reviewing and accepting a proposed plan of correction, apply these criteria:

• Does the facility have a reasonable approach for correcting the deficiencies?
• Is there a high probability that the planned action will result in compliance?
• Is compliance expected timely?

Plans of correction specific to residents identified on the deficiency statement are acceptable only where the deficiency is determined to be unique to that resident and not indicative of a possible systemic problem. For example, as a result of an aide being absent, two residents are not ambulated three times that day as called for in their care plans. A plan of correction that says “Ambulate John Jones and Mary Smith three times per day.” is not acceptable. An acceptable plan of correction would explain changes made to the facility’s staffing and scheduling in order to guarantee that staff is available to provide all necessary services for all residents.

Acceptance of the plan of correction does not absolve the facility of the responsibility for compliance should the implementation not result in correction and compliance. Acceptance indicates the State agency’s acknowledgement that the facility indicated a willingness and ability to make corrections adequately and timely.

Allow the facility up to 10 days to prepare and submit the plan of correction to the State agency, however, follow your SA policy if the timeframe is shorter. Retain the various survey worksheets as well as the Survey Report Form at the State agency. Forward the deficiency statement to the HCFA regional office.

(l) Follow-up Surveys. The purpose of the follow-up survey is to re-evaluate the specific types of care or care delivery patterns that were cited as deficient during the original survey. Ascertain the corrective status of all deficiencies cited on the HCFA-2567. Because this survey process focuses on the actual provision of care and services, revisits are almost always necessary to ascertain whether the deficiencies have indeed been corrected. The nature of the deficiencies dictates the scope of the follow-up visit. Use as many tasks or portions of the Survey Report Form(s) as needed to ascertain compliance status. For example, you need not perform another drug pass if no drug related deficiencies were cited on the initial survey. Similarly, you need not repeat the dining area and eating assistance observations if no related problems were identified. All or some of the aspects of the observation/interview/medical record review, however, are likely to be appropriate for the follow-up survey.

When selecting the resident sample for the follow-up, determine the sample size using the same formula as used earlier in the survey, with the following exceptions:

• The maximum sample size is 30 residents, rather than 50.
• The minimum sample size of 10 residents does not apply if only one care category was cited as deficient and the total number of residents in the facility in that category was less than 10 (e.g., deficiency cited under catheter care and only five residents have catheters).

Include in the sample those residents who, in your judgment, are appropriate for reviewing vis-a-vis the cited substandard care. If possible, include some residents identified as receiving substandard care during the initial survey. If after completing the follow-up activities you determine that the cited deficiencies were not corrected, initiate adverse action procedures, as appropriate.

(m) Role of Surveyor. The survey and certification process is intended to determine whether providers and suppliers meet program participation requirements. The primary role of the surveyor, then, is to assess the quality of care and services and to relate those findings to statutory and regulatory requirements for program participation.

When you find substandard care or services in the course of a survey, carefully document your findings. Explain the deficiency in sufficient detail so that the facility officials understand your rationale. If the cause of the deficiency is obvious, share the information with the provider. For example, if you cite a deficiency for restraints (F118), indicate that restraints were applied backwards on residents 1621, 1634, 1646, etc.

In those instances where the cause is not obvious, do not delve into the facility’s policies and procedures to determine the root cause of any deficiency. Do not recommend or prescribe an acceptable remedy. The provider is responsible for deciding on and implementing the action(s) necessary for achieving compliance. For the restraint situation in the example above, you would not ascertain whether the improper application was due to improper training or lack of training, nor would you attempt to identify the staff member who applied the restraints. It is the provider’s responsibility to make the necessary changes or corrections to ensure that the restraints are applied properly.

A secondary role for the surveyor is to provide general consultation to the provider/consumer community. This includes meeting with provider/consumer associations and other groups as well as participating in seminars. It also includes informational activities, whereby you respond to oral or written inquiries about required outcomes in care and services.

(n) Confidentiality and Respect for Resident Privacy. Conduct the survey in a manner that allows for the greatest degree of confidentiality for residents, particularly regarding the information gathered during the in-depth interviews. When recording observations about care and resident conditions, protect the privacy of all residents. Use a code such as resident identifier number rather than names on worksheets whenever possible. Never use a resident’s name on the Deficiency Statement, Form HCFA-2567. Block out resident names, if any, from any document that is disclosed to the facility, individual or organization.

When communicating to the facility about substandard care, fully identify the resident(s) by name if the situation was identified through observation or record review. Improperly applied restraints, expired medication, cold food, gloves not worn for a sterile procedure, and diet inconsistent with order, are examples of problems which can be identified to the facility by resident name. Information about injuries due to broken equipment, prolonged use of restraints, and opened mail is less likely to be obtained through observation or record review. Do not reveal the source of information unless actually observed, discovered in the record review, or requested by the resident or family.

(o) Team Composition. Whenever possible, use the following survey team model:

**SNF/ICF Survey Team Model**

In facilities with 200 beds or less, the team size may range from 2 to 4 members. If the team size is:

- 2 members: The team has at least one RN plus another RN or a dietitian or a pharmacist.
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• 3-4 member: In addition to the composition described above, the team has one or two members of any discipline such as a social worker, sanitarian, etc.

If the facility has over 200 beds and the survey will last more than 2 days, the team size may be greater than 4 members. Select additional disciplines as appropriate to the facility's compliance history.

Average onsite time per survey: 60 person hours (Number of surveyors multiplied by the number of hours on site)

Preferably, team members have gerontological training and experience. Any member may serve as the team leader, consistent with State agency procedures. In followup surveys, select disciplines based on major areas of correction. Include a social worker, for example, if the survey revealed major psychosocial problems. This model does not consider integrated survey and Inspection of Care review teams, which typically would be larger.

(p) Type of Facility—Application of SNF or ICF Regulations. Apply the regulations to the various types of facilities in the following manner:

• Freestanding Skilled Nursing Facility (SNF)
• Freestanding Intermediate Care Facility (ICF)
• SNF Distinct Part of a Hospital
• ICF Distinct Part of a Hospital
• Dually Certified SNF/ICF
• Freestanding SNF with ICF Distinct Part (Regardless of the proportion of SNF and ICF beds, the facility type is determined by the higher level of care. Therefore, LTC facilities with distinct parts are defined as SNFs with ICF distinct parts.)

(q) Use of Part A and Part B of the Survey Report. (1) Use of Part A (HCFA-525).—Use Part A for initial certification surveys only, except under the following circumstances:

• When a terminated facility requests program participation 60 days or more after termination. Treat this situation as a request for initial certification and complete Part A of the survey report in addition to Part B.
• If an ICF with a favorable compliance history requests to convert a number of beds to SNF level, complete both Part A and Part B for compliance with the SNF requirements. If distinct part status is at issue, also examine whether it meets the criteria for certification as a distinct part.

(i) Addendum for Outpatient Physical Therapy (OPT) or Speech Pathology Services. Use the Outpatient Physical Therapy—Speech Pathology SRF (HCFA-1893) as an addendum to Part A.

(ii) Resurvey of Participating Facilities. Do not use Part A for resurveys of participating SNFs and ICFs. A determination of compliance, based on documented examination of the written policies and procedures and other pertinent documents during the initial survey, establishes the facility's compliance status with Part A requirements. This does not preclude citing deficiencies if they pertain to administrative or structural requirements from Part A that are uncovered incidental to a Part B survey. As an assurance measure, however, each facility at the time of recertification must complete an affidavit (on the HCFA-1516) attesting that no substantive changes have occurred that would affect compliance. Each facility must also agree to notify the State agency immediately of any upcoming changes in its organization or management which may affect its compliance status. If a new administrator is unable to complete the affidavit, proceed with the survey using the Part B form and worksheets; do not use the Part A form. The survey cannot be considered complete, however, until the affidavit is signed. If the facility fails to complete the affidavit, it cannot participate in the program.

(iii) Substantial Changes in a Facility’s Organization and Management. If you receive such information, review the changes to ensure compliance with the regulations. Request copies of the appropriate documents (e.g., written policies and procedures, personnel qualifications, or agreements) if they were
§ 488.115

not submitted. If the changes have made continued compliance seem doubtful, determine through a Part B survey whether deficiencies have resulted. Cite any deficiencies on the HCFA-2567 and follow the usual procedures.

(2) Use of Part B (HCFA-519). Use Part B and the worksheets for all types of SNF and ICF surveys—initials, recertifications, followup, complaints, etc. The worksheets are:

• HCFA-520—Residents Selected for In-depth Review
• HCFA-521—Tour Notes Worksheet
• HCFA-522—Drug Pass Worksheet
• HCFA-523—Dining Area and Eating Assistance Worksheet
• HCFA-524—Observation/Interview/Record Review Worksheet

For complaint investigations, perform a full or partial Part B survey based on the extent of the allegations. If the complaint alleges substandard care in a general fashion or in a variety of services and care areas, perform several tasks or a full Part B survey, as needed. If the complaint is of a more specific nature, such as an allegation of improper medications, perform an appropriate partial Part B survey, such as a drug pass review and a review of selected medical records.
§ 488.115 Care guidelines.

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<thead>
<tr>
<th>SURVEY AREA</th>
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<tr>
<td>Resident Rights</td>
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<tr>
<td>FG 3</td>
<td>SNF 405.1123(k)(1)</td>
<td>ICF 442.311(a)</td>
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<td>FG 4</td>
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<tr>
<td>A. Information*</td>
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<td>FG 5</td>
<td>SNF 405.1123(h)(1)</td>
<td>ICF 442.311(a)(2)</td>
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<td>FG 6</td>
<td>SNF 405.1123(k)(1)</td>
<td>ICF 442.311(a)(3)</td>
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<td>FG 7</td>
<td>SNF 405.1123(h)(2)</td>
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<tr>
<td>FG 8</td>
<td>SNF 405.1123(h)(3)</td>
<td>ICF 442.311(a)(5)</td>
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**INTENT**

To assure that the resident maintains, in so far as possible, those personal rights that are a part of normal adult life, and including the right to personal dignity.

*Information concerning incompetent residents is given in L. Delegation of Rights and Responsibilities.
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<td>F58</td>
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<tr>
<td>SDF 485.1121(k)(2)</td>
<td>4. Resident informed in writing of changes in services and charges for services.</td>
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<td>ICF 482.331(a)(4)</td>
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<td>F59</td>
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<tr>
<td>SDF 485.1121(k)(2)</td>
<td>5. Information to resident of services not covered by Medicare or Medicaid and not covered in the basic rate.</td>
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<tr>
<td>ICF 482.331(a)(4)</td>
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**Ask Resident:**
- If there are changes in services or costs, does someone explain these?

**Ask Administrative Staff:**
- How do residents learn what is expected of them?
- How do they learn about any changes in the facility's procedures and/or costs?
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<tr>
<td>B. Medical Condition &amp; Treatment</td>
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<td>Ask Resident:</td>
<td>If the resident has not been informed of his/her medical condition, physician notes should document that the resident was not informed because it was medically contraindicated. Do care plans or other documentation reflect resident participation in care planning? If resident states he/she has refused treatment or medication, does documentation indicate adherence to violation of resident rights? Review records of residents identified as participating in a clinical research study. Are informed consent forms signed? Do these signed forms list all known risks for the resident? All needed informed consent statements are present and properly signed.</td>
<td>Unless there is documentation that the resident's medical condition should not be discussed with him/her, resident interview/record review should indicate that the resident and physician have discussed his/her medical condition. If you cannot confirm that this has occurred, interview staff to get further clarification. Almost all residents who are able to participate to some extent in their care planning do so. You should find evidence of this for the majority of the residents (e.g., care planning interview, nurses' notes, social worker progress notes).</td>
<td>Patient Care Management § 482.112(b)(d) 442.310 442.341</td>
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Residents do have the right to refuse medication or other treatment, but you would expect that the facility would discuss the implications of this refusal with the resident and possibly do some "gentle persuasion."
LONG TERM CARE SURVEY

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<tr>
<td>f60-64 (cont'd)</td>
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<td>However, except in an emergency situation force should never be used to compel a resident to accept medication or treatment.</td>
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<td>Deceit is also a violation of resident rights, except in the case of therapeutically indicated placebos ordered by the physician.</td>
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<td>Any resident participating in research studies should fully understand the implication of the study.</td>
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<td>The facility is not in compliance with the resident rights regulation if the resident consents to participate in a clinical study without full knowledge of the study. (Record review only as other nonclinical studies may not require informed consent).</td>
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<td>C. Transfer and Discharge</td>
<td>Look for residents that may be inappropriately placed physically— an alert resident roaming with a confused, noisy resident; very ill resident placed far from the nurses station; residents not compatible with each other. (e.g., different life-styles, habits, etc.).</td>
<td>\textbf{Ask Resident:} - How well do you get along with your roommate? - Have you ever been moved from one room to another? If yes, why? - How were you involved in the decision to move? - How much time was there between the time they told you you were to be moved, and when you were moved? - Have you asked for your room to be changed? \textbf{Ask Direct Care and Other Staff:} - What are some of the reasons residents rooms are changed? - What are some of the reasons for discharge of residents or transfer to a hospital or LTC facility? - How are residents involved in the decision to move? - If a resident requests a room change, how is this handled? - When a resident requests a room change are the following areas of consideration presented and discussed:</td>
<td>Nursing, physician, and/or social service progress notes should indicate reason for transfer and discussion with resident and/or family/guardian. If staff interviews give you cause to feel that transfers and discharges may be in violation of these regulations, review a sample of closed records for transfer information on how it was handled.</td>
<td>To be in compliance with transfer and discharge regulations the facility must be able to confirm that all discharges/transfers were for medical or resident welfare reasons, or non-payment. Welfare reasons include physical, emotional, and social issues. Transfers and discharges made solely for the convenience of the facility are unacceptable. Relocation to accommodate contagious or other disorders requiring isolation procedures are not for the convenience of the facility.</td>
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<tr>
<td>F65-68 SNF 485.112(c)(4) ICF 482.311(c)</td>
<td>\textbf{Status Change} 482.112(f) \textbf{Medical Records} 485.112(c)(6) 482.318(c)(4) \textbf{Transfer Agreement} 485.113(e)(2) 482.317(b)(1)(2)</td>
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| D. Exercising Rights | Do residents appear comfortable when speaking to the surveyors as opposed to being afraid that someone may see them or overhear their conversation? | **Ask Resident:**  
- Do you belong to, or have representation on the resident council?  
- Are you informed of changes in the facility that will affect you?  
- Are you given a chance to express views on these changes prior to their implementation?  
- Does the facility assist in arranging for you to vote either at the polls or via absentee ballot?  
- Are you assisted in obtaining legal or Social Services if needed?  
- Do you feel comfortable in expressing yourself freely or are you concerned about retaliation?  
- Is staff/administration responsive to complaints? Do you know who to complain to?  
**Ask Staff:**  
- What arrangements are made for residents to vote?  
- How do you handle it if someone needs a lawyer or other service that you don’t provide? | Review resident council documentation, as available, to determine level of activity. | Compliance determinations will be made based primarily on resident/staff interviews and the correlation of interview information with documentation in the Medical record. If residents ask, they should be allowed to speak to the surveyor without facility personnel being present. However the resident has the right to have a third party of their choosing present during an interview. | Social Services 485.113(b) 442.364 |
|             |             |              |               |                   |                 |

Social Services
485.113(b)  
442.364
## Long Term Care Survey

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<tr>
<th>Survey Area</th>
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<th>Record Review</th>
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</table>
| F72-78      | Ask Residents:  
- Are you able to take care of your own financial affairs?  
- Does the facility keep some money for you that you can have when you request it?  
- When you ask for this money, how quickly do you get it?  
- Do you know the amount of money you have available at this time?  
- If the facility pays bills for you do they periodically provide an itemized listing of the transactions they have made?  
- When did you receive the last itemized statement?  
- Are you comfortable that your funds are taken care of correctly?  
- If you deposit money or valuables with the facility, do you receive a receipt for this deposit?  
- Are you or your family able to review your financial records when you request to do so?  
- Have you ever had money or anything else stolen? If so, what was done about it? |  | |  |  |  |
| SNF 485.1121(k)(6) | A copy of the statement should be in the resident's financial record and given to the resident at least quarterly. Receipts, account logs showing deposits/withdrawals, authorizations/reasons for withdrawals, and interest earned should be reviewed. If resident indicates there may be a problem, an in-depth interview should be conducted.  |  |  |  |  |
| ICF 485.1111(e) | Resident records indicate separate financial records from facility records. |  |  |  |  |
| 442.310(e) | Residents should have reasonable access to their funds (may not be available at 2 a.m.) and should have at least a quarterly accounting of their funds.  |  |  | Social Services 485.1130(a) |  |
### LONG TERM CARE SURVEY

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| F. Freedom from Abuse and   | - How many residents are physically restrained?  
- What type or restraints are used?  
- Are they applied correctly?  
- What is the apparent physical/mental condition of those residents restrained?  
- Do you observe the release of restraints every 2 hours and the provision of at least 10 minutes exercise for the resident?  
- Do staff respond to request for water, assistance to bathroom, etc., from a resident who is restrained?  
- What is the interval between request and response?  | Ask Resident:  
- Why are you wearing this?  
- How often is this worn?  
- Do you know what would happen if it were removed?  
- How often is it removed?  
- What is done for you when the restraint is removed?  
- For nonrestrained resident—  
- Have you ever been restrained?  
- For what reason?  
- What explanation was given for the restraint?  
- Do you ever feel that you receive medication when you don’t need it?  | Look for a physician’s order for the restraint.  
Review nurses’, physicians’ progress notes re: reason for restraints and resident reaction to them.  
Also any alternative methods tried.  
What time of day are restraints most often applied?  
Review schedule of releasing restraints.  | There must be a physician’s order for all restraints, including “safety devices” which are defined in some State laws.  
Progress notes should show evidence that methods other than restraints were initially used to protect the resident from injury, and that restraints were used only when other methods were not adequate  
If used in an “emergency” the reason for use must be documented and show that:  
a. Its use was necessary to protect the resident from injury.  
b. Its use was necessary to protect others from injury.  
If appropriate are the Social Service or activities departments involved in providing different directions for resident attention?  | Nursing Services  
483.1124(c)(5)  
Rehabilitation  
483.1124(e)  
Patient Care  
Management  
483.1124(d) |
### Observation

- How often are residents restrained by staff?
- Observe effects on residents. Do you see what may be signs of overmedication?
- How often is this observed?
- Residency should be free from mental and physical abuse.
- Observe interaction of staff and residents for any signs of harassment, humiliation or threats.
- Do residents appear comfortable with staff?
- Look for numbers of residents with bruises or other injuries (skin of the elderly bruises easily, do not automatically assume abuse or injury).
- Observe resident to resident interactions and staff response to any physical or mental abuse of one resident to another.

### Interviewing

- Ask staff:
  - What is the facility policy regarding restraint?
  - What is considered an "emergency" need for restraint?
  - What is the most common reason for use of restraint?
  - Do you try any alternative measures before using restraints?
  - What information do you give the physician to help him make the decision to order restraints?
  - What do you routinely do for the resident when you periodically release the restraints?
  - Does use of restraints increase on evenings or nights when there are fewer staff members?
  - Have you had any accidents or incidents in the last year while residents were restrained?
  - How do you define the difference between a "safety device," a "device," and a "restraint"?
  - How do your policies differ in regard to "safety devices" and restraints?

### Record Review

- Who authorizes the use of restraints in an emergency?
- Do progress notes indicate that a professional staff member authorized the use of "emergency" restraints?
- There should be documentation that the use of "emergency" restraint has been promptly reported to the residents physician.
- Review incident and accident reports to identify any problematic trends.
- Does the drug regimen review indicate appropriate use of psychotropic drugs?
- Are there resident complaints documented?
- What is the resolution of these complaints?

### Evaluation Factors

- The restraint must be applied correctly.
- If the use of restraints increased during evening and night hours, review progress notes, nursing notes, and staffing to make a determination as to whether the restraints are justified or if they are for staff convenience.
- Care plans should plan not only for care while the resident is restrained but should show effort to find alternative treatments to restraints, or there should be documentation in the medical record that no alternative is appropriate.
- An appropriate drug regimen review should be conducted on the resident.
- Your observations should show interaction between residents and staff to be, except in unusual situations, free from tension and hostility.
- Staff should step into situation where one resident may be abusing another.
### Long Term Care Survey

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<tr>
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<tbody>
<tr>
<td>F79-83 (cont'd)</td>
<td>- Observe for evidence of resident neglect. Residents left in urine/feces without cleaning.</td>
<td>Ask Resident: &lt;br&gt;- Do you feel safe in the facility? &lt;br&gt;- Do you ever feel intimidated, harassed, or otherwise abused? &lt;br&gt;- How are confused residents treated? &lt;br&gt;- Is anyone ever hit or treated roughly? &lt;br&gt;- Do you feel as if you are treated with respect /dignity? &lt;br&gt;- Is the staff/administration responsive to complaints? &lt;br&gt;- Do you know who to complain to?</td>
<td></td>
<td>Resident should feel free to voice complaints. If no complaints are noted in records or on record review, why not? &lt;br&gt;Residents should seem comfortable in relating how they are treated.</td>
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<td>G. Privacy</td>
<td>- Observe interactions between staff and residents for indications of respect, consideration, dignity, and individuality. - How do staff members enter a resident's room or go behind a privacy curtain? - Are privacy curtains used or do doors shut when personal care needs or treatments are rendered? - Are there areas for residents to be alone or meet in private with visitors?</td>
<td>Ask Resident: - Do you feel that you are treated as a worthwhile, adult individual? - When you are being cared for, are you comfortable? - What is the degree of privacy and respect you receive? - Do you feel comfortable that if the door to your room is closed staff will knock or otherwise make their presence known before entry? - Do you have a private place to make telephone calls? - Can you see your record if you ask? - Has any information about your condition been given to someone outside of the facility without your permission?</td>
<td>Review progress notes for indications that staff see resident as an individual, i.e., resident eats breakfast in bed because he/she enjoys it. Signed consent for release of information. Do maintenance of and content of medical records indicate that confidentiality is practiced.</td>
<td>Observations and interviews will give you information to determine if residents are respected and treated as individuals. In privacy available--e.g., access to a private place to meet or make phone calls, ability to shut door when having visitors, etc. Medical records should not be left where unauthorized personnel can read them and there should be identification codes needed to access computerized records. Married residents should be sharing rooms if they desire to do so unless there are appropriate contradictions.</td>
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## Long Term Care Survey

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</table>
| F84-89 (cont'd) | - Are medical records kept in their assigned spots not carelessly left for unauthorized persons to view?  
- Are married residents sharing rooms?  
- Observe for negative attitudes toward aging-infantilization and patronizing of residents.  
- If residents undress in public area, how does staff handle this?  
- Listen to staff conversation in public places (elevator, lobby). Are resident issues being discussed? | For Married Residents:  
- When your husband/wife visits can you shut your door and be assured of privacy?  
- Can you ask that you not be disturbed and have that request respected?  
Ask Staff:  
- What is done to assure that each resident maintains his/her dignity and individuality?  
- How are medical records kept secure? Who has access?  
- Do you have married couples here?  
- Do they share rooms?  
- If not, why?  
- What arrangements do you make for spouses or significant others to visit?  
- Do you allow their door to be closed?  
- Can you adhere to a request that they not be disturbed?  
- How are residents' medical records and conditions kept confidential? | | | |

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<td>M. Work</td>
<td>Are residents doing any type of work such as picking up dirty trays, pushing laundry hampers, etc.?</td>
<td>Ask Resident: Are you ever asked to help out in the facility such as pick up dirty trays or stamp mail?</td>
<td>If residents are performing services for the facility, is that included in their care plan with specific therapeutic goals defined?</td>
<td>Are results documented in progress notes?</td>
<td>Services performed by a resident should be part of the resident's plan of care and should be done only if the resident is in full agreement.</td>
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<td>- What about clerical work?</td>
<td>- If yes, do you do this?</td>
<td>- If appropriate does the family concur?</td>
<td>What service (activities, nursing, etc.) is responsible for planning reevaluating and adjusting work activity?</td>
<td>Service rewards are specifically identified and not obtained using the residents own funds.</td>
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<td></td>
<td>Ask Staff: Are residents asked to help with facility staff if you are shorthanded?</td>
<td>- What is their reaction?</td>
<td>- Look for physician's orders for approval or disapproval of work activity or restrictions on this activity. Look for evidence that the resident is given opportunities to refuse to do the work.</td>
<td>- What useful work is available for residents who want/need to be usefully employed?</td>
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<td>- What useful work is available for residents who want/need to be usefully employed?</td>
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<tr>
<td>I. Freedom of Association and Correspondence</td>
<td>- Are there areas in the facility—e.g., small lounges, etc., where residents can and do meet privately?</td>
<td>Ask Residents:  - Can you have visits from anyone?  - Can you find a private place to visit?  - Do you receive your mail opened or unopened?  - Are there telephones you have access to?  - Does the staff or volunteers assist you in reading or mailing mail?  - How timely is your mail delivered?  - How do you receive incoming calls? Ask Staff:  - Where do residents go when they want privacy?  - What telephones are available to residents?  - What is the facility visiting policy?</td>
<td>Physician orders and care plans for indications of restrictions on visitors and/or receiving and sending mail.</td>
<td>All residents may have access to and maintain contact with the community and members of that community have access to them. Subject to reasonable scheduling restrictions, residents may receive visits from anyone they wish. A particular visitor may be restricted by the facility for one of the following reasons:  - The resident refuses to see the visitor.  - The resident’s physician documents specific reasons why such a visit would be harmful to the resident’s health.  - The visitor’s behavior is unreasonably disruptive of the functioning of the facility (reasons are documented and kept on file). Decisions to restrict a visitor are reviewed and reevaluated each time the resident’s plan of care and medical orders are reviewed by the physician and nursing staff or at the resident’s request.</td>
<td>Resident Rights 420.1121(b)(6) 442.3111(g)</td>
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<td>F91-92 (cont'd)</td>
<td>Do the available telephones accommodate the physically handicapped (e.g., wheelchair bound, hearing impaired, etc.).</td>
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<td>Space is provided for residents to receive visitors in reasonable comfort and privacy. Telephones, consistent with ANSI standards 40.1134(c), are made available and accessible for residents to make and receive calls with privacy. Residents who need help are assisted in using the phone. The fact that telephone communication is possible, as well as any restrictions, is made known to residents. Arrangements are made to provide assistance to residents who require help in reading or sending mail.</td>
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<td>2. Activities</td>
<td>- What planned activities are occurring?</td>
<td>Ask Residents: <em>What do you like to do?</em> <em>What did you do yesterday? (compare answers)</em> <em>Is participation in activities optional?</em> <em>Are you encouraged to participate?</em> <em>Is pressure exerted on you to attend specific activities?</em> <em>Which ones? (Surveyors should be aware of special encouragement—&quot;gentle persuasion&quot;, which might be important for the depressed or withdrawn resident.)</em> - Are residents notified of community activities? - Are arrangements made for transportation, etc. so that residents can participate? - Can residents go to religious services if they wish? - What opportunities are you given to make choices in your life within the facility? <em>(e.g., are all residents &quot;put to bed&quot; at the same time)?</em></td>
<td>Care plans or other documentation should indicate resident preferences for both facility and non-facility planned activities.</td>
<td>Compliance with this element is determined by evidence that residents are given the opportunity to participate in available activities they choose unless medically contraindicated.</td>
<td>Patient Activities 405.113(c) 442.305(a)(c)</td>
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<td>- What unplanned activities are occurring—individual, 2 or 3 persons or a larger group.</td>
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<td>- If there is a facility chapel, is it open?</td>
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<td>- Are activities posted at wheelchair level and kept up to date?</td>
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<td>- Are residents lined up in front of a T.V. in a common room for hours?</td>
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<td>- Are activities offered during the evening and on weekends?</td>
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Residents must not be forced to participate against their wishes.
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<tr>
<td>K. Personal Possessions</td>
<td>Are residents wearing their own clothing or facility nightgowns, robes, etc.?</td>
<td>Ask Residents: - What clothing and personal belongings can you have? - Is there a place that you can secure any valuables that you may not want to keep in your room?</td>
<td>Admission notes on personal property inventory (e.g., the record should indicate a list of any personal property secured by the facility). The record should indicate how personal clothing will be laundered.</td>
<td>Residents are permitted to keep reasonable amounts of personal clothing and possessions for their use while in the facility and such personal property is kept in a safe location which is convenient to the patient. The amount that is reasonable will be dependent on space available in the facility. Patients are advised, prior to or at admission, of the kinds and amounts of clothing and possessions permitted for personal use, and whether the facility will accept responsibility for maintaining these items (e.g., cleaning and laundry). Any personal clothing or possessions retained by the facility for the patient during his stay is identified. The facility is responsible for secure storage of such items and they are returned to the patient promptly upon request or upon discharge from the facility.</td>
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<td>L. Delegation of Rights and</td>
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<td>Responsibilities</td>
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<td>SNF 405.112(k)</td>
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<td>ICF 402.312</td>
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<td>Ask Administrative Staff:</td>
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<td>- When do you have relatives make decisions for residents? i.e., how do you decide when the resident isn't capable of making decisions himself?</td>
<td>Review physician progress notes—incapability must be documented.</td>
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<td>- Have any legal steps been taken?</td>
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<td>Ask Resident and/or</td>
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<td>Guardian:</td>
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<td>- Do you feel that you are given all pertinent information?</td>
<td>The fact that a resident has been judged incompetent, is medically incapable of understanding, or exhibits a communication barrier, does not absolve the facility from advising the resident of their rights to the extent the patient is able to understand them. If the resident is incapable of understanding their rights, the facility advises the guardian or sponsor and acquires a statement indicating an understanding of resident's rights.</td>
<td>Resident Rights 405.112(k)(1) (a) 402.311(a)</td>
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<td>- What opportunities do you have to make decisions regarding clothing, meals, bathing schedules, etc.?</td>
<td>Are pertinent consents/documents signed by appointed guardian?</td>
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<td>- For guardian: are you notified/informed in a timely manner as appropriate?</td>
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## LONG TERM CARE SURVEY

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<tr>
<td>F100 Staff</td>
<td>How do staff relate to residents? Does the facility reflect adaptations for the elderly, i.e., information given in large print, floors covered with materials that allow for ease of movement with walkers, wheelchairs, etc.?</td>
<td>Ask Residents - Does staff know how to take care of you? - What things do they do to help you accommodate your (poor vision, unsteady walking, arthritis, etc.)?</td>
<td>Care plans reflect staff's knowledge of the problems and needs of the residents and special adaptations that are needed.</td>
<td>Facility staff adjusts care to needs/problems of resident.</td>
<td>ECF 442.313 (a) (b) (c) (d) (e) (f)</td>
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<td>Staff is knowledgeable concerning facility policies and procedures.</td>
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<td>Staff practices correct techniques, i.e., infection control rehabilitation, nursing techniques, etc.</td>
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<td>Staff interacts and treats residents in a kind, caring way.</td>
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<td>F102 Facility staff practices proper technique in providing care to the aged, ill and disabled.</td>
<td>Is resident care given using accepted professional standards?</td>
<td>Ask Staff - What, if any, training have you had here to learn about unique problems and needs of the aged? - What training have you had during the last 12 months? - How have you learned about facility policies and procedures? - Does the facility ask your needs when they develop a training program? - In what areas would you like to have training?</td>
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<td>Are housekeeping staff courteous and responsive to resident needs?</td>
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### CROSS REFERENCE
- **Residents Rights**
  - SNF 405.1121(a)
  - ECF 442.313
- **Infection Control**
  - SNF 405.1135(a)(b)(c)
  - ECF 442.327(b)
- **Physical Environment**
  - SNF 405.1134(a)
  - ECF 442.315(b)(c)
  - ECF 442.326(a)(c)
- **Nursing Services**
  - SNF 405.1124(a)(c)(e)
  - ECF 442.330(a)(e)
- **Social Services**
  - SNF 405.1138(a)

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**Health Care Financing Administration, HHS**

$\text{\$88.115}$

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<td>F102 (cont'd) and safety, accident prevention, confidentiality of resident information, and preservation of resident dignity including protection of privacy and personal and property rights.</td>
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**INTENT**

To assure that facility provides ongoing training to staff so that they will be knowledgeable in current practices, use proper techniques, and interact with residents in a kind, caring way.
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| Status Change Notifications | - Clean  
- Well groomed  
- Well adjusted  
- Casts  
- Bruises  
- Decubitus Ulcer  
- Multiple sites of edema  
- Aberrant behavior, e.g., abusive, disruptive, not reasonable, etc. | Ask Resident:  
- Have you been injured since you have been in the facility?  
- If you are injured or become ill, is your physician called?  
- Are your relatives notified?  
- Do you know who is notified if administrative changes such as changes in charges, billings, etc. occur?  
- Are you notified if a resident is injured or has a change in condition?  
- Who do you notify if a resident is injured or has a change in condition?  
- When would they be notified?  
- Does the facility have a policy regarding how soon a relative or responsible party would be notified?  
- Do you notify them of actual changes in resident condition and also if resident's condition is getting progressively worse? | - Progress note should document injury/change in condition plus notification of physician and appropriate family member/guardian.  
- Changes in charges should be documented. Ask facility where this is located.  
- Review accident and incident reports for in-depth sample. | - All injuries and changes in condition must be documented. The resident's physician and family must be notified of significant changes. This should be documented, but this notification should be confirmed by the resident if possible. | Resident Supervision by Physician 485.1123(b)(3)  
Neighborhood Services 485.1123(c)
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| F106        | Ask Resident:  
- Have you ever been or do you know if others have been transferred or discharged without discussing it with you first?  
- Nursing, physician and social work progress notes should be reviewed for evidence of discussion of transfer/discharge with resident or other designated person.  
- Except in an emergency, all transfers or discharges are first discussed with the resident or next of kin as evidenced by documentation in the medical record or confirmed by asking resident. | - | - | - | - |

**INTENT**

To assure that:
- the resident receives proper treatment in the event of an accident or change of condition;
- resident and/or next of kin or responsible party is aware in advance of any changes;
- resident is not discharged to gain a higher source payment for that bed or facility convenience.
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<td>Physician's Services</td>
<td>Ask All Staff: - Interview nursing staff to determine if they receive transfer information and admission orders on day of admission. - Ask Administrator and Director of Nursing to explain procedure if a resident arrives without sufficient medical information and/or orders.</td>
<td>Review records of residents selected for in-depth review to ascertain that: - There is a referral form from the transferring facility that was received in advance of admission or on date of admission that includes current medical findings, diagnosis and orders from a physician for the immediate care of the residents. - If the medical orders were not obtained from the emergency care physician, there are temporary orders from the admitting physician. - Information on the rehabilitation potential (prognosis) of the resident and summary of the course of treatment followed in the transferring facility. - The summary of treatment should include discharge summaries from therapies or special services when appropriate. For residents admitted directly from the</td>
<td>Examine medical records of the residents selected for in-depth review to determine if date of orders, medical data and other required information is the date of admission or within 48 hours of admission. The facility should receive sufficient information and orders to provide continuity of care for all residents.</td>
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1. There is made available to the facility prior to or at the time of admission, resident information which includes current medical findings, diagnosis, and orders from a physician for immediate care of the resident.

2. Information about the rehabilitation potential of

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DMF 483.1123(a)
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<td>F10 (cont'd)</td>
<td>the resident and a summary of prior treatments are made available to the facility at the time of admission, or within 48 hours thereafter.</td>
<td>community, the attending physician provided current medical findings, diagnosis, prognosis, and orders.</td>
<td>- The order should cover: + Medications and treatments + Diet + Therapies (P.T., O.T., Speech) + Activities (bedrest, ambulatory, able to participate with any specific limitations on activity).</td>
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</table>
| Resident Supervision by Physician | Observe resident for any problem/conditions that should be addressed by physician, e.g., edema, loss of appetite, weight loss, etc. | Ask Resident:  
- How often physician visits?  
- If physician has discussed plan of care and medical treatment?  
- If resident feels treatment and/or plan of care meets his/her needs.  
- What kind of questions do you ask the physician about your health problem? (Cite examples).  
Ask Licensed Nursing Staff:  
- How often physician visits and is it often enough to meet resident's need?  
- Does physician participate in evaluation and reevaluation of resident's plan of care?  
- Does plan of care meet resident's needs?  
- Is physician available in an emergency?  
- Is physician available to discuss resident's treatment and care?  
Ask Administrator:  
- Facility's policy regarding a physician to provide care in the absence of the resident's own physician.  
- Facility's policy on physician visits. | Review medical records of selected for indepth review for:  
- A current plan of care that is based upon physician's orders and resident needs.  
- Evidence that the plan is reviewed and revised as needed.  
- Evidence through physician's progress notes, nurses notes, physician's orders, that the physician participates in the resident's overall plan of care.  
- Evidence that rehabilitation potential is addressed.  
- Long range plans include an estimate of the length of time for skilled nursing care and discharge plan.  
- Physician's orders for medications and treatments on admission and during stay.  
- A medical evaluation completed within 48 hours of admission unless done within 5 days prior to admission that includes attention to needs such as diet, vision, hearing, speech | Medical records should provide evidence that the residents are under the supervision of a physician by the coordination of physician's orders and progress notes with the resident's plan of care and observations of resident's needs. There is evidence that the physician reviews and revises the plan of care as needed. There is evidence that the physician services are available to the residents when the residents need such services. An alternate schedule for physician visits may be established if the attending physician determines that the resident need not be seen every 30 days. Justification for the decision is placed in the resident's medical record and is reviewed by the U.K. Committee and State medical review team. Where there is a change in the resident's condition and the physician has failed to document his findings or evaluation of the condition, the physician has failed to provide |
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<td>F114 (cont'd)</td>
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<td>F115</td>
<td>3. A physician is available to provide care in the absence of any resident's attending physician.</td>
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<td>F116</td>
<td>4. Medical evaluation is done within 48 hours of admission unless done within 5 days prior to admission. NOT ICFs.</td>
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<td>F117</td>
<td>5. Each SW resident is seen by their attending physician at least once every 30 days for the first 90 days after admission.</td>
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<td>level of activity, emotional adjustment.</td>
<td>Evidence of his evaluation of resident needs and supervised care. A physician is available to respond within a reasonable time when a resident needs medical attention.</td>
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<td>Evidence in care plans and treatment records that physician's orders are being implemented.</td>
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<td>Discrepancies in medication record, diet order, intake, and output records.</td>
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<td>Evidence that an alternate physician provided care if applicable.</td>
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<td>Progress notes by physician at least every 30 days for first 90 days (ICF at least every 60 days).</td>
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<td>Review of medications and treatments every 30 days or 60 days if an alternate schedule of visits has been approved.</td>
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<td>Documentation of physician observations, actions, and plans for treatment.</td>
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<td>Justification for alternate schedule of visits.</td>
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<td>A few closed records should be reviewed to determine if residents were appropriately discharged by an order written by the attending physician. Also review</td>
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<td>Although medical evaluation can be noted as a revision of the previous MRP. A statement such as &quot;no change&quot; when in conflict with the status of the</td>
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<td>F117 (cont'd)</td>
<td>discharge plans to assure that they were adequate and implemented. Verbal medication orders are countersigned by a physician. Physician is reviewing all medication orders every quarter. resident on this admission to the facility, does not constitute a medical evaluation. Verbal medication orders must be countersigned with 48 hours.</td>
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**Exception:** Only medications must be reviewed quarterly for ICF residents.

**F110:** Progress notes are written and signed by the physician at the time of each visit, and all orders are signed by the physician.

**F120:** Alternate physician visit schedules that exceed a 30-day schedule adopted after the 90th day following admission are justified by the attending physician in...
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<tr>
<td>F128 (cont'd)</td>
<td>the medical record.</td>
<td>These visits cannot exceed 60 days or apply to patients who require specialized rehabilitation schedules.</td>
<td>Exception: ICF residents must be seen every 60 days unless justified otherwise documented by the attending physician.</td>
<td>C. Emergency Services</td>
<td>F121 SNF 485.1123(c)</td>
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<tr>
<td>F121 Emergency services from a physician are available and provided to each resident who requires emergency care</td>
<td>Ask Staff:</td>
<td>- Are you aware of physician reporting procedures and medical protocols to be followed during a fire emergency?</td>
<td>- If records document an accident or a medical emergency, was the patient seen by a physician or was the physician notified promptly of the emergency?</td>
<td>- Review physician's orders to see if specific medications or treatments were ordered to treat emergency situation if applicable.</td>
<td>- Surveyor verifies that there are readily available written procedures for securing a physician in case of emergency. - Names and telephone numbers are posted or on rolodex. - An alternate physician is designated.</td>
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<td>1122 (cont'd)</td>
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<td>- Review physicians progress notes to see if emergency situation was addressed.</td>
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**INTENT**: To assure that a physician has overall responsibility for the management and supervision of the residents care.

- There is provision for:
  - Notification of attending physician/emergency and other responsible person.
  - Arrangements for transportation.
  - Preparation of reports.
  - There is evidence in the medical records that proper procedures have been carried out.
  - Residents with sudden changes in condition have been evaluated by the physician.
### Long Term Care Survey

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<td><strong>Nursing Services</strong></td>
<td>F123 SNF 405.1124</td>
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<td>F124</td>
<td>SNF 405.1124(c)</td>
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<tr>
<td>F126</td>
<td>ICF 442.1124(c)</td>
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A facility provides nursing services sufficient to meet the needs of all residents every hour of each day.

#### Basic care provided to residents:
- Surveyors observe the basic care provided by staff to the residents. Listed below are suggested areas of attention which may provide evidence of the quality of personal care:
  - Eyes/Mouth
  - Presence/Absence of:
    - Secretions forming around eyelids, redness or irritation of eyes.
    - Eyeglasses worn when appropriate are clean, in good repair and fit properly.
    - Backs of ears scaly, obvious wax build-up, discharge, odor.
    - Hearing aid worn when appropriate, is in good repair and working.
    - Dried food particles or drool, etc. around mouth.

#### Ask Resident:
- If the resident’s clothing is inappropriate, ask:
  - Did you choose your clothing today? Is this what you want to wear?
  - Do you have other clothing available?
  - If the resident is not clean, poorly groomed, or inappropriately groomed, ask the resident:
    - Have you had any help in caring for yourself today (e.g., washing your face, brushing your teeth, etc.)?
    - How often do you have a bath/shower?
    - How often is your hair washed?
    - How often do you brush your teeth/clean your dentures?

#### Nursing notes, flow sheets or bathing records should indicate that the care plan for grooming and personal hygiene is being followed. For example:
- Bathing schedules are being followed (including the use of any soaps or special lotions).
- Assistance instruction and/ or supervision is being provided as identified for each activity.

Nursing documentation should also indicate resident response or any changes in the resident’s behavior, reaction to an activity, or the ability to carry out grooming and personal hygiene activities. Look for indicators of progress toward a goal or further determination of resident functioning.

Refer to information on observation. A pattern of evidence of poor personal care indicates non-compliance unless the care plan specifically deals with this and appropriate planning and implementation is occurring.

The regulations require that individual preferences are taken into account when providing for grooming and personal hygiene and that residents are encouraged in self-care activity. On your patient interview substantiate compliance with the regulations!
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<td>F127 (cont'd)</td>
<td>Dentures worn when appropriate and in good repair.</td>
<td>- Odors presence/absence of: Body odors - Hair/Scalp - Clean and free of rashes - Hair combed - Nails are clean and appropriate length - Clothing is appropriate, clean, and in good repair. - Extremities elevated as necessary while in chair or wheelchair. - Appropriate techniques to prevent infection. - Use of whirlpool as a treatment modality as available and appropriate. - With resident’s permission check: - Heels, feet and toes - Talus joint mobility - Scapular area - Sacrum - Buttocks - Bony prominences in contact with braces - Condition of stump (especially diabetic)</td>
<td>resident is participating in dressing retraining program? - Special consideration might be given to the demented patient who frequently “borrows” clothes and for whom removal may elicit catastrophic reaction whether clothing “matches” may not be the most important issue in the care of these patients. Ask Direct Care Staff: - How do you choose what clothing each of your residents wear each day? - Do you have a specific schedule for washing residents’ hair? - How did you learn to bathe residents? - How did you learn to wash residents’ hair? - How did you learn to shave residents? - How do you handle situations when residents want to wear dirty clothes, or mismatched clothes? - How much care do you let the residents do on their own?</td>
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<td>F127 (cont'd)</td>
<td>&lt;br&gt;Observe with residents' permission: &lt;br&gt;- General condition of skin &lt;br&gt;- Redness &lt;br&gt;- Blanching &lt;br&gt;- Soft/dry/rough etc. &lt;br&gt;- Rash/irritation &lt;br&gt;- Bruises &lt;br&gt;- Scabs &lt;br&gt;- Free of above &lt;br&gt;- Measures taken to prevent skin breakdown. &lt;br&gt;- Pressure sores &lt;br&gt;- Pressure sores Rx &lt;br&gt;- Factors contributing to prevention of pressure sores &lt;br&gt;- Overall cleanliness and maintenance of dry and aerated skin (uncompromised by urine/face/perspiration) &lt;br&gt;- Padding for pressure points and bony prominences including padding on bed/chair &lt;br&gt;- Proper gentle massage to bony areas several times a day.</td>
<td>Ask Residents: &lt;br&gt;- Are your feet usually swollen? &lt;br&gt;- Do you know what causes the swelling? &lt;br&gt;- What do you do to alleviate it? &lt;br&gt;- Is this discoloration normal for you? &lt;br&gt;- How did this wound/bruise develop? &lt;br&gt;- Are the treatments done about the same time every day? &lt;br&gt;- What staff person has looked at your skin recently?</td>
<td>Look at nursing notes and P.O.C. for evidence of: &lt;br&gt;- Planned preventive measures &lt;br&gt;- Treatments/intervention including nutrition &lt;br&gt;- Routine assessment/evaluation of skin condition &lt;br&gt;- Documentation of specific skin problems with location number, severity, measurements as appropriate, and cause &lt;br&gt;- Progress or lack of progress in healing &lt;br&gt;- Assessment/Reevaluation of interventions with alterations in plan &lt;br&gt;- Appropriate nutritional plan &lt;br&gt;- Methods to control edema of lower extremities</td>
<td>Preventable pressure sores are not occurring. Ulcers present are treated on a routine basis according to P.O.C. Is skin clean? Is resident dry? Is turning schedule adhered to? Are linens clean and smooth? Do personnel know preventive measures and practice these? Has a nutritional assessment been done, and if appropriate, recommendations implemented?</td>
<td>&lt;br&gt;Dietetic Services 485.1725(1)(c)(v) 442.332(a)(1)(i)(b)(i) &lt;br&gt;Activities 485.1124(b) 442.345(a) &lt;br&gt;Patient Care Management 485.1124(c) 442.341 &lt;br&gt;Rehabilitation Nursing 485.1124(e) 442.362 &lt;br&gt;Supervision of Patient Nutrition 485.1124(f) 442.332(b)(2)</td>
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### Long Term Care Survey

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</table>
| F12B-129 (cont'd) | * Regular assistance for resident to turn or shift weight (bed rails, footboards, trapeze)  
* Bed linens, clothing, underpads smooth and free from wrinkles.  
* Elastic bandages or hose are smooth and wrinkle free.  
* Elastic bandages wrapped smooth with appropriate overlap.  
* Dietary/Nutritional support for skin integrity. (See Guidelines for Dietary/Nutrition  
* Prevention of shearing force when resident's position altered by staff.  
* Turning and repositioning as needed.  
* Care and treatment:  
* Turning and repositioning every two hours or as needed (e.g., alternative approach that is justified by the facility.)  
* Positioning of the ulcer site or protection of affected areas.  
* Use of effective pressure relief devices. | Ask Direct Care Staff:  
- What can you tell me about Mr./Mrs.____  
- Swollen feet/wounds/bruises/etc.?  
- What do you do for them?  
Ask Charge Nurse:  
- How did _______ get cuts, bruises, etc.?  
- What is being done to prevent further occurrence?  
- What treatment is he/she receiving? | | | Resident Super- vision by Physician 485.112(b) |
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| Wounds/Wound Dressings F626 | - Condition of dressing - i.e., clean, firmly secured unless contraindicated.  
- Observe, if possible, and with resident’s permission, a dressing change  
- Pre-dressing Removal Equipment and supplies organized  
- Hands washed  
- Residents provided with privacy  
- Dressing  
- Old dressing observed for drainage?  
- Wound examined  
- Appropriate technique used  
- Proper disposal of old dressing?  
- Post dressing  
- Does staff wash hands?  
- Return resident to comfortable position or previous activity?  
  Ask Resident:  
  - How often is the dressing changed?  
  - By whom is the dressing changed?  
  - Does it seem dressing changes are frequent enough?  
  - Are there any odors from the dressing?  
  - If not, what are the differences?  
  - Do you feel confident that the wound is being well cared for?  
  - Is the area/wound healing?  
  - What caused the ulcer, wound, etc.? Is it healing? Does the staff keep you informed of its status?  
  Ask Staff:  
  - Specific treatment and schedule for each resident?  | - Physician orders for wound care  
- Progress notes detailing condition of wound - i.e., size, drainage, surrounding tissue, odor  
- Treatment provided  
- Progress/change  
- Plan of Care (POC)  
- The plan of care should address:  
  - Area in need of treatment, treatment to be performed, frequency, and responsible staff.  
  - All necessary solutions, ointments, irrigations, types of dressings, and materials.  
  - Any necessary precautions, drains, if present, sutures and tubing.  
  - Specific goals of treatment as well as any problems or limitations imposed as a result of treatment.  | - Physician orders, your observations, progress notes and POC should reflect the same information.  
- Treatment provided over a period of time with no improvement and no re-evaluation also would represent non-compliance unless nursing/physician progress notes address the “no improvement” problem.  
- Compliance is evidenced by:  
  - treatment given according to doctor’s orders and POC  
  - use of appropriate technique when caring for wound/changing dressing (e.g., follows facility’s written procedures)  
  - periodic evaluation of healing process and revision of care plan as needed.  |  |  |
### Long Term Care Survey

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<td>Restraints</td>
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When residents require restraints, the application is ordered by the physician, applied properly, and reviewed at least every two hours. (See also information under Resident rights—freedom from abuse & restraints)

- Direct to evidence of:
  - Proper application
  - Proper use
  - Maintenance of good body alignment
  - Resident observation, release, and exercise

- Observe frequently throughout your visit to validate care. Specific observations should include the following items:
  - Type of restraint: belts, wrist or ankle cuffs, blanket, restraints, vests, bed nets, locked, etc.
  - When locked restraints are used can you readily find the key and/or scissors as well as geriatric chair or goniometer tray in place for prolonged periods.
  - Protective devices and/or safety devices that are used as restraints must be evaluated as restraints.

- Appropriate application: skin protected from injury (restraint neither too loose nor too tight to prevent

- Use of restraints may be precipitated by an "emergency" situation in which there is a threat to the resident's health or safety, or a threat to the health and safety of others due to the resident's behavior.

- Restraint residents may not be coherent or rational enough to respond to questions and caution in interviewing therefore, must be exercised. However, observation of a resident in a goniometer chair with a table or a resident in a wheelchair with restraint for several hours would warrant appropriate questions as to when the staff last assisted him or her to move about or whether the resident would like to get out of the chair. Staff interviews focus on the reason why the resident is restrained.

- Ask direct care staff and charge nurse:
  - Where, why, and how to release and apply restraints?
  - Why is the resident

- Physician orders for restraint: reason, length of time, type

- Progress notes:
  - Describe the resident's status/behavior which prompted the use of the restraint.

- If a chemical restraint, the order should indicate a specific time period for its use as well as a stop date.

- Plan of Care should:
  - Identify other methods or therapies that are being used in conjunction with restraints.
  - What alternatives to restraints have been considered?
  - Identify staff responsible for observing the resident (every 30 minutes), and releasing and exercising the resident (every two hours for at least 10 minutes).

- Time intervals should be identified.

- Indicate involvement and input of other disciplines necessary to overcome the problem.

- Indicate a specific period of time for

- Is there a physician's order, including the circumstances in which they will be used, the length of use, and the type of restraint?

- Is the restraint applied properly?

- Is it released at least every two hours and the resident provided with exercise and toilet facilities if needed?

- Does the staff observe the resident frequently while he/she is restrained?

- Are chemical restraints administered in accordance with physician’s orders?

- Is the order for restraints renewed only after a reassessment of the patient?
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<td>F130 (cont'd)</td>
<td>rubber and blistering or impeded circulation</td>
<td>- Body alignment and support: use of pillows, footboards, and wheelchair footrests to maintain appropriate posture, circulation, and to prevent skin injury or breakdown. - Periodic release and exercise: exercise may include ambulation, range of motion, massage, or other opportunities for motion (at least 2 hours every 2 hours during day and evening hours). - Chemical restraints: residents appear drowsy throughout the day (may indicate tranquilizers or other drugs are being used to limit or control behavior for staff convenience).</td>
<td>- How long has the resident been restrained? - Was the resident given an option of restraint? - When were you taught the use of restraints? - By whom? - If chemically restrained (excessively sedated) - Why is this done? - Whether alternate means of restraint have been attempted, for how long this will continue, etc. This should elucidate from staff whether the chemical restraint is necessary, or whether it is done for staff convenience by controlling resident behavior. - Do you ask the resident for permission before using restraints? - How does the restrained resident summon assistance? - What is the usual time frame for assistance to reach the restrained resident? * Ask Resident: - Why are you restrained? - What would happen if the restraint were removed? - When do you use bed rails? - What purpose do they serve? - How do you gain assistance?</td>
<td>using the restraint.</td>
<td>- Indication of assessment of factors which precipitate residents behavior which has warranted restraints and plans to intervene early enough to prevent occurrence. - Type, duration and frequency of exercise should be documented. - An assessment of why restraints are continued should be documented.</td>
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<td>Bowel and Bladder</td>
<td>There should be a chart/record in the resident's room on which the program is documented accurately. If the room is located a distance from the toiletting room or for residents with problems ambulating, a commode may be present in the room. Verify that a call light is available to the resident if non-ambulatory or restrained. Are fluids available at bedside? Is there roughage on meal tray? Diet is appropriate to enhance elimination? Ask Resident: Suggested questions are: How do you deal with constipation/diarrhea? Are you involved in a special bowel/bladder training program? If so, how does your program work? Any problems with it? Any successes to date? What does the staff do for you in this matter? Are they consistent and timely? How long do you have to wait to be taken to the toilet?</td>
<td>Both the resident and direct care staff should be interviewed and should exhibit a good understanding of the importance of maintaining a regular schedule of elimination. If neither are aware of the intake and toileting schedule, then determine whether they are appropriately planning the resident or carrying out a retraining program. Verify that the resident is aware of the time and when the resident is on a retraining program and knows the content of the program.</td>
<td>Physician orders if required by facility policy. Nursing notes for Assessment. Documentation of techniques and progress. Re-evaluation. Plan of care. The plan of care should clearly address: Goals that resident will aim for. Methods to accomplish the goals. Schedule for fluid intake. Schedule for toileting. Responsible staff. Any limitations the resident may encounter as a result of either incontinence or the training program. Progress notes/Physician orders for cause of incontinence. Laboratory tests of kidney function when available. Treatment for diarrhea/constipation. Resident preference for treatment of constipation. Recently admitted and newly incontinent resident should be thoroughly assessed for any incontinence management program.</td>
<td>Are all incontinent patients assessed for cause of incontinence and ability to be helped by a bowel/bladder rehabilitative training program or an incontinence management program? Are all appropriate residents involved in bowel/bladder training programs or incontinence management and there is a schedule that shows when the program will be started? Is there evidence of follow through on all shifts? For residents not on bowel/bladder retraining programs the plan of care should address specific measures for managing incontinence with a view to prevention of skin and other problems and maintenance of resident dignity.</td>
<td>Nursing Services 405.1124(c) Dietetic Services 405.1125(c)</td>
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### Long Term Care Survey

<table>
<thead>
<tr>
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<tr>
<td>F131 (cont'd)</td>
<td>- When a resident puts on his/her call bell for toileting assistance, how long is it before assistance is given? - Observe pre-meal toileting. - Privacy provided. - Schedule for toileting should allow for resident's normal sleep pattern to avoid disrupted sleep.</td>
<td>Ask Nurse Aides and Charge Nurse: - Will you describe the resident's bowel/bladder (B/B) training program? - How long has it been in effect? - When will you evaluate the results? - If this program is not successful: - What assessment was done to determine B/B status? - For residents not on B/B retraining program: what is the facility program for managing incontinence? at least 7 days for the cause of incontinence and when appropriate an intensive bowel and bladder B/B training program should be instituted. - A trial B/B training program is suggested for all residents with incontinence problems: 1 &amp; 0</td>
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<tr>
<td>Catheter Care</td>
<td>F132 SNF 405.1124(c)</td>
<td>Each resident with a urinary catheter receives proper routine care including periodic evaluation. The indwelling catheter should promote a continuous flow of urine unless ordered otherwise. The surveyor should also observe for the following: - Aseptic supplies for catheter insertion and care. - Proper positioning of the tubing and drainage bag. - Cleanliness of the</td>
<td>Ask Resident: - What is the tubing/catheter for? - Why do you have one? - Does it cause any discomfort? - If it does, what is done about it? - How do you feel about having the catheter? - Is any special care given in relation to the catheter?</td>
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<td>The surveyor should verify that there is a physician's order for an indwelling catheter, including the type and frequency of catheter care. If irrigation is ordered, the order should include type of solution and frequency of irrigation. The record should also indicate the color, consistency, and amount of urinary drainage.</td>
<td><em>The facility should follow accepted professional standards in their catheter care. There should be medical reasons for catheter insertion – staff convenience cannot be justification. Direct care staff should know signs and symptoms of urinary tract</em></td>
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Infection Control

405.1125(a)
<table>
<thead>
<tr>
<th>SURVEY AREA</th>
<th>OBSERVATION</th>
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<th>EVALUATION FACTORS</th>
<th>CROSS REFERENCE</th>
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</thead>
<tbody>
<tr>
<td>F132 (cont'd)</td>
<td>Tubing and drainage bag.</td>
<td>Ask Nursing Aide and Change Nurse:</td>
<td>- Assessment should address:</td>
<td>Infections (U.T.I.s) and these should be reported and treated promptly.</td>
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<tr>
<td></td>
<td>- Color and consistency of urine in bag</td>
<td>- How do you routinely position and secure catheters and drainage bags?</td>
<td>+ Need for an indwelling catheter.</td>
<td>*The Center for Disease Control has developed standards for catheter care which may be used but it is not a requirement.</td>
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<td></td>
<td>- Availability and accuracy of documentation on the MAR sheet if ordered or policy</td>
<td>- How often is each part of the system changed?</td>
<td>+ Resultant problems or limitations.</td>
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<td></td>
<td>- Proper equipment for ambulation - Tag bag if resident is ambulating (if ordered)</td>
<td>- What are the indications for insertion of the catheter?</td>
<td>- Plan of Care should address:</td>
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<td></td>
<td>- Availability of fluids.</td>
<td>- What is the facility's procedure for routine catheter care?</td>
<td>+ Type of catheter and type and frequency of care.</td>
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<td></td>
<td>- When indicated monitor intake to ensure adequate intake and output or conformance with physician orders.</td>
<td>- How do you observe for U.T.I.'s in residents with indwelling catheters?</td>
<td>+ For irritation, the rationale, the type of solution, amount, and frequency of irritation.</td>
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<td></td>
<td>- How many observed residents are on catheter care?</td>
<td>- What is the facility's procedure for the cleansing and storage of reusable catheter equipment and drainage receptacles?</td>
<td>+ Frequency of symptoms which would precipitate catheter change.</td>
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<td>- How do you care for catheter tubing?</td>
<td>+ Line frames of catheter change and responsible staff.</td>
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<td>+ Appropriate increase in oral fluid intake.</td>
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<td>+ Intervention: The record must reflect:</td>
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<td>+ When and by whom the catheter was inserted and for what reason.</td>
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<td>+ Any special care provided</td>
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<td>+ New problems or changes</td>
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<td>+ Only appropriately trained staff should deliver catheter care.</td>
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<td>+ Only licensed staff should insert</td>
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<td>F132 (cont'd)</td>
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<td>indwelling catheter.</td>
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<td></td>
<td>* The specific type and size of equipment used should be noted.</td>
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<td>* Signs and symptoms of urinary tract infections (UTI) should be acted upon and documented as to follow-up.</td>
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<td>* Evaluation/Reevaluation: The record should reflect that the resident:</td>
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<td>* Is assessed for UTI.</td>
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<td>* Has no abdominal distention.</td>
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<td>* Notes should also include:</td>
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<td></td>
<td>* The color and odor of urine and the development of any problems after insertion of indwelling catheter.</td>
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<td>* Verify that catheter is patent.</td>
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### Long Term Care Survey

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</table>
| Injections  | - Observe for preparation of injection - i.e. maintenance of sterility; correct dilution, handwashing, before preparation, etc.  
- Observe injection site for:  
  - Redness  
  - Discoloration  
  - Swelling  
  - Lesions  
- Observe for proper technique when injection is given  
  - Correct site  
  - Correct needle size  
  - Correct volume of drug  
- Sterility maintained  
- Resident is observed for any adverse reaction  
- What is the disposal method for used needles or syringes?  
- Ask Nurse:  
  - What is your plan for alternating injection sites? Show me.  
  - What is the medication for and what are potential adverse reactions?  
  - Is there nonspecific pain at the injection site or shooting pains down a limb?  
  - Is there skin irritations or lumps under the skin?  
  - If adverse reaction occurs, how soon are they reported?  
  - Could this be given by any other route? | - Physician order sheet  
- Nursing notes for:  
  - Resident response to medication if appropriate  
- Any problems noted at injection site  
- Any other adverse reactions  
- Site of injection  
- Plan of care  
- Injection site  
- Care for any special problems related to the injection.  
- Infection Control reports for any infections connected with injections.  
- Is the medication administered according to the physician's order?  
- Is proper technique used in preparation and administration including site rotation?  
- Does the nurse administering the medication know the expected action of the drug?  
- If infection control reports show infections reported at injection sites.  
- Is the resident's response to the medication noted in the progress notes? |
| F155/SN 1124(c) | | | | |

*Staff Development 405.112(1b)  402.314  
Infection Control 405.1135(1b)*
LONG TERM CARE SURVEY

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<tbody>
<tr>
<td>Parenteral fluids</td>
<td>The surveyor should observe that parenteral fluids are administered with safe, aseptic technique providing fluids as ordered by the physician. Safety and comfort measures are to be taken insuring maximum protection and optimum hydration of the resident.</td>
<td>Ask Resident:</td>
<td>- Physician's order for parenteral therapy specifying type of fluid, rate of infusion/hour, and additives, if any, is available and current.</td>
<td>- Is the parenteral fluid administered according to the physician's order and in accordance with accepted nursing practice?</td>
<td>Patient Care Management 485.124(d) 442.341</td>
</tr>
<tr>
<td>SNF 405.1124(c)</td>
<td>- The surveyor should note the following items:</td>
<td>- Why the resident is receiving I.V. therapy?</td>
<td>- Any adverse reactions are noted in the medical record.</td>
<td>- Are infusions noted in a timely manner before a large amount of fluid is infused?</td>
<td>Resident Care Policies 485.1123(1)</td>
</tr>
<tr>
<td></td>
<td>- Labeling of the solution bottle/Bag</td>
<td>- What is the drip rate in (the amount of fluid to be received per hour)?</td>
<td>- Any adverse reactions are noted in the medical record.</td>
<td>- Is the facility procedure for care of the IV site and tubing changes followed for all patients unless contraindicated?</td>
<td>Infection Control 485.1125(b)</td>
</tr>
<tr>
<td></td>
<td>- Rate of infusion (cc/ml)</td>
<td>- How often the dressing is changed?</td>
<td>- Record indicates:</td>
<td>- Does documentation reflect what the patient received, any problems, and the patient's response to the parenteral fluid?</td>
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<td></td>
<td>- Date and time started</td>
<td>- How often the tubing is changed?</td>
<td>- Infusion started by whom; time, rate of flow</td>
<td>- Have any adverse effects been caused by administration of IV fluid?</td>
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<td>- additive, if any</td>
<td>- What are possible side effects?</td>
<td>- Note is made of observation of pain or swelling at infusion site.</td>
<td>- If yes, were those preventable?</td>
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<td></td>
<td>- Any signs of swelling or redness at site</td>
<td>- How often is the site changed?</td>
<td>- The need or reason for parenteral fluids.</td>
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<td></td>
<td>- Site dressing is clean, dry and dated</td>
<td>- How often is the infusion checked for drip rate and the remaining volume to be administered?</td>
<td>- Response to the therapy.</td>
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<td></td>
<td>- Accurate 360 of parenteral and P.O. fluids</td>
<td></td>
<td>- Problems and limitations encountered by the resident as a result of receiving parenteral fluids.</td>
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<td></td>
<td>- If splint (armboard) is used, it is applied to prevent movement but not impede circulation</td>
<td></td>
<td>- Plan of Care:</td>
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<tr>
<td></td>
<td>- Positioning of I.V. tubing</td>
<td></td>
<td>- The plan of care should include:</td>
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<td></td>
<td>- Comfort of restraint used to allow for maximum resident freedom while preventing movement of I.V. site.</td>
<td></td>
<td>- Type, rate of infusion /hour, and additives (if ordered).</td>
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**LONG TERM CARE SURVEY**

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<tr>
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<tr>
<td>F133 (cont'd)</td>
<td>specified goals for correction, time frame, and responsible staff. Documentation must include time administered and by whom, the amount of fluid infused, and any other special care administered as a result of IV therapy (i.e., mouth care assistance with ABIs, etc.). The record must reflect: + Conditions of site and any inflations, phlebitis, necrosis, etc. noted, along with measures taken to correct these. + The resident's response to therapy + Changes in laboratory studies. <em>Plan of care would not be modified for a one-time IV infusion.</em></td>
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<tr>
<td>Colecotomy/Ileostomy F133 SHF 485.1124(c)</td>
<td>The surveyor should ascertain that the facility is providing appropriate nursing care to those residents who have had bowel surgery resulting in a colostomy or ileostomy. It is recommended that the surveyor, with the resident, determine that: Ask Resident: - Why was the ostomy performed? - How do you feel about the ostomy? - Does it ever cause you problems (e.g., pain, skin problems, odors, accidents)? If so, what compliance would be indicated if residents are physically and emotionally comfortable with the ostomy with minimal or no skin problems; If residents are not comfortable with the ostomy, are having skin or other problems, the facility</td>
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### Health Care Financing Administration, HHS § 488.115

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<tr>
<td>Colostomy/Fistostomy F133 (cont'd)</td>
<td>dents permission, observe care being given to determine that proper techniques are being used. The following steps should be taken to assure that proper ostomy care is being provided.</td>
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<td>- The ostomy dressing should be changed or the bag emptied and thoroughly cleaned promptly after each bowel evacuation or more frequently, if drainage continues.</td>
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<td>- The peristomal skin should be cleansed and dried, and appropriate measures taken to prevent excoriation and infection.</td>
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<td>- The resident's privacy should be considered while providing care.</td>
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<td>- The resident should be provided with information and instruction in self-care at the appropriate level of understanding.</td>
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<td>- The resident should be observed for signs of withdrawal, distress, anxiety, or other emotional responses which may be related to his/her conditions.</td>
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<td>- Ask Staff:</td>
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<td>- What does the staff generally do with or for the ostomy? Are they consistent and timely?</td>
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<td>- Has staff talked to you about doing some of the care for this? If so, what was the outcome? If not, is this something you'd be interested in learning more about?</td>
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<td>- Ask Other Nursing Staff:</td>
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<td>- Is there a facility procedure for ostomy care?</td>
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<td>- Do you have skin problems with your ostomy?</td>
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<td>INTERVIEWING</td>
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<td>documented as established through management of diet, fluid intake, exercise, and the use of prescribed laxatives, suppositories, and/or irrigations.</td>
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<td>- Ostomy care is documented in the resident's record along with a description of the excreta.</td>
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<td>- Problems in irregularity, skin breakdown, or other observable concerns are documented and reported to the physician.</td>
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<td>- Documentation indicates that nursing measures are taken to assist the resident who is experiencing problems in understanding and/or accepting the presence of the ostomy.</td>
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<td>- Documentation of nursing measures to maintain skin integrity.</td>
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<td>- Assessment The assessment should indicate:</td>
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<td>- Needs, problems, and limitations as a result of an ostomy.</td>
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<td>- Specific degree of...</td>
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<tr>
<td>Ostomy/Fistula F135 (cont'd)</td>
<td>- Her acceptance of the ostomy/pouching system.</td>
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</table>

- Self-care performed or assistance needed.
- Special skin care needs.
- Special dietary needs (emotional support).
- Medications and treatments if needed.
- Plan of care. The plan of care should clearly address:
  - Specific goals to overcome or improve the problem(s) identified.
  - Methods to accomplish the goal (training, assistance, supervision, treatments, emotional support).
  - Services necessary and who will perform the services.
  - Time frame for accomplishing goals.
### LONG TERM CARE SURVEY

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<tr>
<td>Respiratory Therapy</td>
<td>Aerosol Compressor or IPPB (Intermittent Positive Pressure Breathing Machine)</td>
<td>The surveyor must determine that the facility is providing respiratory therapy as ordered by the physician. Observation for this indicator should focus on the necessary equipment as well as on the resident. In order to determine that the necessary equipment is available, the surveyor must look for the following:</td>
<td>- Respiratory/oxygen therapy is performed or administered by appropriately trained staff.</td>
<td>- Staff Development 405.1121(h)</td>
<td>Staff Development 405.1121(h)</td>
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<td>- There is a physician's order for therapy, and it is specific as to rate of delivery, etc.</td>
<td>Infection Control 405.1133(b)</td>
<td>Infection Control 405.1133(b)</td>
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<td>- If the physician's order is for prn therapy, it should specify for what symptoms.</td>
<td>Patient Care Management 405.1124(d)</td>
<td>Patient Care Management 405.1124(d)</td>
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<td>- Any information gained from resident or staff is verified in the record.</td>
<td>442.341</td>
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<td>- Assessment: The assessment should address both the need or reason for therapy and any necessary reconsideration.</td>
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<td>- The kind, amount, frequency, and/or duration of therapy based on the physician's order.</td>
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<td>- Specific goals to overcome to improve any identified</td>
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**Physical Observations:**
- While interviewing the resident, observe for sounds of congestion. Note color of lips and nail beds.

**Ask Resident:**
- Do you ever feel short of breath?
- If yes, what is done when this occurs?
- Is the therapy helping you to feel better?
- Are there any problems with it?
- If so, how does the staff respond?
- Is the therapy consistently performed - both concerning time and method of providing it.

**Ask Staff:**
- What is the reason the resident is getting this therapy?
- What are the expected results?
- Can you demonstrate how you use the equipment?
- How often is the equipment cleaned?
- What are the infection control procedures in regard to use of res-
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</table>
| Respiratory Therapy F15 (cont'd) | stored wet. If it is not attached to the tubing, ask to see it. The mouthpiece is connected to the nebulizer cup. The surveyor should also check that all involved equipment is clean. | - Oxygen therapy The surveyor must establish that the facility is meeting the oxygen needs of the resident. When the facility does not have wall units, check that:  
  - There are enough cylinders for oxygen delivery.  
  - There should be flow meters and regulators for tanks in use.  
  - A wrench should be attached or stored close by.  
  - If using large cylinders (size 6 or H), look for a carrier since these tanks cannot be transported without it. | - Where is the emergency oxygen supply? | problems and/or limitations.  
  - Specific methods to accomplish the goals (observation, supervision, training, etc.).  
  - Who is responsible to perform therapy or assist in accomplishment of goal.  
  - Intervention - The record should display evidence that:  
    - The plan of care is functional.  
    - The therapy was administered in accordance with physician's order for the specified reason(s) by an appropriately trained staff member.  
    - Change in condition is documented and acted upon promptly.  
  - Evaluation/Nevailuation The record should reflect:  
    - The resident's response to therapy.  
    - If response was undesirable, evidence of further intervention.  
    - Any progress, deterioration, or development of new problems. | |
## LONG TERM CARE SURVEY

<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td>Respiratory Therapy F133 (cont'd)</td>
<td>the carrier, sitting on a metal skirt, or otherwise secured.</td>
<td>+ Based on the above information, possible modification of goals.</td>
<td></td>
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</tr>
</tbody>
</table>
<pre><code>                                                                                                                      | + There should be other necessary equipment available such as humidifiers, nebulizers, masks, nasal cannulas, l-pieces, etc., all should be dry and clean when stored.                                                                 |                                                                                                                                            |                                                                |                  |
                                                                                                                      | + Check to see that non bed-bound residents are not limited to their own chair/room when using oxygen (portable units will prevent social isolation).                                                                 |                                                                                                                                            |                                                                |                  |
                                                                                                                      | + Water reservoir is appropriately filled per manufacturer's instructions.                                                                                                                               |                                                                                                                                            |                                                                |                  |
                                                                                                                      | + Check to make certain the tank is not empty and that any tank is labeled as such.                                                                                                                                 |                                                                                                                                            |                                                                |                  |
                                                                                                                      | + Check for good oral hygiene of resident.                                                                                                                                                    |                                                                                                                                            |                                                                |                  |
                                                                                                                      | + The room should be posted with a &quot;No Smoking&quot; sign.                                                                                                                                             |                                                                                                                                            |                                                                |                  |
                                                                                                                      | - Residents on respirators:                                                                                                                                                                              |                                                                                                                                            |                                                                |                  |
                                                                                                                      | - Are alarm systems turned on?                                                                                                                                                                            |                                                                                                                                            |                                                                |                  |
                                                                                                                      | Residents on Respirators                                                                                                                                                                                 |                                                                                                                                            |                                                                |                  |
                                                                                                                      | Ask Staff (all levels):  What training have you had in caring for                                                                                                                                       |                                                                                                                                            |                                                                |                  |
</code></pre>
### LONG TERM CARE SURVEY

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<tr>
<td>Respiratory Therapy</td>
<td>• Is sufficient oxygen supply available?</td>
<td>residents on respiratoryators?</td>
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<tr>
<td>F133 (cont'd)</td>
<td>• Is the ventilator accessible to an emergency outlet?</td>
<td>- Can you show me how the alarm system works?</td>
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<tr>
<td></td>
<td>• Is the resident in a location that allows for frequent observation by staff?</td>
<td>- What is your procedure for pulmonary care?</td>
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<td></td>
<td>• How does the resident communicate with staff?</td>
<td>- What is your procedure for changing tubing and the water reservoir?</td>
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<td></td>
<td>• What level of staff (CNA, LPN, RN) caring for the resident?</td>
<td>- What happens if the power goes off?</td>
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<td></td>
<td>• Is such equipment at bedside?</td>
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<td>• Is there reserve back-up equipment?</td>
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<td></td>
<td>• What is the condition of the resident's skin around intubation tube/trephostomy?</td>
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<td></td>
<td>• Does the care given use appropriate techniques in caring of the patient?</td>
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<tr>
<td>Survey Area</td>
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<tr>
<td>Tracheostomy Care</td>
<td>Satisfactory tracheostomy care is a procedure which promotes a clean, unobstructed air passageway and maintains the skin integrity surrounding the tracheostomy site. The surveyor should determine whether: - Adequate supplies are available for the care of the tracheostomy such as tracheostomy kits, hydrogen peroxide, normal saline or sterile water, suction machine, catheter, sterile gloves, and clean dressings. - The resident is breathing without difficulty and is comfortable. - The dressing is clean, dry, and intact; the cannula is clean, in the proper position, and secured. - The skin surrounding the tracheostomy is clean and dry with no redness or inflammation. - The resident has adequate oral hygiene. - An extra tube, the same size as the one in the trachea, is available. Resident interviews must be guided by the resident’s communication ability. Ask Resident: - How long will you have it? - What care can you do for yourself? - What do you need help with? - Who helps you? - Is someone always available to suction him/her when needed? - Is the suction equipment always available in working order? - Is the dressing kept clean and comfortable? - Is the tube kept clean and changed as needed? - How often are the tubes and dressings changed? - Does he/she feel confident in the personnel caring for his/her tracheostomy? - What is communicating with staff and other residents like? - Are staff patient and do they allow you enough time to express your needs/thoughts/feelings? Ask Staff: - Why does resident have - The surveyor should determine that tracheostomy care is done as scheduled and as needed following the proper procedure. - Any special solutions that are needed should be addressed in the physician’s orders. - Assessment: The record should reflect that the need for tracheostomy care was assessed in terms of: + Frequency + Skin integrity surrounding the tracheostomy, noting redness, inflammation, and/or excoriations. - Plan of Care should include: + Specific times of tracheostomy and the responsible, appropriately trained person performing this task. + Specific problems relating to skin and breathing as well as the goals set to overcome these problems listing the appropriate personnel responsible. + Time frames for resolving problems</td>
<td></td>
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<td></td>
<td>Stoma and surrounding skin should be in good condition and if not, there should be treatment directed to resolving this problem. All staff caring for the tracheostomy must be trained and emergency procedures be known. All needed equipment must be available and in working order. Resident must at all times have readily available a means of communicating with the staff in an emergency.</td>
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<tr>
<td>SURVEY AREA</td>
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<tr>
<td>Tracheostomy Care</td>
<td>F73 (cont'd)</td>
<td>tracheostomy?</td>
<td>listed in goals.</td>
<td>Plan for periodic assessment of appropriateness of resident's own self care re: teaching or nursing assuming more responsibility as appropriate.</td>
<td></td>
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<tr>
<td></td>
<td>place, is available at bedside.</td>
<td>what training were you given to enable you to care for tracheostomy?</td>
<td>Intervention - The surveyor should look for documentation of:</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>- Does resident have an adequate method of communicating with the staff?</td>
<td>what is the procedure for tracheostomy care?</td>
<td>- Teaching or nursing assuming more responsibility as appropriate.</td>
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<td></td>
<td>- Does staff allow enough time for residents to communicate?</td>
<td>how often is the tube changed?</td>
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<td></td>
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<td>what do you do if the tube comes out?</td>
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<td>may I watch you do a dressing change?</td>
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<td>if not convenient, describe what you do.</td>
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<td></td>
<td>how do you communicate with a tracheostomized resident?</td>
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</table>
### Long Term Care Survey

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<tbody>
<tr>
<td><strong>Tracheostomy Care</strong></td>
<td></td>
<td></td>
<td>since this may require additional care planning.</td>
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<tr>
<td><strong>Suctioning</strong></td>
<td>Suctioning is necessary for any resident who is unable to cough up secretions</td>
<td>Suctioning is necessary for any resident who is unable to cough up secretions</td>
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<td></td>
<td>that are obstructing his airway.</td>
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<td>Suctioning may occur via the oropharynx, nasal route, or tracheostomy tube.</td>
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<td>Attempts should be made to observe a resident being suctioned should such an</td>
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<td>opportunity arise. If so, observe that a clean/aerobic technique is observed</td>
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<td>throughout and that the resident tolerated the procedure. The suction should</td>
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<td></td>
<td>not be bloody aspirated, cyanotic, or bronchospasm. Check that equipment</td>
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<td>is in good working order, frequency of procedure, etc.</td>
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<td>Resident observations which indicate need for intervention include:</td>
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<td>- Secretions are draining from a resident's mouth or trach and the resident</td>
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<td>is unable to</td>
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<td>Ask Resident:</td>
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<td></td>
<td>- How are you feeling now after the suctioning?</td>
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<td>- Does the suctioning seem to help?</td>
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<td></td>
<td>- Has staff explained to you the need for suctioning?</td>
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<td></td>
<td>- How often?</td>
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<td>- Who performs the suctioning? (i.e., nurses or nurses aides?) Do you feel</td>
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<td>safe with the staff performing the suctioning?</td>
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<td>- Does everyone do it the same way?</td>
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<td></td>
<td>Ask Staff:</td>
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<td></td>
<td>- When and where did you team for suctioning?</td>
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<td></td>
<td>- Tell me what procedure you use when you suction a resident.</td>
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<td></td>
<td>- Do you always have enough suction machines and catheters?</td>
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<td>- How frequently is suction tubing changed?</td>
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<td>- What provisions do you have for suctioning if the electricity is lost?</td>
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<td></td>
<td>- Assessment - The record should reflect that:</td>
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<td></td>
<td>- The resident is frequently observed for suctioning needs.</td>
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<td>- Any limitations a resident has as a result of his suctioning needs should</td>
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<td>be specifically noted.</td>
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<td>- Any problems resulting must be specified.</td>
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<td></td>
<td>Plan of Care should include:</td>
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<td></td>
<td>- Awareness of the resident's suctioning needs, goals, approaches, and</td>
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<td></td>
<td>responsible staff needed to improve the problem or at least to maintain the</td>
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<td>resident at his present status without further deterioration. The plan must</td>
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<td>clearly indicate specific approaches towards:</td>
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<td></td>
<td>- Prevention of skin problems around the trach if one exists.</td>
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<td></td>
<td>- Correction of any existing skin prob-</td>
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## Long Term Care Survey

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<tr>
<td>Suctioning</td>
<td>Cough or clear himself.</td>
<td>- Where are your emergency electrical outlets?</td>
<td>Blows.</td>
<td>Provision of good oral hygiene including a rigid schedule for mouth care, schedules, or procedures for maintaining clean equipment at bedside, as well as disposal of used (dirty) equipment.</td>
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<tr>
<td></td>
<td>- There are audible crackles or wheezes and/or diminished breath sounds.</td>
<td>- What is your procedure for disposing of the secretions from suctioning?</td>
<td>- How often does Mrs./Mr. need to be suctioned?</td>
<td>Route of suctioning (i.e., oral/nasal/trach).</td>
<td></td>
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<tr>
<td></td>
<td>- The resident is dyspneic.</td>
<td>- May I observe you when you suction Mrs./Mr.?</td>
<td>- Intervention - The record should indicate clearly that:</td>
<td>- The plan of care is being implemented. Documentation should reflect:</td>
<td></td>
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<tr>
<td></td>
<td>- Restlessness or agitation may also be an indication that suctioning is needed.</td>
<td></td>
<td>+ The number of times the resident required suctioning for what specific reason, and by whom the resident was suctioned.</td>
<td>+ Any special treatment the resident received in conjunction with suctioning.</td>
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<tr>
<td>Suctioning</td>
<td>F133 (cont’d)</td>
<td></td>
<td>(i.e., oral hygiene, skin care, etc.).</td>
<td>Evaluation/Reevaluation: The record should reflect:</td>
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<td></td>
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<td></td>
<td>- How well the resident tolerates suctioning procedures.</td>
<td>- Any bloody aspirate, cardiovascular abnormality, cyanosis, or bronchospasm.</td>
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<td>- Further interventions utilized to overcome or improve these.</td>
<td>- The amount of sputum as well as its color and consistency.</td>
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<td></td>
<td>- Any progress or lack of progress, deterioration, and/or the development of new problems.</td>
<td>- The evaluation should determine whether goals are being reached or if new goals must be addressed.</td>
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**Tube Feedings**

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<tr>
<td>F133</td>
<td>SNF 405.1120(c)</td>
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<tr>
<td></td>
<td>- Staff use proper technique in administering feedings and medications. Check to see that staff checks for location of tube before feeding and that tubing</td>
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<td></td>
<td>If the resident is able to be interviewed, suggested questions may be:</td>
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<td></td>
<td>Do you feel comfortable/safe with all the staff who perform the feeding?</td>
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<td></td>
<td>Tube Feeding Review:</td>
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<tr>
<td></td>
<td>- Plan of care</td>
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<td></td>
<td>- Must document tube placement and formula potency prior to each feeding.</td>
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<td>- Has the feeding been ordered by a physician?</td>
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<td>- Is tube feeding nutritionally adequate?</td>
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<td>- Have attempts been made to discontinue tube feeding if indicated?</td>
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**Nursing Services**

405.1120(d)(1)(v) 442.320(a)(1) 442.333(c)
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<tr>
<td>Tube Feedings F133 (cont'd)</td>
<td>Is irritated before and after addition of medication.</td>
<td>- The tube is clean and formula flows freely.</td>
<td>- Nasal gastric tube must be secured in a manner that avoids creating pressure on the nose and nasopharynx.</td>
<td>- Is skin free from irritation; much care is given several times daily? (More frequent mouth care in the case of continuous feeding.)</td>
<td>G215-1125(c)</td>
</tr>
<tr>
<td></td>
<td>- The equipment is clean and protected. If dressings are ordered, they are in place, clean, and dry.</td>
<td>- The nasal tube is securely but comfortably secured on the face with skin maintained intact and without irritation.</td>
<td>- Identify frequency, amount, and time span over which each feeding is accomplished.</td>
<td>- Have changes in resident condition been noted and addressed (weight loss, constipation, diarrhea, skin condition)?</td>
<td></td>
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<td></td>
<td>- The skin around the gastrostomy is kept clean and free from irritation or infection. It should be checked carefully for leakage of gastric contents.</td>
<td>- The skin around the gastrostomy is kept clean and free from irritation or infection. It should be checked carefully for leakage of gastric contents.</td>
<td>- Motivation and treatment records.</td>
<td>- Have observed problems been coordinated with other departments and resolved?</td>
<td></td>
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<tr>
<td></td>
<td>- A resident who has a NG tube for a prolonged period of time should be observed for possible complications, such as nasal erosion, sinusitis, esophagitis, gastric ulceration, and pulmonary infection.</td>
<td>- A resident who has a NG tube for a prolonged period of time should be observed for possible complications, such as nasal erosion, sinusitis, esophagitis, gastric ulceration, and pulmonary infection.</td>
<td>- Fluid intake records.</td>
<td>- Is feeding being monitored to ensure that feeding is occurring at the ordered/appropriate rate?</td>
<td></td>
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<td></td>
<td>- Resident is fed slowly with head elevated to 45 degrees during feeding and at least 2 hours post-feeding.</td>
<td>- Resident is fed slowly with head elevated to 45 degrees during feeding and at least 2 hours post-feeding.</td>
<td>- Number of calories as well as amount of additional water.</td>
<td>- Varied supplements as preferences allow?</td>
<td></td>
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</tbody>
</table>
### LONG TERM CARE SURVEY

<table>
<thead>
<tr>
<th>SURVEY AREA</th>
<th>OBSERVATION</th>
<th>INTERVIEWING</th>
<th>RECORD REVIEW</th>
<th>EVALUATION FACTORS</th>
<th>CROSS REFERENCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tube Feedings F13 (cont'd)</td>
<td>- Supplies for mouth care are in evidence, observe if possible for technique; mouth shows evidence of good care (i.e., moist, clean.)</td>
<td></td>
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</tbody>
</table>

**Nursing Services**

F137  
SNF (405.1124)  
ICF (442.318)  
B. Twenty-four hour nursing.

1. Assigned duties consistent with their education and experience based on the characteristics of the resident load.

F138  
2. Weekly time schedules are maintained.

F139  
3. There is a sufficient number of nursing staff.

<table>
<thead>
<tr>
<th>Question:</th>
<th></th>
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</thead>
<tbody>
<tr>
<td>Are personnel performing duties that are allowed under the State Nurse Practice Act?</td>
<td></td>
</tr>
<tr>
<td>Do you observe care being rendered in an appropriate, competent manner?</td>
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<tr>
<td>Does the time schedule posted indicate that at least the minimum required personnel are scheduled and actually on duty?</td>
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<tr>
<td>What is the usual response time before a call bell is answered?</td>
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<tr>
<td>In SNF's is an RN on duty during the day?</td>
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<tr>
<td>Are licensed staff and aide staff functioning in appropriate roles?</td>
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<tr>
<td>Where are staff spending their time?</td>
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<tr>
<td><strong>Ask Resident:</strong></td>
<td>- Do residents generally feel that people taking care of them know what they are doing?</td>
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<tr>
<td></td>
<td>- If no, explain.</td>
</tr>
<tr>
<td></td>
<td>- Are your treatments done in a consistent manner?</td>
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<tr>
<td></td>
<td>- If no, explain.</td>
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<tr>
<td></td>
<td>- Do you feel that there are enough people here to take care of you?</td>
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<tr>
<td></td>
<td>- If no, explain.</td>
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<tr>
<td></td>
<td>- How long do you usually wait for help when you put your call light on?</td>
</tr>
<tr>
<td></td>
<td>- Is there anything that doesn't get done as often as it should?</td>
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<tr>
<td><strong>Ask Staff:</strong></td>
<td>- Do you feel qualified to do all the work you are assigned to do?</td>
</tr>
<tr>
<td></td>
<td>- If no, explain.</td>
</tr>
<tr>
<td></td>
<td>- Do you feel you have enough training to keep up with the care the residents require?</td>
</tr>
</tbody>
</table>

- Review progress notes to determine who is giving care.
- Review care plan to determine who the facility has assigned to care responsibility to.
- Check staffing sheets for minimal requirements and time and attendance for actual staffing.
- Review charts maintained for ARD medications, I & O, restrictions, etc., to assure that sufficient staff are available for carrying out responsibilities as specified in patient care plans.

- All nursing personnel must function within their State Nursing Practice Act levels of staffing meet at least minimum requirements.
- Nursing care needs must be identified by the facility & documentation, resident and staff interview should determine if these needs are met.
- All nursing staff should have education or training to prepare them for the care they perform.

**Patient Rights**

405.112(k)(11)

**Patient Care Policies**

405.112(k)(g)

**Medical Records**

405.112(1)

442.318(a)(c)

**Patient Care Management**

405.112(4)

442.341

**Staff Development**

405.112(b)

442.314
<table>
<thead>
<tr>
<th>SURVEY AREA</th>
<th>OBSERVATION</th>
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</tr>
</thead>
<tbody>
<tr>
<td>F139 (cont'd) available to meet the total needs of all residents.</td>
<td>Check for staff who are actually on duty.</td>
<td>- If no, what else do you need?</td>
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<tr>
<td>F140</td>
<td>There is a registered nurse on the day tour of duty 7 days a week (for SNF only).</td>
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<td></td>
<td>Intent</td>
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<tr>
<td>Survey Area</td>
<td>Observation</td>
<td>Interviewing</td>
<td>Record Review</td>
<td>Evaluation Factors</td>
<td>Cross Reference</td>
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<tr>
<td>Patient Care Management</td>
<td>Observe resident level of physical, mental, emotional and social functioning. Note problems, potential problems, needs, using observation/ interview/record review work sheet.</td>
<td>Ask Resident: - Are you aware that you have a plan of care? - Did you participate in developing a plan of care? - Do you know what the plan is and what it entails? (e.g., diet, ambulation, dressing, etc.) - Do you attend and participate in plan of care meetings? - Who else attends the plan of care meetings? - When did you last attend the meeting for your plan of care? - Does the staff assist you in achieving the goals on the plan of care? If not, who does or why not? - Do you have all necessary assistive devices and equipment? - Is there anything that is not part of your plan of care that you think should be included? What happens if you question any treatment or procedure? Can you give an example?</td>
<td>Review: - Plan of care The content of the plan of care is of primary importance rather than the format. Separate care plans are not required for each discipline, but may be acceptable if there is evidence that the various disciplines coordinate their planning. - Nursing assessment/record and notes. - Physician orders. - Physician notes. Assessments/evaluations and progress notes from all professional disciplines as appropriate. - Medication and treatment records as applicable. - Lab reports, as applicable.</td>
<td>- Are all resident's needs/problems identified? - Is the plan developed to meet these needs? - Does the plan demonstrate an interdisciplinary approach, and include: - Goals stated in measurable/observable terms? - Approaches (staff action) to meet the resident action goals? - Responsible disciplines/staff responsible for approaches to assist resident in achieving goal/goals? - Is plan being reassessed and adjusted as needed to reflect current status? - Does plan of care accurately reflect information gained from observation, interview and record review?</td>
<td>(485.1123)</td>
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<td>Medical Records</td>
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<td>Resident Rights</td>
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<td>24 Hour Nursing Service</td>
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<td>Specialized Rehabilitation Services</td>
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<td>Training</td>
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<td>Resident Rooms</td>
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<td>Infection Control</td>
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## Long Term Care Survey

<table>
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<tr>
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</tr>
</thead>
</table>
| F 170 (cont'd) | Ask Staff:  
- What is your input into resident's plan of care?  
- What aspect of the resident plan of care are you carrying out?  
- What is this particular resident's plan of care?  
- How do you assist the resident in carrying out the plan of care?  
- Who attends the care planning meeting?  
- Is the plan of care useful to you in caring for the resident?  
- Is there anything the resident needs that is not addressed in the plan of care?  
- How often is it reassessed? | | | | Social Services  
405.1333  
405.1330(a)  
442.344(d)  
Activities  
405.1331  
442.345  
Dietetic Services  
442.1135  
442.332 |
### Long Term Care Survey

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| **Restorative Nursing Activities of Daily Living**
| F331-176 | SNF 405.1104(e) | ICF 442.342 | 442.342(3)(e)(c) |

**INTENT**

To assist the resident to attain or maintain his/her maximum level of independence and function.

<table>
<thead>
<tr>
<th>A. Observe residents in need of assistance.</th>
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<tbody>
<tr>
<td>1. Is needed assistance provided?</td>
</tr>
<tr>
<td>2. Is resident provided assistance and instruction, as appropriate, in all ADL's to increase his/her level of independence?</td>
</tr>
<tr>
<td>3. Does staff minimize pain/discomfort while assisting resident?</td>
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<tr>
<td>4. Is resident taught transfer techniques?</td>
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<tr>
<td>5. Is resident assisted to toilet in timely manner?</td>
</tr>
<tr>
<td>6. Resident personal equipment available &amp; within reach?</td>
</tr>
<tr>
<td>Glasses</td>
</tr>
</tbody>
</table>

**Ask Resident:**

- What assistance do you need with bathing and/or dressing? Who helps you?
- Does the staff plan with you your dressing/bathing schedule?
- Do the nursing and activities staff coordinate your schedule so that you have the opportunity to participate in favorite activities?
- Are you able to dress/bathe at time convenient for you?
- Are you bathed consistently? (i.e., on the days scheduled does the bath get performed?)
- Where are you bathed? (bed, shower, tub?)
- Are there adequate clothes available for you to wear?
- Do they come back from laundry in appropriate condition?
- How do you get in and out of bed?
- If staff assists you, do they seem to be able to do their job appropriately? Do you always feel safe when

**Review:**

- Plan of care
  - Reflects assessment, goals, methods to reach goals, service providers, evaluation, and achievement.
  - Addresses restorative nursing assessment, program initiation, implementation and evaluation of the progress over a reasonable time period.
  - Professional judgement determines the assessment of appropriate time frames.
  - Identifies planning for potential discharge for all residents to determine a disposition on home care or an alternate level of care.

**Nursing Notes**

- Demonstrate evidence of assessment, intervention, response to treatments/teaching, and their progress toward independence, a maintenance level or a deterioration.
- Provide evidence of interdisciplinary conferences.

**Are patient needs identified?**

- Verify that the plan of care addresses resident needs and is implemented as scheduled and that all appropriate information is documented.
- If goals are not reached, has a reevaluation been performed and goals revised?
- Does restorative nursing assist the resident to acquire a higher level of independence?
- Is sufficient time allowed to resident for learning to increase his/her level of independence?
- Are assistive devices used regularly as per plan and are they in good repair?
- Is there an assessment, and if appropriate, a plan for each ADL that the resident needs to gain independence in?
- Maintenance goals should be noted as appropriate.
<table>
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</thead>
<tbody>
<tr>
<td>F171-176 (cont’d)</td>
<td>Prosthetic devices (e.g., braces, artificial limbs)</td>
<td>Are staff members encouraging you to do things for yourself?</td>
<td></td>
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<tr>
<td></td>
<td>(e.g., built-up spoons, reachers)</td>
<td>Do you have any problems getting to the bathroom on time?</td>
<td></td>
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<tr>
<td></td>
<td>Orthotic devices (e.g., splints, AFO’s)</td>
<td>Do you have any problems with leakage when you sneeze, laugh, or at any other particular time?</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>Restraints (e.g., vest, wrist, ankle, mitts, nets, geri-chairs)</td>
<td>How does the staff help you with these problems?</td>
<td></td>
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<tr>
<td></td>
<td>Grooming items (e.g., comb, brush, shaver)</td>
<td>Are they aware of the problems?</td>
<td></td>
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<td></td>
<td>Oral hygiene (e.g., toothbrush, toothpaste, mouthwash, denture cup)</td>
<td>Do you bowl’s move regularly?</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>Self-feeding devices</td>
<td>If not, what do you/your staff do about this?</td>
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<tr>
<td></td>
<td>Assistive devices for special sensory loss needs (e.g., communication boards, large print books, magnifiers, writing tablets, picture cards, talking books)</td>
<td>Are you able to feed yourself?</td>
<td></td>
<td></td>
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<td></td>
<td>Training/re-training</td>
<td>Are you able to get to the dining room by yourself? If not, why? In that case, what does your staff do about this?</td>
<td></td>
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<tr>
<td></td>
<td>Prosthetic management Stroke adapted A0L’s</td>
<td>How long have you been up today?</td>
<td></td>
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<tr>
<td></td>
<td>Self-injections of medications</td>
<td>Do you usually lie down for a rest?</td>
<td></td>
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<tr>
<td></td>
<td>Bowel/bladder</td>
<td>If you need help getting into or out of bed, is staff available to help you when you need it?</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>Self-feeding</td>
<td>Where do you spend most of your time - in your chair, wheelchair or in bed?</td>
<td></td>
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<tr>
<td></td>
<td>Self-grooming</td>
<td>being helped?</td>
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<td></td>
<td>Ambulation</td>
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<tr>
<td>F171-176 (cont'd)</td>
<td>Cuts/groats</td>
<td>Does anyone move your arms or legs or help you with exercises?</td>
<td>- Have your sleeping habits changed since you came to the nursing home? If yes, in what way?</td>
<td>- Are you able to get help during the night if needed?</td>
<td>- Is staff response timely?</td>
</tr>
<tr>
<td></td>
<td>Oxygen inhalation</td>
<td>- Do you feel confident to assist you?</td>
<td></td>
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<tr>
<td></td>
<td>Speech</td>
<td>- Is your family involved in assisting you or if learning to help you?</td>
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<tr>
<td></td>
<td>Mobility</td>
<td>- Do you feel there is adequate staff at this facility?</td>
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<tr>
<td></td>
<td>Upper extremity dressing</td>
<td>- If not, can you give me an example of why you feel this way?</td>
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<tr>
<td></td>
<td>Lower extremity dressing</td>
<td>- Is your family involved in assisting you or if learning to help you?</td>
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<td></td>
<td></td>
<td>- Do you feel there is adequate staff at this facility?</td>
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<td></td>
<td></td>
<td>- If not, can you give me an example of why you feel this way?</td>
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<td></td>
<td></td>
<td>- Does staff assist and/or encourage activities (e.g., R.D.H., ambulation ADL, communication program, feeding)?</td>
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<td></td>
<td></td>
<td>- How often does staff assist in activities?</td>
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</tbody>
</table>
| | | - Is there anything resident would like to do?
## LONG TERM CARE SURVEY

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<tr>
<td>F171-176 (cont'd)</td>
<td>- Does he/she know why he/she is in a chair?</td>
<td>- Is resident comfortable to use bathroom?</td>
<td>- Does he/she see therapist? (O.T., Speech, P.T.) and how often?</td>
<td>- Does resident go to a</td>
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</table>

*Ask Activities Staff*
- Do you provide information to nursing staff about time and place of activities, plus names of residents who are to attend or those who might be interested in attending?

*Chair-bound Resident*
- Ask Resident:
  - Does he/she know why he/she is in a chair?
  - Is resident assisted to use bathroom?
  - Is resident comfortable?
  - Does he/she see therapist? (O.T., Speech, P.T.) and how often?
  - Does resident go to a
<table>
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</thead>
<tbody>
<tr>
<td>F171-176 (cont'd)</td>
<td>- therapy area or does therapist come to resident?</td>
<td>- Is able to reach items needed?</td>
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<td></td>
<td>Ask Nurse's Aide</td>
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<td></td>
<td>- Who gives you information about the time and place of activities and which</td>
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<td></td>
<td>residents are to attend?</td>
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<td>- How are you given this information?</td>
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<td>- How do you encourage a resident to do the most for themselves?</td>
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<td></td>
<td>Wheelchair Resident</td>
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<td>Ask Resident</td>
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<tr>
<td></td>
<td>- Does he/she know why he/she needs a wheelchair?</td>
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<td>- Is resident trained and/or encouraged in independent W/C ambulation and</td>
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<td></td>
<td>activity?</td>
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<td></td>
<td>- Does resident know how to lock and unlock wheelchair?</td>
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<td></td>
<td>Ask Staff</td>
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<td>- How is the resident set up for independent C/W ambulation?</td>
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<td>- Nurse Aide</td>
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<td></td>
<td>- Has resident received instruction in transfer techniques?</td>
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<td>for Bed Bound Resident</td>
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<td>In addition to appropriate interview questions above</td>
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<td>SURVEY AREA</td>
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<tr>
<td>F171-176 (cont'd)</td>
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</table>

**Ask Resident:**
- How do you spend your day?
- Can you do some things for yourself?
- Does the staff give you a chance to learn self-care skills?

**Ask Nurse:**
- If the resident had access to a recliner chair, would he/she be able to be out of bed?
- Is the time out of bed coordinated with the activity schedule and necessary care?

**Ask Nurses' Aide:**
- Does this resident do any self-care? Why not?
- If no, has anyone tried to teach him/her to do some care?
LONG TERM CARE SURVEY

<table>
<thead>
<tr>
<th>SURVEY AREA</th>
<th>OBSERVATION</th>
<th>INTERVIEWING</th>
<th>RECORD REVIEW</th>
<th>EVALUATION FACTORS</th>
<th>CROSS REFERENCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positioning</td>
<td>Observe residents in bed, chairs, restrained, or in &quot;protective devices&quot; for body alignment</td>
<td>Ask Resident: How often are you turned/repositioned by the staff? Is that often enough? Are you comfortable now? Do you have any pain or discomfort? Where? How long have you had joint stiffness (contractures)? What kind of exercise do you do every day, including range of motion (ROM)? How long does the exercise last and how frequently do you exercise each week? Do you wear special devices? How often? Consistently? Are they always applied and removed appropriately and promptly? How often? By whom?</td>
<td>MD orders for non-mq interventions/treatments. Plan of care should include at least one minimum: Restorative goals. Specific joint to be exercised. Frequency of treatment or repositioning. Frequency of treatment or repositioning. Resident teaching information. Services responsible for carrying out the procedures. Time frames for reaching goals. Nursing notes indicate: Progress toward goals. Response to information from revamping. Look for actual turning/repositioning schedule.</td>
<td>Plan of care should be complete (addressing resident positioning needs) and plan is implemented on a daily basis. Caregivers are knowledgeable re plan content. Residents are turned as scheduled. In good body alignment with proper assistive devices &amp; equipment. Contractures are prevented and/or treated. Plan is reviewed, reevaluated and revised at least quarterly, but must be done as often as patient condition dictates. Ask aide assigned to demonstrate the hand holds he/she uses for ROM. If aide doesn’t know, ROM is probably not being done. Do it &quot;at bath time&quot; is not sufficient.</td>
<td>Rehabilitative Services 485.1124(e) 442.343(c)(2) MD Orders Activities Resident Rights Nursing-Staffing Inservice Social Service Dietary</td>
</tr>
</tbody>
</table>

Intent:
To assure that the resident is positioned at all times to promote maximum therapeutic benefit and comfort, as well as safety.

Specific Observations for the Bed Resident (as appropriate to condition):
- Positioning
- Resting splints & correct application
- Foot positioning boards
- Tapes
- Hand rolls
- Elbow/leg splints & correct application
- Restraints
- Siderails (padded)
- Special mattresses

Special Notes:
- MD orders for non-mq interventions/treatments.
- Plan of care should include at least one minimum:
  - Restorative goals
  - Specific joint to be exercised
  - Devices to be used in positioning
  - Frequency of treatment or repositioning

Plan of care should be complete (addressing resident positioning needs) and plan is implemented on a daily basis. Caregivers are knowledgeable re plan content. Residents are turned as scheduled. In good body alignment with proper assistive devices & equipment. Contractures are prevented and/or treated. Plan is reviewed, reevaluated and revised at least quarterly, but must be done as often as patient condition dictates. Ask aide assigned to demonstrate the hand holds he/she uses for ROM. If aide doesn’t know, ROM is probably not being done. Do it "at bath time" is not sufficient. 

Rehabilitative Services 485.1124(e) 442.343(c)(2) MD Orders Activities Resident Rights Nursing-Staffing Inservice Social Service Dietary

§488.115
### Long Term Care Survey

<table>
<thead>
<tr>
<th>Survey Area</th>
<th>Observation</th>
</tr>
</thead>
</table>
| F175 (cont'd) | Blankets/pillows  
Clean, smooth linen  
Clean, appropriate bed wear  
Turning schedules  
N.B. schedule  
O.B.B. (as tolerated)  
Water available  
All adaptive devices are clean and in good repair.  
All assistive supportive devices are clean and in good repair.  
**Specific Observation for the Bed Resident in Chair** (e.g., chair, lounge chair, in room, as appropriate to condition)  
Arrangement of room facilitates residents optimal independence (e.g., independent eating, grooming, I.V., radio, water).  
Positioning/body alignment.  
Blankets/blanket, pillows, foot stool.  
Hand rolls, splints.  
Clean, dry attire.  
Pressure relief device.  
Restraints, with release & activity schedule.  
Call bell available. |  
| Interviewing | - When?  
- Does staff answer call bell promptly? How soon?  
- Is resident able to reach items (e.g., water call bell, urinal, emesis basin, tissues)?  
- How much confidence do you have when the nurses are helping you transfer, or turn and so on?  
- Does resident go to therapy area or does therapist come to resident?  
**Bed Resident ASK Staff**  
- How often is position changed?  
- What activity is done at the time (e.g., A.M., I.M., toileting, O.B.B., grooming)?  
- What can resident do independently?  
- Is equipment available?  
- Who maintains and cleans the equipment?  
- What is the schedule for this?  
- What training have you had to learn to position patients correctly? |  
| Record Review |  
| Evaluation Factors |  
| Cross Reference |  

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42 CFR Ch. IV (10–1–00 Edition)
<table>
<thead>
<tr>
<th>SURVEY AREA</th>
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<tbody>
<tr>
<td>F175 (cont'd)</td>
<td>Specific Observation for the Wheelchair Resident (as appropriate to condition, including deliberate alterations made to equipment for specific reasons.)</td>
<td>- Proper fit &lt;br&gt; - Good working condition &lt;br&gt; - Appropriate arm rest, footrest, leg support, lap tray &lt;br&gt; - Proper positioning &lt;br&gt; - Pressure relief aids, (e.g., gel flotation pads, egg crate mattress, sheepskin) &lt;br&gt; - Set up for independent WC ambulation &lt;br&gt; - Functional adapted toilet area &lt;br&gt; - Transfer techniques &lt;br&gt; Observe how staff wheel the resident (e.g., do they inform before starting movement)? Are patients moved wheeling forward and facing elevator doors? Observe staff for: - verbal cues &lt;br&gt; - physical support &lt;br&gt; - body mechanics</td>
<td>- Was there any part of your orientation when you first came to work here that addressed positioning? &lt;br&gt; - Do you have any periodic reviews/updates on positioning?</td>
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<tr>
<td></td>
<td>Specific Observation for the Ambulatory Resident (as appropriate to condition)</td>
<td>- Gait (steady/unsteady) &lt;br&gt; - Appropriate devices for</td>
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<tr>
<td>SURVEY AREA</td>
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<tr>
<td>F175 (cont'd)</td>
<td>ambulation (e.g., cane, crutches, hemi-sling - Posture - Appropriate staff assistance in ambulation - Grab bars (hall, bath/shower area) - Functionally adapted toilet area</td>
<td>you deal with it? - Is there something resident would like to do that he/she is not allowed to do (e.g., shave self, apply make-up, style own hair)? - What training have you had in learning to position residents and do range of motion? - What opportunity do you have for ongoing training? - Who does the actual training? Check question placement under Interviewing. May be more appropriate for resident’s rights section. Observe wheelchair technique used by staff.</td>
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### Nursing Services 8. Administration of Drugs

- **Observation:** Observe a drug pass with at least 20 residents receiving medication. See 406 Appendix N, Transmittal No. 174 for details of the Surveyor Methodology for Detecting Medication Errors.

- **Ask Resident:** Do you always receive your medication on time? - If not, what is the problem? - Do you receive the correct medication? - What does it look like? - Who explained your medications to you? - What reactions do you have? - What happens if you have a question or refuse to take your medication? - Who gives you your medication? - Do your medications change in appearance?

- **Review the medication administration record.** (as appropriate) See 406 Appendix N, Transmittal No. 174 for details of the record review.

- **If the combined total of significant & non-significant errors is 5% or above, a deficiency is present.** Any significant error is cause for a deficiency. See Appendix N for details.
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</table>
| F187        | 2. Drugs and biologics are administered as soon after doses are prepared. | - Do the nurses stay with you when you take your medication?  
- Do any of the medications bother you?  
Ask Staff:  
- Do you generally have available the medications you need?  
- Are there any problems in administering medications?  
Note drug doses refused by resident and how handled by staff. | | | |
| F188        | 3. Administered by same person who prepared the doses for administration except under single unit dose packet distribution system.  
Exception: ICF residents may self-administer medications with their physician's permission. | | | | |
## Long Term Care Survey

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<th>Evaluation Factors</th>
<th>Cross Reference</th>
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<tr>
<td>H. Conformance with Physician Drug Orders</td>
<td>Combine with observation of drug pass.</td>
<td></td>
<td>- Review the latest recap of the physicians orders</td>
<td>See Appendix N for details</td>
<td>Physician Services 485.112-3(b)(1)</td>
</tr>
</tbody>
</table>

### Intent
All residents receive medications as ordered by the physician.
<table>
<thead>
<tr>
<th>Survey Area Cross Reference</th>
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<th>Record Review</th>
<th>Evaluation Factors</th>
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</thead>
<tbody>
<tr>
<td>Dietetic Services (Condition of Participation)</td>
<td>Specific Observations which might be indicative of possible nutrition problem: Clinical - underweight/overweight - dehydration - anemia - cracked lips - palor - dull or dry hair - swollen or red tongue - bleeding gums - decubitus ulcers - infections</td>
<td>Ask dietitian to explain the procedure for making substitutions and recording the changes. If menu usually followed? Ask Resident: 1. How are your meals? 2. Are there foods you are not allowed to have? 3. Are you on a special diet? 4. Do you receive foods that are not appropriate for your diet? If so, what do you and the staff do about that? 5. What time do you receive breakfast, lunch and supper? Do you always receive a meal at mealtime? If not, why? What happens then? 6. Do you like the taste of the food? 7. Is the temperature appropriate (i.e., milk chilled, coffee hot, etc.)? 8. Do you get enough to eat? What do you do if you're still hungry after a meal?</td>
<td>Review Nutrition assessment for the following documentation: o Usual/ideal body weight/height o Dietary allergies/constitutions, ability to chew and swallow regular foods without difficulty o Full or partial dentures o Mental and emotional condition o Physical appearance, skin condition o Appetite and food preference o Vitamin and mineral supplements, etc. o Food and fluid intake in measurable terms and frequency of meals o Degree of assistance needed in eating, related mobility, vision, or other identified problems o Medications (e.g., diuretics, insulin, antibiotics, etc.) o Related laboratory findings (e.g., fasting blood sugar, cholesterol, sodium, potassium, hemoglobin, BUN, serum albumin, transferrin or creatinine-height index if available).</td>
<td>o Are physician diet orders followed? o Did nursing plan for feeding and assistance at mealtime? o Is there rehabilitative use of assistive devices, if appropriate? o Is modification of consistency of meals made if resident has a problem or change in condition? o Are between meal and bedtime snacks provided as needed? o Is socialization at meals provided? o Has dietitian provided counseling of resident and family as needed? (related to diet)? o Usual body weight is maintained/supported? o Is there evidence that the plan is being carried out (e.g., documentation in the resident's chart, observation by the surveyor and resident/staff interview)? If the resident refuses meals or does not respond to intervention, the notes in the chart should indicate efforts to intervene or provide counseling.</td>
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<td>F193</td>
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<td>SNF (405.1125)</td>
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<td>A. Menus and Nutritional Adequacy</td>
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<td>F194</td>
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<td>SNF (405.1125(b))</td>
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<td>F194</td>
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<td>ICF 442.332(a)(1)</td>
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<td>F196</td>
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<tr>
<td>Menus are planned and followed to meet the nutritional needs of each resident in accordance with physicians' orders and, to the extent medically possible, based on the recommended dietary allowances of the Food and Nutrition Board of the National Research Council, National Academy of Sciences.</td>
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<td>SURVEY AREA</td>
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<tr>
<td>F196(cont'd)</td>
<td>- Excessive food likes and dislikes</td>
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<td></td>
<td>- Refusal to eat</td>
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<td>- Selected biochemical changes which might indicate changes in nutritional status:</td>
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<td></td>
<td>- Visceral protein status</td>
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<td>- Serum albumin</td>
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<td></td>
<td>- Transferrin</td>
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<td>- BUN</td>
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<td></td>
<td>- Serum electrolytes</td>
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<td></td>
<td>- During mealtime observe the resident for:</td>
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<td>- Adherence to food preferences</td>
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<td></td>
<td>- Adequate space for eating</td>
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<td></td>
<td>- Self-feeding skills</td>
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<td></td>
<td>- Proper position for eating</td>
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<td></td>
<td>- Ability to eat foods served</td>
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<td></td>
<td>- Use of adaptive feeding devices</td>
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<td></td>
<td>- Amount of food actually eaten</td>
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<td></td>
<td>- Protection of resident's clothes</td>
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<td></td>
<td>- Amount of time resident is allowed to chew and swallow</td>
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<td></td>
<td>- Assistance provided as needed to and from dining area</td>
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<td></td>
<td>- All beverages are covered</td>
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<td></td>
<td>- Do you receive nourishment in the evening? Do you have a choice about what you want to eat?</td>
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<td></td>
<td>- Do you receive medications during meals? If yes, do you know what it is or what it is for?</td>
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<td></td>
<td>- Do you get food from outside of facility that you buy or family brings? How often? What kind of food?</td>
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<td></td>
<td>- How often does anyone from the kitchen come to ascertain your feelings and opinions on the food service, your portion size, etc.?</td>
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<td></td>
<td>- Where do you eat (e.g., dining room, your room, etc.)? Is this your choice? Do you have a choice of where you eat?</td>
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<td></td>
<td>- How often have you seen a therapist for your swallowing difficulties? How has the therapist instructed you/staff/family on methods to improve your swallowing?</td>
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<td></td>
<td>- Ask dietitian</td>
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<td></td>
<td>- Describe the meal planning input you receive from residents.</td>
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<td></td>
<td>- Food/drug interactions</td>
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<tr>
<td></td>
<td>- Mental/emotional assessment as it relates to resident's food habits. Review:</td>
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<tr>
<td></td>
<td>- Plan of Care</td>
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<tr>
<td></td>
<td>- Nursing Notes Review:</td>
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<tr>
<td></td>
<td>- Physicians orders</td>
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<td></td>
<td>- Progress notes</td>
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<td>- Notes from other professional disciplines as appropriate.</td>
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<tr>
<td></td>
<td>- Nutritional status depends not only on adequacy of menu planning but also whether the resident eats the food and how the body uses it. While the surveyor is not responsible for individual nutritional assessments of residents, when specific information is needed during the survey to make a compliance decision, the surveyor will utilize the following minimum assessment guidelines: Menu Evaluation:</td>
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<td></td>
<td>- Adequate in energy and nutrients - Protein - Calories</td>
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<td>- Is there evidence that the resident's progress is regularly observed (e.g., awareness of food and fluid intake such as acceptance of foods, food consumed, and resident's appetite)?</td>
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<td></td>
<td>- Is fluid intake for resident encouraged, Foley catheter, problem feeders monitored?</td>
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<td></td>
<td>- Is there general evidence as to whether poor resident conditions are due to poor care or whether the facility has taken appropriate measures to prevent or resolve problems?</td>
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<td>- Is there indication of progress toward desired outcomes? If not, is the evidence of re-evaluation available within specified time frames?</td>
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<td></td>
<td>- When the anthropometric and clinical data do not correlate with dietary data (food intake, dietary supplements) the surveyor should take note that the problem may not be nutritional.</td>
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| F196(cont'd) | Assistance being provided in case of choking, incontinence, falling, or other emergencies. Nursing Staff supervision of dining areas including residents' rooms during meal times. | | | - Vitamin C  
- Calcium | |

Selected evaluation of residents for in depth review:

A checklist can be used to evaluate daily menus for basic foods:

- Use standard serving portions

Daily food plan should include:

**Milk Group**
1 pt milk

**Milk Group**
5 equivalents:* 1 equivalent equals 1 oz. of milk (edible portion) weighed after cooking (this includes eggs, milk, cheese, nuts, and all milk, fish and poultry).

**Vegetable and Fruit Group**
5 servings or more, including a dark green or deep yellow vegetable for vitamin A value every other day and a citrus fruit or other fruit rich in vitamin C daily.

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*Note:* This table is from the Health Care Financing Administration, HHS, and includes guidelines for evaluating residents in long-term care settings.
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<tr>
<td>F196 (cont'd)</td>
<td>Observe serving portions sizes on all menu items:</td>
<td></td>
<td></td>
<td>BREAD-CEREAL-POATATO-LEGUME-PASTA GROUP</td>
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<tr>
<td>MILK GROUP</td>
<td>- 1 pint daily</td>
<td></td>
<td></td>
<td>7 servings</td>
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<td></td>
<td>Source of: Protein Calcium</td>
<td></td>
<td></td>
<td>FATS AND SWEETS</td>
<td>(Without this group the diet contains 1,415 Kcal)</td>
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<td></td>
<td>Phosphorus B Complex</td>
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<td>Diets should be adapted from facility's currently approved diet manual.</td>
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<tr>
<td>MEAT GROUP</td>
<td>- 5 lean meat equivalents (1 meat equivalent = 1 oz meat, poultry, fish, cheese &amp; eggs; also dried peas, beans, and nuts)</td>
<td></td>
<td></td>
<td>Menus are dated and contain minimum portion sizes.</td>
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<tr>
<td></td>
<td>Source of: Protein Iron</td>
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<td></td>
<td>Are substitutions noted on the file copy?</td>
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<tr>
<td></td>
<td>Vitamin B12</td>
<td></td>
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<td>Are substitutions made within the same food group i.e., meat for another source of protein in the meat group, or vegetable of similar nutritional value?</td>
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<tr>
<td>VEGETABLE AND FRUIT GROUP</td>
<td>- 5 servings or more (1/2 cup = 1 serving)</td>
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<td></td>
<td>Source of: Vitamin A, C, K, folate, fiber</td>
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<tr>
<td>BREAD-CEREAL-POATATO-LEGUME-PASTA GROUP</td>
<td>- 7 servings (1 serving = 1 slice bread, 1/2 cup other; 3/4 cup kale-type cereal)</td>
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<tr>
<td>F196 (cont'd)</td>
<td>FATS AND SWEETS (to increase caloric intake)</td>
<td></td>
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<tr>
<td></td>
<td>CODIFIED SALT (unless contraindicated)</td>
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<tr>
<td></td>
<td>Adequate fiber in diet</td>
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</table>

**INTERVIEWING**

- Documentation of decision to withdraw or begin artificial feeding and hydration.
- Check menus for variety.
- Are they specific (i.e., state kinds of fruit, juice, vegetables)?

**RECORD REVIEW**

- Nutritional Requirement: Record review
- N.B. The basal energy expenditure (BEE) and calorie requirement using Harris-Benedict formula recognizes the variation in energy needs for individuals.

1. **Anthropometry - Height/Weight**

**NOTE:** The following simple formulas and guidelines are not the only acceptable guides available. The surveyor should ask to use the assessment guidelines used by the facility before using the ones provided here.

- Important indicator of nutritional outcome.
- Disease state can have adverse effect on desired body weight.
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<tbody>
<tr>
<td>F196 (cont'd)</td>
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</table>

2. **Height for Height Calculation**

   - **Females:**
     - Allow 100 lbs. for first 5 ft. of height plus 5 lbs. for each additional inch.
   - **Males:**
     - Allow 100 lbs. for first 5 ft. of height plus 6 lbs. for each additional inch.

3. **Estimating Caloric Needs**

   - **FORMULA:** Harris-Benedict Equation

   - **Men:**
     - $66.5 + (13.7 \times \text{Wt. in Kg}) + (5 \times \text{Ht. in cm})$
     - $- (6.8 \times \text{Age}) \times \text{BEE}$
   - **Women:**
     - $65.5 + (9.6 \times \text{Wt. in Kg}) + (1.7 \times \text{Ht. in cm})$
     - $- (4.7 \times \text{Age}) \times \text{BEE}$

   - **Parenteral Anabolic:**
     - $1.75 \times \text{BEE}$
   - **Oral Anabolic:**
     - $1.5 \times \text{BEE}$ (Kcal/s)
### Long Term Care Survey

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<tbody>
<tr>
<td>F196 (cont'd)</td>
<td>Oral Maintenance: 1.20 x BEE (kCal/s)</td>
<td>Metric Conversions (Approx): Pounds (lb.) x 0.45 = Kilograms (Kg)</td>
<td>Inches (in.) x 2.5 = Centimeters (cm)</td>
<td>Estimating Protein Needs: 1. Allow 0.8 gram protein per kilogram of ideal body weight. 2. Increase to 1.2 - 1.5 gm/kg for patients with depleted protein stores (decubitus, draining wounds, fractures, etc.). Fluid Requirement: Based on actual body weight: Over 55 years with no major cardiac or renal diseases: (NOTE: 1.2 lbs. equals 1 kg of body weight)</td>
<td></td>
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## Long Term Care Survey

<table>
<thead>
<tr>
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<tr>
<td>F100 (cont'd)</td>
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</table>

### Record Review

**Example:**
- 120 lbs/2.2 lbs. = 54.5 kg (55 kg)
  - 55 kg x 30 cc = 1,650 cc/day

**Note:** Isotonic Standard Tube Feeding:
- Approximately 80% water.

### Evaluation Factors

**Amputation % of Body Weight**
- Leg 20%
- Below Knee 10%
- Arm 6%
- At Elbow 3.6%

**Suggested Standards for Evaluating Significance of Weight Loss**
- % of body weight loss
  - Inter- Significant Severe
  - 1 week 1-2% 2%
  - 1 month 5% 5%
  - 3 months 7 1/2% 7 1/2%
  - 6 months 10% 10%

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</thead>
<tbody>
<tr>
<td>F196 (cont'd)</td>
<td>Lab Indices for Visceral Protein</td>
<td>Mild Deficiency</td>
<td>Moderate Deficiency</td>
<td>Severe Deficiency</td>
<td></td>
</tr>
<tr>
<td>Albumin (g/dL)</td>
<td>3.5-3.2</td>
<td>3.2-2.8</td>
<td>2.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Lymphocyte Count (cells/mm³)</td>
<td>1500-1600</td>
<td>1500-900</td>
<td>900</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Transferrin (If Available)</td>
<td>200-100</td>
<td>100-160</td>
<td>160</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SURVEY AREA</td>
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<tr>
<td>B. Therapeutic Diets</td>
<td>System for the provision of diets:</td>
<td>Ask Staff:</td>
<td>Review:</td>
<td>Nursing Services 405.1124 405.1124(c) Patient care plan (f.) Supervision of patient nutrition</td>
<td></td>
</tr>
<tr>
<td>FIQD</td>
<td>o Dietetic service Kardex or file</td>
<td>o Number, type of therapeutic diets?</td>
<td>- Physician diet orders in medical record</td>
<td></td>
<td></td>
</tr>
<tr>
<td>FINP</td>
<td>o Therapeutic menus</td>
<td>o Time of nourishment activity, who’s responsible?</td>
<td>- Nurses’ Kardex</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ICF</td>
<td>o Nourishment preparation and service</td>
<td>o Nourishment provided for day of survey?</td>
<td>- Dietary Kardex</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ICFI</td>
<td>o Adequacy of nourishment</td>
<td>o Individual menus or diet cards</td>
<td>- Therapeutic diet menu</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ICFI</td>
<td>o Individual menus or diet cards</td>
<td>Note:</td>
<td>- Diet cards</td>
<td></td>
<td></td>
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<tr>
<td>SPOF</td>
<td>o Number of residents having difficulty in speaking or swallowing with the tube in place o Tube feeding feeding - Some residents should be considered for in-depth reviews</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>SPOF</td>
<td>o Tube feeding review</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>SPOF</td>
<td>o Plan of care</td>
<td></td>
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<tr>
<td>SPOF</td>
<td>o Identify frequency, amount of feeding based on the physician’s order and the time span over which each feeding is accomplished</td>
<td></td>
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<tr>
<td>SPOF</td>
<td>o Medication and treatment records</td>
<td></td>
<td></td>
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<tr>
<td>SPOF</td>
<td>o Fluid intake records</td>
<td></td>
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<tr>
<td>SFOF</td>
<td>o Number of calories as planned in writing, prepared, and served as ordered with supervision from the dietitian and advice from the attending physician whenever necessary</td>
<td></td>
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<tr>
<td>SFOF</td>
<td>o Unused milk-based tube feeding should be discarded in a timely manner</td>
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<tr>
<td>SFOF</td>
<td>o Staff use proper technique in administering feedings and medications. Check to see if staff checks for location of tube before feeding and that tubing is irrigated before and after addition of medication.</td>
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</table>

**SPOF:**
- Consider appropriateness of special diet—updated and reviewed since admission.
- Progress notes reflect reevaluation of resident’s progress on diet.
- Selected number of residents on therapeutic diets should be considered for in-depth reviews.

**SPOF:**
- Ordered by physician
- Prepared fresh daily
- Same calories and/or food groups as if served whole

**On Pureed diets:**
- Pureed foods are coordinated with general/regular menu.

**On Tube Feeding:**
- Has the feeding been ordered by physician?
- Is tube feeding nutritionally adequate?
- Have attempts been made to progress tube feeding (if indicated)?
- Have changes in resident condition been noted and addressed?
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<tr>
<td>F197-199 (cont'd)</td>
<td>4. Does the staff help you in feeding? Do you feel comfortable/safe with all the staff who perform the feeding? If not, what happens?</td>
<td>- Well as amount of additional water</td>
<td>- Periodic reassessment of ability to swallow</td>
<td>weight loss, constipation, diarrhea, skin condition?</td>
<td>On Diabetic Diets and Other Therapeutic Diets</td>
</tr>
<tr>
<td></td>
<td>5. Are you losing or gaining weight? What is your goal?</td>
<td>- Record should indicate measures taken to prevent diarrhea and constipation and to treat if they have developed.</td>
<td></td>
<td>o Have observed problems been coordinated with other departments and resolved?</td>
<td>o Varied nourishments as preferences allow?</td>
</tr>
<tr>
<td></td>
<td>6. How often is the tube changed? Who does this? Do you feel comfortable/safe with all staff who perform this procedure?</td>
<td>Interview staff regarding knowledge of diabetic diets.</td>
<td></td>
<td>o Is feeding being monitored to ensure that feeding is occurring at the ordered/appropriate rate?</td>
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<tr>
<td></td>
<td></td>
<td>o What nourishment does the diabetic patient receive?</td>
<td></td>
<td>o Varied, nutritionally adequate</td>
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<tr>
<td></td>
<td></td>
<td>o If diabetic patient refuses the meal, what is done to supplement the meal?</td>
<td></td>
<td>o Individualized to suit resident</td>
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<tr>
<td></td>
<td></td>
<td>If resident is able to be interviewed, suggested questions:</td>
<td></td>
<td>o Re-evaluation indicates diet meets objectives. If not appropriate, documentation is provided</td>
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<tr>
<td></td>
<td></td>
<td>1. How long have you been on your diabetic diet?</td>
<td></td>
<td>o Laboratory results support diagnosis</td>
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<td></td>
<td></td>
<td>2. Do you know some of foods you must avoid? What are they?</td>
<td></td>
<td>o Between meals nourishment provided as needed and recorded in measurable amounts.</td>
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</table>
### Long Term Care Survey

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<tbody>
<tr>
<td>F197-199 (cont'd)</td>
<td>Observe tray/meal service:</td>
<td>3. Do you receive a nourishment between meals or before going to bed?</td>
<td></td>
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<tr>
<td>F198</td>
<td>Therapeutic diets prescribed by the attending physician</td>
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<tr>
<td>F199</td>
<td>Therapeutic menus are planned in writing, prepared and served as ordered with supervision from the dietician and advice from the physician whenever necessary.</td>
<td>Functioning system to provide the needed nutrients:</td>
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<tr>
<td></td>
<td>- Resident's general appearance</td>
<td>- Resident's general appearance</td>
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<td></td>
<td>- Meal service</td>
<td>- Meal service</td>
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<td></td>
<td>- Food acceptance</td>
<td>- Food acceptance</td>
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<td></td>
<td>- Adherence to food preferences</td>
<td>- Adherence to food preferences</td>
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<td></td>
<td>- Food supplement</td>
<td>- Food supplement</td>
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<td></td>
<td>- Type of service</td>
<td>- Type of service</td>
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<td></td>
<td>- Method of service</td>
<td>- Method of service</td>
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<td></td>
<td>- Assistance provided</td>
<td>- Assistance provided</td>
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<td></td>
<td>- Timely provision as ordered</td>
<td>- Timely provision as ordered</td>
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<td></td>
<td>- Portion sizes</td>
<td>- Portion sizes</td>
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<td></td>
<td>- Conforms to physicians orders</td>
<td>- Conforms to physicians orders</td>
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#### For the Resident with Decubitus Ulcers

| Ask Staff | 1. Identify residents with conditions that immobilize or prevent voluntary body movement. | | | | |
| | - What do you do when this resident refuses meals, meals, bread, etc.? | 2. Identify location, number, size and depth of decubitus ulcers. | | | |
| | - What happens when a weight loss is noticed with this resident? | 3. Calculations of kilocaloric and protein levels as needed. | | | |
| | - What is the best diet for this resident? | 5. Progress notes | | | |
| | - How do you monitor weight? | - Monitor weight | | | |
| | - How do you monitor healing of decubitus ulcers? | - Monitor healing of decubitus ulcers | | | |
| | - How do you monitor the condition of your resident? | 6. Pertinent Laboratory Data | | | |
| | - How do we ensure the resident's comfort? | - Hemoglobin/Hematocrit | | | |
| | - How do we ensure the resident's nutritional intake? | - Serum Albumin | | | |
| | - How do we ensure the resident's fluid intake? | - Total Lymphocyte Count | | | |
| | - How do we ensure the resident's hydration? | - Fluid Intake | | | |
| | - How do we ensure the resident's support? | - Sufficient to maintain hydration | | | |

A system is in place to provide the type and amount of nutritional support needed by the residents who have developed decubitus ulcers. Food and supplementation are provided in a method to ensure intake of nutrients needed by residents with decubitus ulcers. Nutritional intervention is assessed and reassessed to ensure appropriate intervention for acceptable health care outcome.

---

Nursing Service

483.1124 (d) Patient Care Plan

(f) Supervision of Patient Nutrition
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<tr>
<td>197-199 (cont'd)</td>
<td><strong>Renal Review</strong></td>
<td>Interview staff regarding knowledge of renal diets: 1. What foods should be restricted when a patient has kidney problems? 2. What nourishments are given to these patients? 3. Are fluids restricted? Ask Resident: 1. Are you on a special diet? 2. What foods must you avoid? 3. Do you feel hungry? 4. Do you eat everything at mealtime? 5. Are the foods the kitchen sends you the correct ones for your diet? 6. Has the dietitian explained your diet to you?</td>
<td>Renal Patient Diet Review - Pertinent Laboratory Data + Serum Sodium + BUN + Serum Potassium + Albumin + Hematocrit + Creatinine - Pertinent Medications + Vitamin/Mineral + Supplements - Weight gains/losses</td>
<td>On Renal Diets - Ordered by physician - Written menu nutritionally complete in so far as medically possible, including calories - Individualized to suit resident - Laboratory testing as needed - Coordination with dialysis unit to determine effectiveness of diet</td>
<td>Nursing Service 195.1124</td>
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<tr>
<td>C. Preparation</td>
<td><strong>F204</strong></td>
<td>SNF 405.1125(a)</td>
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<td></td>
<td><strong>F205</strong></td>
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<tr>
<td>1. Food is prepared by methods that conserve its nutritive value and flavor.</td>
<td>Observe:</td>
<td>Feeding assistance is provided or not provided by staff.</td>
<td>Review:</td>
<td>The facility has kitchen and dietetic service areas adequate to meet the food service needs. These areas are properly ventilated, arranged, and equipped for sanitary refrigeration, storage, and preparation of food. Equipment and storage areas are clean, well maintained, within proper temperatures ranges, and safe.</td>
<td>Proper temperatures:  (Fabriheat) Frozen food storage -- 0 or below Cold food storage -- 40-45 degrees Hot food holding equipment -- 140 degree minimum Dishwasher wash cycle -- 150 - 150 degrees Dishwasher rinse cycle -- 160-180 degrees or a color change in thermopaper; or adherence to manufacturers recommendations</td>
</tr>
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<td></td>
<td><strong>F206</strong></td>
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<tr>
<td>2. Meals are palatable, served at proper temperatures. They are cut, ground, chopped, pureed or in a form which meets individual resident needs.</td>
<td>Observe:</td>
<td>Food is served soon after cooking or refrigerated. Trays are free of spillage of foods or liquids. Foods are appropriately covered and kept at a proper temperature. Cooking and service utensils are clean, sanitary and greaseless. Refrigerated foods must be covered. Leftover and pre-cooked foods must be dated and labeled. All cooked food stored above raw meals in refrigerator. Temperature gauge on or in refrigerator to record temperature. Shelving to allow air circulation. Food not stored in refrigerator must be stored off the floor. (This is applicable to food stored in walk-in refrigerator and freezer.)</td>
<td>Review:</td>
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<td><strong>F207</strong></td>
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<tr>
<td>3. If a resident refuses food served, appropriate substitutes of similar nutritive value are offered.</td>
<td>Observe:</td>
<td></td>
<td>Review:</td>
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<tr>
<td>F207 (Cont'd)</td>
<td>- No rust on shelves - No dripping or spillover on shelves and floors - Degree to which diet modification is commensurate with residents' tolerance and capability - Residents for meal satisfaction - Observe appearance of food color, texture, aroma, and flavor - Less than 75% of meal is consumed - Type of substitutions provided</td>
<td>- Progress notes - Diet card - Day's menu substitute record</td>
<td>Dietary personnel are clean and free of infectious disease. They practice acceptable techniques and procedures to keep foods at proper temperatures and protected against contamination. Is dietary information pertinent to dietary modification? Has resident been assessed for eating program to maintain independence? The food substitute is of similar nutritive value as the refused item (e.g., milk refused, alternate of calcium rich food should be provided.</td>
<td></td>
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<tr>
<td>SNF 495.1125(e)</td>
<td>To provide foods that are safe and nutritious</td>
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<tr>
<td>D. Frequency</td>
<td>o Menus as under A on page 63</td>
<td>Interview various residents about the nourishment service:</td>
<td>o Menu as under A o Nourishment list</td>
<td>Three meals or their equivalent are served daily with not more than a 16-hour span between the evening meal and breakfast. The nourishment service is more difficult to evaluate, must find evidence that patients are offered nourishments on a planned basis and documented.</td>
<td></td>
</tr>
<tr>
<td>F208 SNF 405.1124(d)</td>
<td>o Who serves nourishments o Nourishment list and schedule</td>
<td>o Are nourishments offered routinely? o At what time are they offered? o By whom? o What kind of nourishments are offered?</td>
<td></td>
<td></td>
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<tr>
<td>F209 ICF 442.331(a)</td>
<td></td>
<td></td>
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<tr>
<td>F210</td>
<td>At least three meals are served daily at regular hours with not more than a 16-hour span between a substantial evening meal and breakfast.</td>
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<tr>
<td>F211</td>
<td>In the extent medically possible, bedtime nourishments are offered to all residents</td>
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<tbody>
<tr>
<td>E. Staffing</td>
<td>- Food service personnel are on duty for all defined dietary responsibilities: - Supervision - Food Preparation - Dishwashing - Cleaning</td>
<td>- Interview personnel to verify that they are aware of their responsibilities and job descriptions.</td>
<td>-</td>
<td>- from an assessment of the total dietetic service operation: - The dietetic supervisor is capable of the overall management and supervision of the dietetic service. - There are dietetic personnel on duty over a 12-hour period who demonstrate ability to perform tasks adequately. - Dietetic personnel receive appropriate orientation and training consistent with their duties and responsibilities. There is evidence that the dietetic staff are knowledgeable about food service policies and procedures and apply these accepted professional practices in their daily work. - Services provided are consistent with the size, scope and facilities available.</td>
<td>HHS § 488.115 (a)</td>
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### Intent

Persons are providing services commensurate with their level of training and at the level of sophistication needed by the residents.
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<tr>
<td><strong>Specialized Rehabilitation</strong></td>
<td>Observe residents as per &quot;Restorative Nursing Activities of Daily Living&quot;</td>
<td>Ask resident (or ask staff if resident has severe communication problems):</td>
<td>Review: - Plan of care - Doctors' orders - Nursing assessment and progress notes - Adequate assignment sheets - Therapy assessments/evaluations (includes a minimum of): - name, age, date, diagnoses - Referring physician and reason for referral - History, precautions, limitations - Objective documentation (e.g., tests, measurements) - Rehabilitation potential - Treatment plan (includes a minimum of): - Specific rehabilitation needs and objectives - Treatment to meet specific measurable rehabilitative goals - Type, amount, frequency, duration, modalities - Name of therapist(s) who will provide treatment - Restorative nursing follow-up (recommendations for plan of care)</td>
<td>- Are rehabilitation services integrated with restorative nursing? - Do therapists participate in development of resident plan of care? - Do observations and interview indicate that services are provided in conjunction with 24-hour nursing, and in accordance with the overall plan of care regarding restorative nursing and specialized rehabilitative services?</td>
<td>- Nursing Services 442.338 - 442.339 - 442.341 - Physician Services 442.346 - Medical Records 442.345 - Activities Program 442.347 - Resident Rights 442.311</td>
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<tr>
<td>F214 SNF 405.1120</td>
<td>Observe residents in therapy areas: - Is privacy provided during treatment, as applicable (e.g., cubicle curtains, room dividers, one to one areas?) - Is there appropriate, courteous resident/staff interaction? - Are therapy areas appropriate to treatment goals (e.g., small, quiet area for speech therapy/auditory testing and sessions, large for PT, exercise and therapy groups, OT, perceptual testing/splinting, A.D.L. adaptations, area, as applicable)? - Is equipment clean and in good working condition? Is it operating as per manufacturer's instructions (e.g., hydrocollator temp., paraffin, whirlpool, etc.)?</td>
<td>- How many days/hours per week do you provide therapy? - Do you participate in the development of the resident's overall plan of care? In what way? - Do you utilize P.T.?</td>
<td>- Therapy start: - How many days/hours per week do you provide therapy? - Do you participate in the development of the resident's overall plan of care? In what way? - Do you utilize P.T.?</td>
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<td>- Therapy start: - How many days/hours per week do you provide therapy? - Do you participate in the development of the resident's overall plan of care? In what way? - Do you utilize P.T.?</td>
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<td>F21B (cont')</td>
<td>professional practices by qualified therapists or qualified assistants.</td>
<td>&quot;Aides&quot;. In what way (if interviewing the registered physical therapists)? How do you assure carry-over of therapists in your absence? How often do you provide inservice to staff? What topics are covered? Do you have opportunities to attend inservices? How do you communicate patient progress/regression, etc. with physician, nursing personnel, family, other disciplines? How many residents currently are receiving P.T., O.T., Speech-language pathology and audiology therapy (S/P/A)? Do you utilize the services of a certified occupational/therapy assistant (if interviewing the registered occupational therapist)? If so, in what way? Is there space available for the conduct of your therapy? Is equipment readily available to meet resident needs? Is there a coordinated interdisciplinary service?</td>
<td>- Are assistive devices being provided as needed? - Do assistive devices fit well, function and are used properly (e.g., wheelchairs, slings, braces, gaiters, hearing aids, canes, artificial limbs, assistive eating devices)? - Is staff responsive to resident expressions of discomfort?</td>
<td>- Identifies modalities that will be delegated to non-skilled staff</td>
<td>Physical Environment 405.1134 442.324 442.325 442.326 442.328 442.329 442.338 Dietetic Services 405.1125(e) 442.329 442.331(c)</td>
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<td>C. PROGRESS</td>
<td>ICF 442.3451(f)</td>
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LONG TERM CARE SURVEY

| SURVEY AREA | OBSERVATION | INTERVIEWING | RECORD REVIEW | EVALUATION FACTORS | CROSS REFERENCE |
|-------------|-------------|--------------|---------------|-------------------|----------------|}

F220
2. The resident's progress is thereafter reviewed regularly and the plan of rehabilitative care is re-evaluated as necessary. But at least every 30 days by the physician and therapist.

<table>
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<td>If resident's plan must be revised as necessary</td>
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<th>INTENT</th>
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<td>Therapy services are provided that will assist the resident to attain his/her optimal level of function.</td>
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### Long Term Care Survey

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<tr>
<td><strong>Pharmaceutical Services</strong></td>
<td>- Observe residents for excess sedation or adverse effects: drooling, shaving, gait, involuntary movements of limbs, tongue, facial muscles, loss of affect, drowsiness, postural abnormalities, pill rolling movement, observe for depression, agitation.</td>
<td><em>Ask Resident:</em> - Are you aware of the medications you are taking? - Are you on the drug regimen prescribed by the doctor? - Has your physician discussed the medications you are taking? - How many medications are you taking? - How do you feel the medication helps you? - How do medications bother you? (e.g., make you feel nauseated or dizzy?) - Have you told anyone about this?</td>
<td>Review medical record: - to see if pharmacist or nurse has reviewed a drug regimen on a monthly basis. - for evidence that the reviewer has reported irregularities to the physician or another who has authority to correct the irregularities for evidence that the irregularities have been evaluated. - review nurses notes, progress notes, care plan, etc. for any adverse reaction to medication and indication that corrective action was taken. - screen the drug therapy of the residents included in the sample using the indicators (if prepared) outlined in SOM Appendix N Transmittal 4174. - review pharmacists drug regimen monthly reports to determine if pharmacist has corrected any potential irregularities, screened out through this process (need full year).</td>
<td>Reviews were performed in the facility. There was evidence of a review performed on every resident whose record was reviewed indepth. In records reviewed, the average prescription utilization was not substantially over 6.1. If it is, review for appropriateness. Apparent irregularities were identified and reported. <em>Refer to SOM Appendix N in 4174 for further information on drug regimen review.</em></td>
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<td>F221</td>
<td>SNF 405.1127(a)</td>
<td>A. Supervision</td>
<td></td>
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<td>Physicians Services 405.1123(b) 442.336</td>
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<tr>
<td>F222</td>
<td>SNF 405.1127(a)</td>
<td>The pharmacist reviews the drug regimen of each resident at least monthly &amp; reports any irregularities to the medical director and administrator.</td>
<td></td>
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<td>Nursing Services 405.1124 442.338</td>
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A registered nurse may be utilized to perform this monthly review for ICT residents. Also the attending or staff physician must review medication quarterly.
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<tr>
<td>Laboratories and Radiological Services</td>
<td>Observe symptoms of targeted residents, e.g., drainage, odors, jaundice, fevers, edema, etc.</td>
<td>Ask Nursing/Rehabilitative Staff: - What do you do when you think a resident needs laboratory work done - blood work, cultures, etc.? - How long does it take to get lab results back? - What do you do with the results when they do come back? - Do you have any problems with your laboratory services? - How are lab specimens stored? - Do you have any instruction from the lab regarding collection and storage of specimens?</td>
<td>Review the physician's order sheet to see if: - Orders for lab services are signed - That there are orders for tests that have been done. Nursing progress notes are reviewed for documentation of physician notification of lab results. Physician progress notes or other documentation indicating that the physician is aware of lab results. There are lab reports on the medical record for all tests ordered (except if just performed).</td>
<td>There must be signed physician orders for all lab radiology services performed. Record results of all testing in the medical record. There is documentation in nursing or physician notes to indicate the results of lab tests were promptly communicated to the physician. When lab tests are performed the resident should be informed of significant findings and the possible therapeutic alternatives.</td>
<td>Nursing Services 489.1124(a)(b)(c) 442.345 Physician Services 485.1123(b)</td>
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<td>F232</td>
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3. Signed and dated reports of a clinical laboratory, x-ray and other diagnostic services are filled with the patient's medical record.

**INTENT**

To assure that lab tests are performed as ordered and findings are reported to physicians are made aware of symptoms that may require lab tests.
## Survey Area

### Social Services

**F231**

SNF 405.1130

- Behavioral exhibited (dis-orientated, confused, un-cooperative, agitated, withdrawn, isolated, lonely)
- Personal appearance
- Apparent disabilities
- Apparent vision and hearing problems

**F234**

SNF 405.1130(d)

- A. Plan

**F235**

ICF 442.344(d)

- B. Provision of Services

**F237**

1. Services are provided to meet the social and emotional needs of the facility or by referral to an appropriate social agency.

### Observation

**ON**

- Observe resident for:
  - Level of alertness
  - Behavioral exhibited (dis-orientated, confused, un-cooperative, agitated, withdrawn, isolated, lonely)
  - Personal appearance
  - Apparent disabilities
  - Apparent vision and hearing problems

- Talk to staff, other residents, family, visitors
- Participation in group activities
- Independence in activities, decision making

**INT**

- Therapeutic staff intervention: constructive reaction to resident's behavior
- Resident's participation on policy-making bodies and committees of facility, e.g., resident councils.

### Interviewing

- **F231**
  - How long have you been in the facility?
  - Can you explain to me why you are here?
  - Have you had any problem adjusting to the facility, i.e., loss of independence?
  - Have you had any other problems?
  - Have staff been helpful, e.g., financial?
  - Do you have any family or any other visitors?
  - Do you have any problems with which this facility has not been helpful?
  - If exhibiting disruptive behavior, agitated, enervous, etc., how? Do you notice that you are upset? Quiet, nervous, unhappy today? Can you tell me what has bothered you?
  - Does staff respond to your suggestions about your own care?
  - Did you participate in planning what care you will receive and who will give it to you?
  - Do you make use of the dining, activity, community room, and/or outdoor area?

### Record Review

**R2**

- Review medical records of residents selected for in-depth review to determine that:
  - Assessment and plan of care identifies residents medically related social and emotional needs and/or problems.
  - Resident's family and home situation, information related to medical and nursing requirements, and community resources are considered in making decisions regarding the resident's care.
  - Medical records contain current specific information that highlights the social and emotional needs of the resident and significant findings and actions are entered promptly in the medical record.

### Evaluation Factors

- The residents' social and emotional needs are identified, the plan of care addresses these needs.
- The plan of care is being followed, reviewed, and revised as necessary. The family's needs and concerns are addressed if applicable. There is referral to appropriate agencies if necessary. Sufficient space is provided for private meetings and discussions.
- While it is not a program requirement, a social worker or other staff may contribute to the resident's care plan by identifying personal strengths that can be used to build upon.

### Cross Reference

- **Nursing Services**
  - SNF 405.1130
  - ICF 442.344(d)

- **Activities**
  - SNF 405.1133
  - ICF 442.344(a)(c)(d)

- **Physician Services**
  - SNF 405.1131(b)
  - ICF 442.344(c)

- **Patient Care Management**
  - SNF 405.1131(d)
  - ICF 442.344(d)

- **Physical Environment**
  - SNF 405.1130(b)
  - ICF 442.344(c)
### Survey Area: F233-21B (cont'd)

#### 2. If financial assistance is indicated, arrangements are made promptly for referral to an appropriate agency.

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| F233-21B    | - Can you tell me about your life here? What do you do in a usual day? | - Ask Social Work/Nurse: When the social worker is readily available, delete "ask the nurse.
- How often is the resident seen by a social worker? | - Plan of care, social service notes, reflect the current status of the resident. | - There is evidence that the resident needs mental status has been considered when plan of care was developed. | |
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| F233-238 (cont'd) | - How is physician notified and involved in plan of care?  
- Ask social service staff about their role, function, and services they provide.  
- Ask staff what referral services are available.  
- If services are being provided by outside resource, are resources documented in work service reports?  
- Ask social service staff about their background and education.  
- If there is a consultant, ask staff:  
  - How often does the person come?  
  - How long do they stay?  
  - What does the person do while in the facility?  
  - What assistance, consultation is being provided?  
- Ask social service staff if adequate space is provided for them by the facility to conduct private interviews and meetings. | - The time period between date of referral and date of services is reasonable and if not, there is evidence of follow-thru by staff.  
- The outside agency has documented their involvement and activities.  
- Plan of care demonstrates awareness of behavior, articulates the reasons for it, and indicates in the plan of care an approach to the behavior.  
- Assessment should contain:  
  - A flexible approach to each resident (should be individualized),  
  - Awareness of a mental status evaluation.  
  - Resident history.  
  - Family availability for planning, resident support, etc.  
  - Identification of problems resulting from placement.  
  - Recent social adjustment.  
  - Discharge planning.  
- The record reflects | - There is documentation of collaboration between nursing and social work for meeting emotional needs. | Patient Care Management: 405.12A(d) |
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<td>F233-238 (cont'd)</td>
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<td>Social Service intervention with family and resident, i.e., grief and bereavement counseling.</td>
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<td>Review integrated plan of care for: - Family for concerted social services. - Plan for supportive services for adjustment. - Adjustment goals. - Interventions for specific conditions.</td>
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### Activities

| F239 | SNF 405.1131 | | | | |
| F240 | SNF 405.1131(b) | | | | |
| F241 | ICF 442.345 | | | | |
| F242 | | | | | |

1. An ongoing program of meaningful activities is provided based on identified needs and the level of activity of the residents, as well as in specifically designated areas.

- How many residents are lying on their beds or sitting in chairs staring at the walls during waking hours?
- What is the level of residents' interest in activities they are doing?
- Are residents positioned correctly for activity?

### Activities Assessment

- Interests of the resident (past and present) are identified as to resident's current capabilities and necessary adaptations to pursue their interests.

### Activities Intervention

- Are each resident's personal interests known? If not, what actions are being taken to identify them? Residents in facility 60 days should not be without some identified interests.
- Are each resident's needs identified? If not, what actions are being taken to identify them?
- Have medical contraindications been identified in the care plan?
- Needs and contraindications of residents in the facility more than 30 days should be known and/or have a plan of action.
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<td>F242-(cont'd) Interests of each resident. It is designed to promote opportunities for engaging in normal pursuits, including religious activities of their choice, if any.</td>
<td>Are needed personal equipment (e.g., splints, glasses) and adaptations for limitations and safety (e.g., carholder, goggles, footrests) used in activities?</td>
<td>- Does resident get out of facility to activities?</td>
<td>- Needs of the resident in the following areas are identified: - social interaction + creative expression + work and service opportunities + intellectual stimulation or activities adaptation + physical exercise + spiritual or religious expression</td>
<td>Does each resident’s activities promote his physical, social and mental well-being?</td>
<td>Physical Environment 405.1134 442.399 Infection Control 405.1135 442.328 Resident Rights 405.1121(b) 405.311 Medical Records 405.1132 405.318 Patient Care Management 405.1124(c) 442.361</td>
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<tr>
<td>2. Unless contraindicated by the attending physician, all residents are encouraged to participate in activities.</td>
<td>- Does resident have problems getting to activities? If so, does the staff assist?</td>
<td>- Does the staff encourage residents to go to activities?</td>
<td>- Does resident participate in Resident Council?</td>
<td>- Does resident have free choice of activities?</td>
<td>- What kind of activities do best for residents engage in? Ask Resident: Have you ever had difficulty in having private visits? Give examples.</td>
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<td>F244 3. The activities promote the physical, social and mental well-being of the residents.</td>
<td>- Does resident participate in activities?</td>
<td>- Needs of the resident in the following areas are identified: - social interaction + creative expression + work and service opportunities + intellectual stimulation or activities adaptation + physical exercise + spiritual or religious expression</td>
<td>- Plan of care Used all available information about: + interests + needs + indications and contraindications for activities from other assessments + physician orders and progress notes</td>
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<td>F205 4. Equipment is maintained in good working order.</td>
<td>Is lighting adequate throughout the facility for activities in which residents are engaged? Do men and women have activities of interest to them? Do residents communicate with each other in activities? Are methods of communicating upcoming activities appropriate to the resident population? Specific observation for physically impaired residents: Activities adapted to meet specific needs of the resident. Alert residents have activities of interest and at their cognitive functional level. Specific observations for confused/disoriented, emotionally disturbed and mentally retarded residents. There are current calendars, clocks and patients.</td>
<td>Ask Nursing/Activity Staff: - Do they know the interests of residents under their care? - Programs they like? Activities they want to participate in today/this week? - Do they know the personal equipment needed (e.g., glasses, hearing aids, walker)? Do they know the adaptive equipment used by residents for specific activities (e.g., talking books, built up tools)? - Do they talk to residents to identify new interests and report these and &quot;dislikes&quot; to activities personnel? How? What is staff's involvement with individual and group activities of residents in their care? How do they determine interests of residents who have difficulty communicating? What activities does resident participate in regularly? Which activities does he/she enjoy most/least?</td>
<td>Activities notes spell out implementation of plan, resident's reactions to specific activities, approaches, and people. Residents' participation in individual and group self-started and organized structured and unstructured activities timespent. Evaluation of plan of care for: changes in interests; changes in precautions, changes in needs, new problems, approaches, etc. Plans are revised as needed.</td>
<td>Are equipment and supplies to meet residents' interests available and maintained in good working order? Are residents evaluated periodically with emphasis on participation levels and desire for new activities? Are plans readdressed if they do not reach desired outcomes? Residents in the facility more than 60 days should have at least two activities per week of interest to them personally.</td>
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### Long Term Care Survey

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<td>F246 (cont'd)</td>
<td>and patients names or symbols visible to all the residents.</td>
<td>- If he/she does not participate, why?</td>
<td>- Which activities appear to relax/ calms the resident?</td>
<td>- How does staff manage maladaptive behavior (e.g., obsessional, disruptive, combative)?</td>
<td>- Is direct care staff involved in resident activities? How? When (weekends, evenings)?</td>
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<td>Staff consistently use techniques such as reality orientation, empathy, and/or validation therapy as per each individual's needs.</td>
<td>- How does staff manage</td>
<td>- Does resident have assistance in activities?</td>
<td>- How many residents have activities a day of interest to them as individuals?</td>
<td>- Why do these residents have so little interest?</td>
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<td>Resident has familiar items available in room (e.g., family pictures, artwork, afghan, chair from home).</td>
<td>maladaptive behavior</td>
<td>- What is your plan to find more activities of interest to them that will meet their needs?</td>
<td>- What types of residents seem not to be interested in activities?</td>
<td>- How many (who) residents have only passive activities?</td>
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<td>Residents in restraints have activities of interest geared to their abilities when restrained (e.g., table-top activity, music, radio, reading, and writing materials); when out of restraints (e.g., walks, exercise, group, visiting).</td>
<td>- How many residents have few activities a day of interest to them as individuals?</td>
<td>- Why do these residents have so little interest?</td>
<td>- How many (who) residents have only passive activities?</td>
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<td>Small group and one-on-one involvement with staff reinforcing appropriate responses.</td>
<td>- Which activities appear to relax/ calms the resident?</td>
<td>- How does staff manage maladaptive behavior (e.g., obsessional, disruptive, combative)?</td>
<td>- Is direct care staff involved in resident activities? How? When (weekends, evenings)?</td>
<td>- Does resident have assistance in activities?</td>
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### F 246 (cont'd.)

**Specific observation for depressed or terminally ill resident:**
- Appropriate items for sensory enrichment in room (e.g., TV, radio, adequate lighting).
- Resident placed in supportive living environment (e.g., around people, in hall, activities room, sunshine, fresh air) when appropriate to the resident needs and consistent with the resident's choice.

- Adequate lighting.
- Functional area is appropriate for activities of interest (e.g., religious services, arts and crafts, cooking, reading, TV watching, card playing, parties, discussion groups, gardening).

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- How do you adapt activities for needs of residents who are:
  - Confused/disoriented
  - Emotionally disturbed
  - Mentally retarded
  - Physically impaired
  - Alert
  - Terminally ill?

- Are community volunteers utilized in the activities program? In what way?
- Are the residents encouraged to offer suggestions for new activities? If so, what activities have been instituted as a result?

- How they manage maladaptive behavior (e.g., abusive, disruptive, combative)?
- How do they help depressed residents (e.g., tearful, emotionally labile)?

Resident may refuse to participate in activity. However if the activities are part of a diagnostic or therapeutic program, the resident is responsible for assisting in the selection of mutually acceptable alternative activities.
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<td>F246 (cont'd)</td>
<td>- Multi-purpose room use and timing of activities does not conflict.</td>
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<td></td>
<td>- Outdoor activity area.</td>
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<td></td>
<td>- Functional furniture, indoors and outdoors.</td>
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<td></td>
<td>- Evidence of free-choice activities:</td>
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<td>- Magazines</td>
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<td>- Record player</td>
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<td>- Sewing</td>
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<td>- Personal visits</td>
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<td>- Church services</td>
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<td></td>
<td>- Activities, equipment, and supplies are appropriate and sufficient to meet interest of residents.</td>
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<td></td>
<td>- Activities equipment clean, safe, and in working order.</td>
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<td></td>
<td>- Residents' rooms contain independent project materials as appropriate.</td>
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<td></td>
<td>- Residents have access to the total activity environment (e.g., lobby, sunroom, day room, porch, dining room).</td>
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### Long Term Care Survey

<table>
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<tr>
<th>Survey Area</th>
<th>Observation</th>
<th>Interviewing</th>
<th>Record Review</th>
<th>Evaluation Factors</th>
<th>Cross Reference</th>
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<tbody>
<tr>
<td>Medical Records</td>
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<td>F248  &lt;br&gt; SNF 405.1132(c)</td>
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<td>F249  &lt;br&gt; CCF 602.318(a)(c)</td>
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<td></td>
<td>F250  &lt;br&gt; The medical record contains sufficient information to identify the resident clearly to justify diagnosis and treatment and to document results accurately.</td>
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<td>All information required is present in the record. Does the record document all observable resident needs/problems?</td>
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<td></td>
<td>F251  &lt;br&gt; The medical record contains the following information.</td>
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<td>a. Identification information.</td>
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<td>F252</td>
<td></td>
<td>b. Admission data including past medical social history.</td>
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<td>F253</td>
<td>c. Transfer form, discharge summary from any transferring facility.</td>
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<td>F255</td>
<td>e. Report of physical examinations.</td>
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<td>f. Reports of physicians' periodic evaluations and progress notes.</td>
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<td>F257</td>
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<td>g. Diagnostic reports and therapeutic orders.</td>
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<td>F258</td>
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<td>h. Reports of treatments.</td>
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<td>F259</td>
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<td>i. Medications administered.</td>
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<td>j. An overall plan of care setting forth goals to be accomplished through each service's designed activities, therapies and treatments.</td>
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<td>Reflects the care</td>
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<td>A. Whenever the</td>
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<td>Ask Staff:</td>
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<td>- What is the</td>
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<td>information you</td>
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<td>provide to a new</td>
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<td>facility when</td>
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<td>Review information</td>
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<td>on medical record</td>
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<td>of resident</td>
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<td>who was</td>
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<td>is again back in</td>
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<td>the facility.</td>
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<td>Look at physician</td>
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<td>and nursing</td>
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<td>progress notes of</td>
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<td>Does facility have</td>
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<td>hospital? Not</td>
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<td>All pertinent</td>
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<td>The resident was</td>
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<td>process.</td>
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<td>Patient Rights</td>
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<td>484.112(b)</td>
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<td>442.311</td>
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<td>F269 (cont'd)</td>
<td>hospital or a facility providing more specialized care and the nursing facility, admission to the new facility shall be effected in a timely manner.</td>
<td>Is transfer form complete with all data, with appropriate signatures? Does the medical record indicate that adequate and pertinent aspects of the discharge planning portion of the patient care plan accompany the patient on transfer?</td>
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<td>F269</td>
<td>B. Information necessary for providing care and treatment to transferred individuals is provided.</td>
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<td>PHYSICAL ENVIRONMENT</td>
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### LONG TERM CARE SURVEY

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<td>F271</td>
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<tr>
<td>A. Nursing Unit</td>
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</table>
|              | 1. Unit properly equipped for preparation and storage of drugs and biologicals. | There is adequate light to prepare medications. | Ask Nursing Staff:  
- What do you use the medication room (area) for?  
- Where is the handwashing sink?  
- Are there enough convenient storage areas for I.V. fluids and medications needing refrigeration? | Medication preparation and storage areas provide adequate space and light to prepare medication and to store medication and needed supplies.  
Light is available where and where the medication cart is in use.  
A medication refrigerator is available and does not contain patient or employee snacks. Juice, etc., used in administering medication is allowed.  
Clean and dirty areas must be separated, preferably in separate rooms.  
Storage space must be available for bulky items and supplies so that they can be stored without blocking corridors and exits.  
Medications are protected from unauthorized use.  
Call bells must be in working order and must be present in all resident bedrooms, toilets and | Nursing Service  
425.1124(c)  
442.337  
Infection Control  
425.1135  
Governor's Body  
442.325  
Resident Rooms  
425.1134(c)  
442.325 |
|              | 2. Utility and storage rooms are adequate size. | There is sufficient space to prepare medications for administration in a safe and effective manner. |              |                   |                 |
|              | 3. The unit is equipped to register resident calls with a functioning communications system from resident areas including rooms and toilets and bathing facility. | There is sufficient space for storage of medications. Unit dose carts are protected from tampering and theft. | Ask Nursing Staff:  
- Are the keys for the medication room and unit dose carts?  
- Do you feel you have adequate storage space for supplies and equipment?  
- If no, what problems does that cause?  
- Does the resident call system function properly? |             |                 |
|              |             | Medications are stored in a locked area. Refrigeration facilities are available for medications. | Ask Residents:  
- Do the call bells in your room and in the toilets and bathing areas always work? |             |                 |
|              |             | There is sufficient storage space for I.V. fluids. Handwashing facilities are readily accessible either in the medication preparation area or adjacent to it. |             |             |                 |
### Long Term Care Survey

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<tr>
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</table>
| F274 (cont'd) | Audible call system is on and working. Long cards are available for chair bound patients. | - If no:  
- How often is it that they do not work?  
- How long does it take to get them fixed? | Bathing areas.  
Audible signals, if in the system, must be in working order and turned on. | Regulations clearly set out conditions for compliance. Refer to the regulations. |  |

#### B. Dining and Activities Area

| F275  
SNF 405.1134(g)  
ICF 442.292 | Area is clean and well maintained.  
There is sufficient space between tables to allow for safe passage of wheelchairs and residents with walkers, canes and other assistive devices. | Ask Residents:  
- Is there enough room between tables to allow you to feel safe in getting to your table?  
- Can you sit comfortably in your wheelchair at the table?  
- How is the lighting and ventilation level for you?  
- Are sitting preferences permitted?  
- Do you go to the dining room for meals? | | Dietetic Services 405.1125  
402.331  
Patient Activities 405.1131  
402.345 |

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The facility provides one or more clean, orderly, and appropriately furnished rooms of adequate size, designed for resident dining and resident activities.  
Lighting and ventilation in the dining/activity areas is provided according to recommended standards.  
A multi-purpose room should not be used for storage of items such as beds, mattresses, boxes, etc.
### Long Term Care Survey

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<tbody>
<tr>
<td>F279</td>
<td>Are dining areas utilized at meal service?</td>
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<tr>
<td>F279</td>
<td>2. Dining and activity rooms are well lighted and ventilated.</td>
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<tr>
<td>F279</td>
<td>3. Any multi-purpose room used for dining and resident activities has sufficient space to accommodate all activities and prevent their interference with each other.</td>
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SNF 405.1134(e) Indicators 4D apply to SNFs
### Resident Rooms

<table>
<thead>
<tr>
<th>F281</th>
<th>ICF 442.325</th>
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</thead>
<tbody>
<tr>
<td>F282</td>
<td>Single rooms have at least 100 sq. ft.</td>
</tr>
<tr>
<td>F283</td>
<td>Multiple resident rooms have no more than 4 residents and at least 80 sq. feet per resident.</td>
</tr>
<tr>
<td>F284</td>
<td>Each room is equipped with or conveniently located near toilet and bathing facilities.</td>
</tr>
</tbody>
</table>

**Observation:**
- Observe rooms and furnishings for maintenance, cleanliness and safety.
- Look for dust/dirt on spots, high surfaces, under heating units, and in corners. Use a flashlight.
- Are beds, lights, plumbing all in working order?
- Observe for all regulatory requirements as noted on the left.
- Are privacy curtains present, and appropriate to maintain resident privacy?
- Test several call lights.
- Are call lights within reach, including emergency lights in toilets and bathing areas?
- Are toilet and bathing facilities appropriate in number, size, and design to meet resident needs?
- What personal belongings do residents have in their rooms? Is there

**Interviewing:**
- Ask Residents:
  - Is your room kept clean? Who cleans it? When, and how often?
  - Is your bed, chair, and other furniture and fixtures kept in good repair?
  - Do you feel you have enough privacy?
  - What personal belongings are you allowed to have?
  - Is the lighting in your room sufficient for you?
  - Is your chair comfortable?
  - When do you permit staff to clean you room?
  - When do you ask staff to give you your medication?

**Refer to the regulations.**

**Resident Rights**
- 405.112(6)(1)(a)(5)
- 442.311(a)(d)(2)
- 442.316(c)(1)(2)

**Physical Environment**
- 405.1134(d)(e)
- 442.326
<table>
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<tr>
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<tbody>
<tr>
<td>F285</td>
<td>4. There is a capability of maintaining privacy in each.</td>
<td>sufficient storage and security for their belongings?</td>
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<tr>
<td>F286</td>
<td>5. There is adequate storage space for each resident.</td>
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<tr>
<td>F287</td>
<td>6. There is a comfortable and functioning bed and chair, plus a functional cabinet and light.</td>
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<td>Survey Area</td>
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<tr>
<td>F290 7. The resident call system functions in resident rooms.</td>
<td></td>
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<tr>
<td>F299 8. Each room is designed and equipped for adequate nursing care and the comfort and privacy of residents.</td>
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<tr>
<td>F290 9. Each room is at or above grade level.</td>
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<tr>
<td>F291 10. Each room has direct access to a corridor and outside exposure.</td>
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Exception: Not required for ICF residents.
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<tr>
<td>0. Toilet and bath facilities</td>
<td>Are there adequate numbers of toilets, baths, and showers for the residents that are accessible to, and functional for all residents?</td>
<td>Ask Residents: - When was your last bath? - The one before? - What safety precautions are used for getting in and out of the bathtub? - What equipment is needed to get in and out of the tub, and how do you feel about it? - How do you get your wheelchair into the toilet or bathroom? - When, if ever, do you refuse to be bathed?</td>
<td>Bathing schedule for patients in your in-depth review.</td>
<td>Privacy is maintained for residents in toilet and bathing areas.</td>
<td>Toilet and bathing areas are clean. Water is removed from floors immediately upon completion of bathing.</td>
</tr>
<tr>
<td>F292</td>
<td>Are these conveniently located in or near resident rooms?</td>
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<tr>
<td>F293</td>
<td>F294</td>
<td>Facilities are clean, sanitary and free of odors.</td>
<td>Check for water on floors of bath and shower rooms. Is privacy provided? Are facilities clean, sanitary and free of unpleasant odors?</td>
<td></td>
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<tr>
<td>F295</td>
<td>Are bathrooms equipped with soap, toilet tissue, towels, etc.? Hot water is between 110-120 degrees or the acceptable State level. Hot water temperature control must be maintained. Simple use disposable towels should be available for handwashing purposes. Note also condition of grab bars, plumbing and fixtures. Bath areas are not used for storage.</td>
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<td>F296</td>
<td>Facilities have grab bars and other safe guards against slipping.</td>
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### Long Term Care Survey

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<tr>
<td>F297</td>
<td>5. Facilities have fixtures in good condition.</td>
<td>Does the social worker have a locked file available?</td>
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<tr>
<td>F298</td>
<td>6. The resident call system functions in toilet and bath facilities.</td>
<td>Where are social service interviews and clerical functions performed?</td>
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<tr>
<td>E. Social Service Area</td>
<td></td>
<td></td>
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<td></td>
<td>Refer to regulations.</td>
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<tr>
<td>F299</td>
<td>SW 405.113(b)</td>
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<tr>
<td>F300</td>
<td>1. Ensures privacy for social service interviewing.</td>
<td>Are rooms in areas easily accessible to residents?</td>
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<tr>
<td>F302</td>
<td>2. Adequate space for clerical and interviewing functions is provided.</td>
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<tr>
<td>F303</td>
<td>3. Facilities are easily accessible to residents and staff.</td>
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</table>
| 1. Therapy area | Therapy areas are accessible to all residents needing the facilities. Space allows for safe maneuvering of residents and equipment and staff. All residents are able to be observed and supervised during therapy. Equipment has labels (stickers, etc.) to indicate proper maintenance. All equipment fastened to floor and walls is secure. | Ask Resident:  - Do you feel that the equipment you use is safe?  - Do you have enough room for your treatment?  
Ask Therapy Staff:  - Is your equipment adequately maintained?  - Do you have enough room to safely and adequately provide treatment? | Refer to regulations. | | |
<p>| F303 SNF 485.1126(a) | | | | | |
| F304 ICF 442.320(a) | | | | | |
| F305 | | | | | |
| 1. Space is adequate for proper use of equipment by all residents receiving treatment | | | | | |
| 2. Facilities for special care | Are therapy areas properly ventilated to effectively reduce heat, moisture and odors? Are private rooms available that meet regulatory criteria. If a resident is infected and in isolation, are precautionary signs posted, and are they legible and understandable? | Ask Supervisor personnel:  - What room(s) do you use for isolation?  - What is your procedure if the room is already occupied when you need it for isolation?  - Will you show me the signs you use to identify the isolation room? | | | |
| F307 SNF 485.1134(f) | | | | | |
| F308 ICF 442.320(b) | | | | | |
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<td>F309</td>
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<tr>
<td>1. Single rooms with private toilet and handwashing facilities are available for isolating residents.</td>
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<td>F310</td>
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<td>2. Precautionary signs are used to identify these rooms when in use.</td>
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<td>H. Common Resident Areas</td>
<td>Use senses - sight, hearing, olfactory when surveying common areas as lounges, lobby, corridors. Are there floors, walls, ceilings, and furnishings that appear clean?</td>
<td>Ask Residents: - Do you think that the lounges and corridors are usually clean? - Do they have any unpleasant odors? - Is the lighting level comfortable for you to read? Is it adequate for you to feel safe at night? - Do you have any difficulty with the noise level? - Is the temperature usually comfortable for you? - Do you feel there is adequate ventilation? - Are there handrails in all of the corridors? - Are they securely fastened to the wall?</td>
<td>- Floors and furniture should appear clean - free of gross contamination. - Residents should have lighting bright enough to safely negotiate corridors, lounges, etc., and in reading area, be bright enough to read. But the brightness should be free of glare. Remember, the elderly need a higher level of lighting as their sight diminishes. - Except for times when a louder level of sound is necessary to communicate, sounds should be unobtrusive and &quot;comfortable&quot;. - Room temperature comfort levels vary widely, and in general the elderly will require a higher temperature for comfort than younger people. Use information from resident interviews your observations to determine if the temperature is &quot;comfortable&quot; for most residents. - All corridors in...</td>
<td>Infection Control 406.1135(c)</td>
<td></td>
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</tbody>
</table>
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<tr>
<td>F316</td>
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<td>resident-used areas are equipped with handrails on each side. These rails securely fastened provide the residents with a firm support. Supervisory staff are able to tell you how they will obtain water for drinking, cleaning/bathing of residents, and other essential functions if their normal water supply is interrupted.</td>
<td>Disaster Preparedness 482.1336 442.313</td>
</tr>
<tr>
<td>F317</td>
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</table>

4. A comfortable room temperature is maintained.

5. There is adequate ventilation through windows or mechanical measures or a combination of both.

6. Corridors are equipped with firmly secured handrails on each side.

7. Staff are aware of procedures to ensure water to all essential areas in the event of loss of normal supply.
### LONG TERM CARE SURVEY

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<tbody>
<tr>
<td>1. Maintenance of Building and Equipment</td>
<td>- Ceiling and floor tile in good condition</td>
<td>Ask Staff: - How many housekeeping staff are available? - How late are housekeepers on duty during the week? - How is weekend coverage different?</td>
<td></td>
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<td>Physical Environment 485.1134(d)</td>
</tr>
<tr>
<td></td>
<td>- Paint in good repair</td>
<td>Ask Resident: - What if any problems have you had with special equipment you need to use?</td>
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<td></td>
<td>- No holes in walls</td>
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<td></td>
<td>- Look for rodent trails outside and inside</td>
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<td></td>
<td>- Preventive maintenance program for all equipment is followed</td>
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<td></td>
<td>- Wheelchairs not stored in hallways, bathrooms, etc.</td>
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<td></td>
<td>- Window screens are in good repair</td>
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<td></td>
<td>- Check overbed tables, wheelchairs, etc., for cleanliness and operation</td>
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<td></td>
<td>- Sufficient storage space is available and used for equipment to ensure that the facility is orderly and safe.</td>
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<td>SURVEY AREA</td>
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<td>F324</td>
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<tr>
<td>4. Resident care equipment is clean and maintained in safe operating condition.</td>
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## § 488.115

### Long Term Care Survey

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<th>Survey Area</th>
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<tbody>
<tr>
<td><strong>Indicator 3</strong>&lt;br&gt;Applying to MDS 3.0&lt;br&gt;Dietetic Service Area F326&lt;br&gt;SNF 405.1134(h)</td>
<td><strong>F327</strong>&lt;br&gt;1. Kitchen and&lt;br&gt;dieterc service areas&lt;br&gt;are adequate to ensure proper&lt;br&gt;timely service for all patients.</td>
<td><strong>Ask Staff:</strong>&lt;br&gt;- What have you been trained to do?&lt;br&gt;- What type of dishwasher machine do you have?&lt;br&gt;- How does it operate?&lt;br&gt;<strong>The proper temperature for the dishwasher wash cycle is 150-160 degrees Fahrenheit. The dishwasher rinse cycle is acceptable at temperature of 180 degrees Fahrenheit or when there is a change in the temperature-sensitive tape (thermocouple). The individual manufacturers' specifications may countermand these instructions, particularly in the case of chemical sanitization.</strong></td>
<td></td>
<td></td>
<td>Diabetic Services&lt;br&gt;405.1135(g)&lt;br&gt;462.331(b)</td>
</tr>
<tr>
<td><strong>F328</strong>&lt;br&gt;2. Kitchen areas&lt;br&gt;are properly ventilated, arranged, and&lt;br&gt;equipped for storage and preparation of food as well as for dish and utensil cleaning, and refuse storage and removal.</td>
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<tr>
<th>SURVEY AREA</th>
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<tbody>
<tr>
<td>Indicator K applies to ICF K. Dietary Staff Hygiene</td>
<td>F329</td>
<td>SNF 405.1125(f)</td>
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<td></td>
<td>F330</td>
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<tr>
<td>1. Dietary service personnel practice hygienic food handling techniques</td>
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<tr>
<td>Indicator L applies to ICF</td>
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<tr>
<td>L. Dietary Sanitary Conditions</td>
<td>F333</td>
<td>SNF 405.1125(g)</td>
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<td></td>
<td>F332</td>
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<tr>
<td>1. Food is stored, refrigerated, prepared, distributed, and served under sanitary conditions.</td>
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<td></td>
<td>F333</td>
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<td>2. Waste is disposed of properly.</td>
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<td></td>
<td>Ask Staff:</td>
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<td></td>
<td>- What happens when you report to work with a cold, a cut or sore on your hand?</td>
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<td></td>
<td>- Where is handwashing sink for dietary staff?</td>
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<td></td>
<td>- Do you use disposable plastic hand covers? If so, when?</td>
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<td></td>
<td>- Where are your serving utensils located?</td>
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<td></td>
<td>- What are temperatures for the refrigerators and freezers? Who is responsible for checking temperatures?</td>
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<tr>
<td></td>
<td>- Do you have thermometers to check water and food temperatures? (ask them to demonstrate how they take temperatures)</td>
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<td>SURVEY AREA</td>
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</tbody>
</table>
| F333 (cont'd) | - check that the refrigerators are equipped with an accurate thermometer  
- food does not have an "off" or bad odor  
- cracked eggs are discarded  
- foods are dated and then stored as to their preparation date.  
Observe that waste is in covered containers, bagged and tied for disposal, and that dumpsters are covered. | | | | |
### Long Term Care Survey

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<tbody>
<tr>
<td>L. Emergency Power</td>
<td>Is an emergency generator available? Test generator under full load conditions.</td>
<td></td>
<td></td>
<td>As per regulations and covered by the Life Safety Code surveyor</td>
<td></td>
</tr>
<tr>
<td>1. An emergency power necessary to protect the health and safety of residents is available.</td>
<td>Check items of emergency power: - lighting - fire detection - alarms - extinguishing systems - life support systems</td>
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<tr>
<td>2. Emergency power is adequate at least for lighting in all means of egress; equipment to maintain fire detection, alarm, and extinguishing systems; and life support systems.</td>
<td>Check for grounded extension cards at nurses stations. Where are emergency outlets?</td>
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</tbody>
</table>
## Long Term Care Survey

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<tbody>
<tr>
<td>Infection Control</td>
<td>- Observation of dressing technique to identify if infection control principles are being adhered to; - Use of gloves</td>
<td>Ask Staff: - What type of dressing changes are you performing? - How often are dressings changed? - Why is resident on isolation precautions? - Do laundry/housekeeping personnel/sides know procedures?</td>
<td>Review records of residents selected for in-depth review for infection.</td>
<td>Compliance will be based mainly on your observations.</td>
<td>Deficiencies will be cited if you see: - Breaks in aseptic or isolation technique - Clutter or unclean conditions that would cause unsafe conditions - Inadequate supplies of linen to provide proper care and comfort for residents - Inadequate techniques for handling clean and dirty linen - Evidence of insect or rodent infestation - Use flash light to check for roaches in closets, cabinets.</td>
</tr>
<tr>
<td>A. Infection Control</td>
<td>- Observation of isolation precautions: - masks - gowns - gloves - handwashing - disposable items - information for visitors</td>
<td>- Ask Resident: - Do you know why you have dressings? - Do you know why you are on isolation precautions? - Do you have clean linen when you need it?</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>B. Sanitation</td>
<td>- Procedures followed by: - Laundry - Housekeeping</td>
<td>How is dirty linen transported to laundry or holding areas?</td>
<td>Do aids wash hands after cleaning dirty linen? How do aids handle clean/gray linen while changing beds?</td>
<td>-</td>
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</tr>
<tr>
<td>C. Linen</td>
<td>-</td>
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<td>SURVEY AREA</td>
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<td>F344</td>
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<td>ICF 442.327</td>
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<tr>
<td>F345</td>
<td>1. The facility has available at all times a quantity of linen essential for proper care and comfort of residents.</td>
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<tr>
<td>F346</td>
<td>2. Linens are handled, stored, processed, and transported in such a manner as to prevent the spread of infection.</td>
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<tr>
<td>F347</td>
<td>D. Pest Control</td>
<td>Look for evidence of insect or rodent presence (mouse or rat droppings, roaches, ants, flies around trash)</td>
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<tr>
<td></td>
<td>ICF 442.315(c)</td>
<td>- Screen doors closed</td>
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<tr>
<td></td>
<td>ICF 442.315(c)</td>
<td>- Windows that can be opened have screens that are in good repair</td>
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<td>F348</td>
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<td>Ask Staff:</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>- Have you seen insects (roaches, ants, flies, etc.)?</td>
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<td></td>
<td>- Have you seen rodents and/or droppings?</td>
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<td>- What foods are residents permitted to keep in their rooms?</td>
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<td>F349</td>
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<td></td>
<td>The facility is maintained free from insects and rodents.</td>
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</table>
### Long Term Care Survey

<table>
<thead>
<tr>
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</tr>
</thead>
</table>
| Disaster Preparedness | - Disaster plan is located at each nursing station.  
- Evacuation plans posted in each smoke compartment. | Ask Residents:  
- Do you know what to do in case of fire?  
- How often do you rehearse it?  
Ask Staff:  
- What are your responsibilities at a fire drill?  
- What is the facility’s disaster plan? (Specify types, i.e., fire, flood, etc.)  
- How you undergone disaster training?  
- Have you participated in a fire disaster drill? When?  
- How frequently are drills held?  
- Have you been trained/ instructed in the use of fire equipment, fire containment methods?  
- Have you been trained in transfer or casualties and routes?  
- How would staff meet emotional needs of residents during/following a "disaster", e.g., fire | | A disaster plan is available and facility staff know their roles. | Physical Environment 485.1134(a)(b) 442.321 |

Indicators A and B apply to ICFs.

#### A. Disaster Plan

<table>
<thead>
<tr>
<th>F350</th>
<th>SNF 405.1136</th>
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</thead>
<tbody>
<tr>
<td>F351</td>
<td>SNF 405.1136(a)</td>
</tr>
<tr>
<td>F352</td>
<td>ICF 442.313</td>
</tr>
</tbody>
</table>

1. Facility staff are aware of plans, procedures to be followed for fire, explosion or other disaster.

2. Facility staff are knowledgeable about evacuation routes.
<table>
<thead>
<tr>
<th>SURVEY AREA</th>
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<td>F355</td>
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<tr>
<td>3. Facility staff are aware of their specific responsibilities in regard to evaluation and protection of residents.</td>
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<td>F356</td>
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<tr>
<td>4. Facility staff are aware of methods of containing fire.</td>
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<tr>
<td>B. Drills</td>
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<td>F357</td>
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<td></td>
<td>SNF 405.1136(b)</td>
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<td>F358</td>
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<tr>
<td>1. All employees are trained as part of their employment orientation in all aspects of preparedness for any disaster.</td>
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<td>SURVEY AREA</td>
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<td>F359</td>
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<tr>
<td>2. Facility staff participate in ongoing training and drills in all procedures so that each employee promptly and correctly carries out a specific role in case of a disaster.</td>
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To ensure a clean, safe environment for residents.
§ 488.201 Reconsideration.

(a) Right to reconsideration. (1) A national accreditation organization dissatisfied with a determination that its accreditation requirements do not provide (or do not continue to provide) reasonable assurance that the entities accredited by the accreditation organization meet the applicable long-term care requirements, conditions for coverage, conditions of certification, conditions of participation, or CLIA condition level requirements is entitled to a reconsideration as provided in this subpart.

(2) A State dissatisfied with a determination that the requirements it imposes on laboratories in that State and under the laws of that State do not provide (or do not continue to provide) reasonable assurance that laboratories licensed or approved by the State meet applicable CLIA requirements is entitled to a reconsideration as provided in this subpart.

(b) Eligibility for reconsideration. HCFA will reconsider any determination to deny, remove or not renew, the approval of deeming authority to private accreditation organizations, or any determination to deny, remove or not renew the approval of a State laboratory program for the purpose of exempting the State’s laboratories from CLIA requirements, if the accreditation organization or State files a written request for a reconsideration in accordance with this subpart.

(c) Manner and timing of request for reconsideration. (1) A national accreditation organization or a State laboratory program described in paragraph (b), dissatisfied with a determination with respect to its deeming authority, or, in the case of a State, a determination with respect to the exemption of the laboratories in the State from CLIA requirements, may request a reconsideration of the determination by filing a request with HCFA either directly by its authorized officials or through its legal representative. The request must be filed within 60 days of the receipt of notice of an adverse determination or nonrenewal as provided in subpart A of part 488 or subpart E of part 493, as applicable.

(2) Reconsideration procedures are available after the effective date of the decision to deny, remove, or not renew the approval of an accreditation organization or State laboratory program.

(d) Content of request. The request for reconsideration must specify the findings or issues with which the accreditation organization or State disagrees and the reasons for the disagreement.

[57 FR 34012, July 31, 1992, as amended at 58 FR 61843, Nov. 23, 1993]

§ 488.203 Withdrawal of request for reconsideration.

A requestor may withdraw its request for reconsideration at any time before the issuance of a reconsideration determination.

§ 488.205 Right to informal hearing.

In response to a request for reconsideration, HCFA will provide the accreditation organization or the State laboratory program the opportunity for an informal hearing as described in § 488.207 that will—

(a) Be conducted by a hearing officer appointed by the Administrator of HCFA; and

(b) Provide the accreditation organization or State laboratory program the opportunity to present, in writing or in person, evidence or documentation to refute the determination to deny approval, or to withdraw or not renew deeming authority or the exemption of a State’s laboratories from CLIA requirements.

§ 488.207 Informal hearing procedures.

(a) HCFA will provide written notice of the time and place of the informal hearing at least 10 days before the scheduled date.

(b) The informal reconsideration hearing will be conducted in accordance with the following procedures—
(1) The hearing is open to HCFA and the organization requesting the reconsideration, including—
   (i) Authorized representatives;
   (ii) Technical advisors (individuals with knowledge of the facts of the case or presenting interpretation of the facts); and
   (iii) Legal counsel;
(2) The hearing is conducted by the hearing officer who receives testimony and documents related to the proposed action;
(3) Testimony and other evidence may be accepted by the hearing officer even though it would be inadmissible under the usual rules of court procedures;
(4) Either party may call witnesses from among those individuals specified in paragraph (b)(1) of this section; and
(5) The hearing officer does not have the authority to compel by subpoena the production of witnesses, papers, or other evidence.

§ 488.209 Hearing officer's findings.
(a) Within 30 days of the close of the hearing, the hearing officer will present the findings and recommendations to the accreditation organization or State laboratory program that requested the reconsideration.
(b) The written report of the hearing officer will include—
   (1) Separate numbered findings of fact; and
   (2) The legal conclusions of the hearing officer.

§ 488.211 Final reconsideration determination.
(a) The hearing officer's decision is final unless the Administrator, within 30 days of the hearing officer's decision, chooses to review that decision.
(b) The Administrator may accept, reject or modify the hearing officer's findings.
(c) Should the Administrator choose to review the hearing officer's decision, the Administrator will issue a final reconsideration determination to the accreditation organization or State laboratory program on the basis of the hearing officer's findings and recommendations and other relevant information.
(d) The reconsideration determination of the Administrator is final.
(e) A final reconsideration determination against an accreditation organization or State laboratory program will be published by HCFA in the Federal Register.

Subpart E—Survey and Certification of Long-Term Care Facilities

SOURCE: 59 FR 56238, Nov. 10, 1994, unless otherwise noted.

§ 488.300 Statutory basis.
Sections 1819 and 1919 of the Act establish requirements for surveying SNFs and NFs to determine whether they meet the requirements for participation in the Medicare and Medicaid programs.

§ 488.301 Definitions.
As used in this subpart—
Abbreviated standard survey means a survey other than a standard survey that gathers information primarily through resident-centered techniques on facility compliance with the requirements for participation. An abbreviated standard survey may be premised on complaints received; a change of ownership, management, or director of nursing; or other indicators of specific concern.
Abuse means the willful infliction of injury, unreasonable confinement, intimidation, or punishment with resulting physical harm, pain or mental anguish.
Deficiency means a SNF's or NF's failure to meet a participation requirement specified in the Act or in part 483, subpart B of this chapter.
Dually participating facility means a facility that has a provider agreement in both the Medicare and Medicaid programs.
Extended survey means a survey that evaluates additional participation requirements subsequent to finding substandard quality of care during a standard survey.
Facility means a SNF or NF, or a distinct part SNF or NF, in accordance with §483.5 of this chapter.
§ 488.303  State plan requirement.

(a) A State plan must provide that the requirements of this subpart and subpart F of this part are met, to the extent that those requirements apply to the Medicaid program.

(b) A State may establish a program to reward, through public recognition, incentive payments, or both, nursing facilities that provide the highest quality care to Medicaid residents. For purposes of section 1903(a)(7) of the Social Security Act, proper expenses incurred by a State in carrying out such a program are considered to be expenses necessary for the proper and efficient administration of the State plan.

(c) A State must conduct periodic educational programs for the staff and residents (and their representatives) of NFs in order to present current regulations, procedures, and policies under this subpart and subpart F of this part.

(d) Required remedies for a non-State operated NF. A State must establish, in addition to termination of the provider agreement, the following remedies or an approved alternative to the following remedies for imposition against a non-State operated NF:

(1) Temporary management.

(2) Denial of payment for new admissions.

(3) Civil money penalties.

(4) Transfer of residents.

(5) Closure of the facility and transfer of residents.

(6) State monitoring.

(e) Optional remedies for a non-State operated NF. A State may establish the following remedies for imposition against a non-State operated NF:

(1) Directed plan of correction.

(2) Directed in-service training.

(3) Alternative or additional State remedies.

(f) Alternative or additional State remedies. If a State uses remedies that
§ 488.308 Survey frequency.

(a) Basic period. The survey agency must conduct a standard survey of each SNF and NF not later than 15 months after the last day of the previous standard survey.

(b) Statewide average interval. (1) The statewide average interval between standard surveys must be 12 months or less, computed in accordance with paragraph (d) of this section.

(2) HCFA takes corrective action in accordance with the nature of the State survey agency's failure to ensure that the 12-month statewide average interval requirement is met. HCFA's corrective action is in accordance with §488.320.

(c) Other surveys. The survey agency may conduct a survey as frequently as necessary to—

1. Determine whether a facility complies with the participation requirements; and

2. Confirm that the facility has corrected deficiencies previously cited.

(d) Computation of statewide average interval. The statewide average interval is computed at the end of each Federal fiscal year by comparing the last day of the most recent standard survey for each participating facility to the last day of each facility's previous standard survey.

(e) Special surveys. (1) The survey agency may conduct a standard or an abbreviated standard survey to determine whether certain changes have caused a decline in the quality of care furnished by a SNF or NF, or causes a SNF or NF to be notified, of the time or date on which a standard survey is scheduled to be conducted is subject to a Federal civil money penalty not to exceed $2,000.

§ 488.307 Unannounced surveys.

(a) Basic rule. All standard surveys must be unannounced.

(b) Review of survey agency's scheduling and surveying procedures. (1) HCFA reviews on an annual basis each State survey agency's scheduling and surveying procedures and practices to ensure that survey agencies avoid giving notice of a survey through the scheduling procedures and the conduct of the surveys.

(2) HCFA takes corrective action in accordance with the nature and complexity of the problem when survey agencies are found to have notified a SNF or NF through their scheduling or procedural policies. Sanctions for inadequate survey performance are in accordance with §488.320.

(c) Civil money penalties. An individual who notifies a SNF or NF, or causes a SNF or NF to be notified, of the time or date on which a standard survey is scheduled to be conducted is subject to a Federal civil money penalty not to exceed $2,000.
(iv) Director of nursing.

(2) The survey agency must review all complaint allegations and conduct a standard or an abbreviated standard survey to investigate complaints of violations of requirements by SNFs and NFs if its review of the allegation concludes that—

(i) A deficiency in one or more of the requirements may have occurred; and

(ii) Only a survey can determine whether a deficiency or deficiencies exist.

(3) The survey agency does not conduct a survey if the complaint raises issues that are outside the purview of Federal participation requirements.

§ 488.310 Extended survey.

(a) Purpose of survey. The purpose of an extended survey is to identify the policies and procedures that caused the facility to furnish substandard quality of care.

(b) Scope of extended survey. An extended survey includes all of the following:

(1) Review of a larger sample of resident assessments than the sample used in a standard survey.

(2) Review of the staffing and in-service training.

(3) If appropriate, examination of the contracts with consultants.

(4) A review of the policies and procedures related to the requirements for which deficiencies exist.

(5) Investigation of any participation requirement at the discretion of the survey agency.

(c) Timing and basis for survey. The survey agency must conduct an extended survey not later than 14 calendar days after completion of a standard survey which found that the facility had furnished substandard quality of care.

§ 488.312 Consistency of survey results.

HCFA does and the survey agency must implement programs to measure accuracy and improve consistency in the application of survey results and enforcement remedies.

§ 488.314 Survey teams.

(a) Team composition. (1) Surveys must be conducted by a multidisciplinary team of professionals, which must include a registered nurse.

(2) Examples of professionals include, but are not limited to, physicians, physician assistants, nurse practitioners, physical, speech, or occupational therapists, registered professional nurses, dieticians, sanitarians, engineers, licensed practical nurses, or social workers.

(3) The State determines what constitutes a professional, subject to HCFA approval.

(4) Any of the following circumstances disqualifies a surveyor for surveying a particular facility:

(i) The surveyor currently works, or, within the past two years, has worked as an employee, as employment agency staff at the facility, or as an officer, consultant, or agent for the facility to be surveyed.

(ii) The surveyor has any financial interest or any ownership interest in the facility.

(iii) The surveyor has an immediate family member who has a relationship with a facility described in paragraphs (a)(4)(i) or paragraph (a)(4)(ii) of this section.

(iv) The surveyor has an immediate family member who is a resident in the facility to be surveyed. For purposes of this section, an immediate family member is defined at § 488.301 of this part.

(b) HCFA training. HCFA provides comprehensive training to surveyors, including at least the following:

(1) Application and interpretation of regulations for SNFs and NFs.

(2) Techniques and survey procedures for conducting standard and extended surveys.

(3) Techniques for auditing resident assessments and plans of care.

(c) Required surveyor training. (1) Except as specified in paragraph (c)(3) of this section, the survey agency may not permit an individual to serve as a member of a survey team unless the individual has successfully completed a training and testing program prescribed by the Secretary.

(2) The survey agency must have a mechanism to identify and respond to in-service training needs of the surveyors.
(3) The survey agency may permit an individual who has not completed a training program to participate in a survey as a trainee if accompanied on-site by a surveyor who has successfully completed the required training and testing program.

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§ 488.318 Inadequate survey performance.

(a) HCFA considers survey performance to be inadequate if the State survey agency—

(1) Indicates a pattern of failure to—

(i) Identify deficiencies and the failure cannot be explained by changed conditions in the facility or other case specific factors;

(ii) Cite only valid deficiencies;

(iii) Conduct surveys in accordance with the requirements of this subpart;

or

(iv) Use Federal standards, protocols, and the forms, methods and procedures specified by HCFA in manual instructions; or

(2) Fails to identify an immediate jeopardy situation.

(b) Inadequate survey performance does not—

(1) Relieve a SNF or NF of its obligation to meet all requirements for program participation; or

(2) Invalidate adequately documented deficiencies.

§ 488.320 Sanctions for inadequate survey performance.

(a) Annual assessment of survey performance. HCFA assesses the performance of the State's survey and certification program annually.

(b) Sanctions for inadequate survey performance. When a State demonstrates inadequate survey performance, as specified in § 488.318, HCFA notifies the survey agency of the inadequacy and takes action in accordance with paragraphs (c) and (d) of this section.

(c) Medicaid facilities. (1) For a pattern of failure to identify deficiencies in Medicaid facilities, HCFA—

(i) Reduces FFP, as specified in paragraph (e) of this section, and if appropriate;

(ii) Provides for training of survey teams.

(d) Medicare facilities. For all survey inadequacies in Medicare facilities, HCFA—

(1) Requires that the State survey agency submit a plan of correction;

(2) Provides for training of survey teams;

(3) Provides technical assistance on scheduling and procedural policies;

(4) Provides HCFA-directed scheduling; or

(5) Initiates action to terminate the agreement between the Secretary and the State under section 1864 of the Act, either in whole or in part.

(e) Reduction of FFP. In reducing FFP for inadequate survey performance, HCFA uses the formula specified in section 1919(g)(3)(C) of the Act, that is 33 percent multiplied by a fraction—

(1) The numerator of which is equal to the total number of residents in the NFs that HCFA found to be noncompliant during validation surveys for that quarter; and

(2) The denominator of which is equal to the total number of residents in the NFs in which HCFA conducted validation surveys during that quarter.

(f) Appeal of FFP reduction. When a State is dissatisfied with HCFA's determination to reduce FFP, the State may appeal the determination to the Departmental Appeals Board, using the procedures specified in 45 CFR part 16.

§ 488.325 Disclosures of results of surveys and activities.

(a) Information which must be provided to the public. As provided in sections 1819(g)(5) and 1919(g)(5) of the Act, the following information must be made available to the public, upon the public's request, by the State or HCFA for all surveys and certifications of SNFs and NFs:

(1) Statements of deficiencies and providers' comments.

(2) A list of isolated deficiencies that constitute no actual harm, with the potential for minimal harm.

(3) Approved plans of correction.

(4) Statements that the facility did not submit an acceptable plan of correction or failed to comply with the conditions of imposed remedies.
§ 488.330 Certification of compliance or noncompliance.

(a) General rules—(1) Responsibility for certification. (i) The State survey agency shall conduct surveys of all facilities for compliance with requirements for long term care facilities. The survey results are made available to the public in accordance with §488.320.

(2) Reports of adverse actions specified at §488.406 imposed on a facility.

(b) Written response by the provider. (1) Each resident in the facility with respect to which such finding was made; and

(ii) The name and address of his or her attending physician.

(2) Failure to disclose the information timely will result in termination of participation or imposition of alternative remedies.

§ 488.406 Certification of noncompliance.

(a) General rules—(1) Responsibility for certification. (i) The State survey agency shall conduct surveys of all facilities for compliance with requirements for long term care facilities. The survey results are made available to the public in accordance with §488.320.

(2) Reports of adverse actions specified at §488.406 imposed on a facility.

(b) Written response by the provider. (1) Each resident in the facility with respect to which such finding was made; and

(ii) The name and address of his or her attending physician.

(2) Failure to disclose the information timely will result in termination of participation or imposition of alternative remedies.
Health Care Financing Administration, HHS § 488.330

(A) The State certifies the compliance or noncompliance of non-State operated NFs. Regardless of the State entity doing the certification, it is final, except in the case of a complaint or validation survey conducted by HCFA, or HCFA review of the State's findings.

(B) HCFA certifies the compliance or noncompliance of all State-operated facilities.

(C) The State survey agency certifies the compliance or noncompliance of a non-State operated SNF, subject to the approval of HCFA.

(D) The State survey agency certifies compliance or noncompliance for a dually participating SNF/NF. In the case of a disagreement between HCFA and the State survey agency, a finding of noncompliance takes precedence over that of compliance.

(ii) In the case of a validation survey, the Secretary's determination as to the facility's noncompliance is binding, and takes precedence over a certification of compliance resulting from the State survey.

(2) Basis for certification. (i) Certification by the State is based on the survey agency findings.

(ii) Certification by HCFA is based on either the survey agency findings (in the case of State-operated facilities), or, in the case of a validation survey, on HCFA's own survey findings.

(b) Effect of certification—(1) Certification of compliance. A certification of compliance constitutes a determination that the facility is in substantial compliance and is eligible to participate in Medicaid as a NF, or in Medicare as a SNF, or in Medicare and Medicaid as a dually participating facility.

(2) Certification of noncompliance. A certification of noncompliance requires denial of participation for prospective providers and enforcement action for current providers in accordance with subpart F of this part. Enforcement actions must include one of the following:

(i) Termination of any Medicare or Medicaid provider agreements that are in effect.

(ii) Application of alternative remedies instead of, or in addition to, termination procedures.

(c) Notice of certification of noncompliance and resulting action. The notice of certification of noncompliance is sent in accordance with the timeframes specified in §488.402(f), and resulting action is issued by HCFA, except when the State is taking the action for a non-State operated NF.

(d) Content of notice of certification of noncompliance. The notice of certification of noncompliance is sent in accordance with the timeframes specified in §488.402(f) and includes information on all of the following:

(1) Nature of noncompliance.

(2) Any alternative remedies to be imposed under subpart F of this part.

(3) Any termination or denial of participation action to be taken under this part.

(4) The appeal rights available to the facility under this part.

(5) Timeframes to be met by the provider and certifying agency with regard to each of the enforcement actions or appeal procedures addressed in the notice.

(e) Appeals. (1) Notwithstanding any provision of State law, the State must impose remedies promptly on any provider of services participating in the Medicaid program—

(i) After promptly notifying the facility of the deficiencies and impending remedy or remedies; and

(ii) Except for civil money penalties, during any pending hearing that may be requested by the provider of services.

(2) HCFA imposes remedies promptly on any provider of services participating in the Medicare or Medicaid program or any provider of services participating in both the Medicare and Medicaid programs—

(i) After promptly notifying the facility of the deficiencies and impending remedy or remedies; and

(ii) Except for civil money penalties, during any pending hearing that may be requested by the provider of services.

(3) The provisions of part 498 of this chapter apply when the following providers request a hearing on a denial of participation, or certification of noncompliance leading to an enforcement remedy (including termination of the provider agreement), except State monitoring:

(i) All State-operated facilities;
§ 488.331 Informal dispute resolution.

(a) Opportunity to refute survey findings. (1) For non-Federal surveys, the State must offer a facility an informal opportunity, at the facility's request, to dispute survey findings upon the facility's receipt of the official statement of deficiencies.

(2) For Federal surveys, HCFA offers a facility an informal opportunity, at the facility's request, to dispute survey findings upon the facility's receipt of the official statement of deficiencies.

(b)(1) Failure of the State or HCFA, as appropriate, to complete informal dispute resolution timely cannot delay the effective date of any enforcement action against the facility.

(2) A facility may not seek a delay of any enforcement action against it on the grounds that informal dispute resolution has not been completed before the effective date of the enforcement action.

(c) If a provider is subsequently successful, during the informal dispute resolution process, at demonstrating that deficiencies should not have been cited, the deficiencies are removed from the statement of deficiencies and any enforcement actions imposed solely as a result of those cited deficiencies are rescinded.

§ 488.332 Investigation of complaints of violations and monitoring of compliance.

(a) Investigation of complaints. (1) The State survey agency must establish procedures and maintain adequate staff to investigate complaints of violations of participation requirements.

(2) The State survey agency takes appropriate precautions to protect a complainant's anonymity and privacy, if possible.

(3) If arrangements have been made with other State components for investigation of complaints, the State must have a means of communicating information among appropriate entities, and the State survey agency retains responsibility for the investigation process.

(4) If, after investigating a complaint, the State has reason to believe that an identifiable individual neglected or abused a resident, or misappropriated a resident's property, the State survey agency must act on the complaint in accordance with § 488.335.

(b) On-site monitoring. The State survey agency conducts on-site monitoring on an as necessary basis when—

(1) A facility is not in substantial compliance with the requirements and is in the process of correcting deficiencies;
§ 488.335 Action on complaints of resident neglect and abuse, and misappropriation of resident property.

(a) Investigation. (1) The State must review all allegations of resident neglect and abuse, and misappropriation of resident property and follow procedures specified in § 488.332.

(2) If there is reason to believe, either through oral or written evidence that an individual used by a facility to provide services to residents could have abused or neglected a resident or misappropriated a resident's property, the State must investigate the allegation.

(3) The State must have written procedures for the timely review and investigation of allegations of resident abuse and neglect, and misappropriation of resident property.

(b) Source of complaints. The State must review all allegations regardless of the source.

(c) Notification—(1) Individuals to be notified. If the State makes a preliminary determination, based on oral or written evidence and its investigation, that the abuse, neglect or misappropriation of property occurred, it must notify in writing—

(i) The individuals implicated in the investigation; and

(ii) The current administrator of the facility in which the incident occurred.

(2) Timing of the notice. The State must notify the individuals specified in paragraph (c)(1) of this section in writing within 10 working days of the State's investigation.

(3) Contents of the notice. The notice must include the—

(i) Nature of the allegation(s);

(ii) Date and time of the occurrence;

(iii) Right to a hearing;

(iv) Intent to report the substantiated findings in writing, once the individual has had the opportunity for a hearing, to the nurse aide registry or appropriate licensure authority;

(v) Fact that the individual's failure to request a hearing in writing within 30 days from the date of the notice will result in reporting the substantiated findings to the nurse aide registry or appropriate licensure authority.

(vi) Consequences of waiving the right to a hearing;

(vii) Consequences of a finding through the hearing process that the alleged resident abuse or neglect, or misappropriation of resident property did occur; and

(viii) Fact that the individual has the right to be represented by an attorney at the individual's own expense.

(d) Conduct of hearing. (1) The State must complete the hearing and the hearing record within 120 days from the day it receives the request for a hearing.

(2) The State must hold the hearing at a reasonable place and time convenient for the individual.

(e) Factors beyond the individual's control. A State must not make a finding that an individual has neglected a resident if the individual demonstrates that such neglect was caused by factors beyond the control of the individual.

(f) Report of findings. If the finding is that the individual has neglected or abused a resident or misappropriated resident property or if the individual waives the right to a hearing, the State must report the findings in writing within 10 working days to—

(1) The individual;

(2) The current administrator of the facility in which the incident occurred; and

(3) The administrator of the facility that currently employs the individual,
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if different than the facility in which
the incident occurred;
(4) The licensing authority for indi-
viduals used by the facility other than
nurse aides, if applicable; and
(5) The nurse aide registry for nurse
aides. Only the State survey agency
may report the findings to the nurse
aide registry, and this must be done
within 10 working days of the findings,
in accordance with § 483.156(c) of this
chapter. The State survey agency may
not delegate this responsibility.
(g) Contents and retention of report of
finding to the nurse aide registry. (1) The
report of finding must include informa-
tion in accordance with § 483.156(c) of
this chapter.
(2) The survey agency must retain
the information as specified in para-
graph (g)(1) of this section, in accord-
ance with the procedures specified in
§ 483.156(c) of this chapter.
(h) Survey agency responsibility. (1) The
survey agency must promptly re-
view the results of all complaint inves-
tigations and determine whether or not
a facility has violated any require-
ments in part 483, subpart B of this
chapter.
(2) If a facility is not in substantial
compliance with the requirements in part 483, subpart B of this chapter, the
survey agency initiates appropriate ac-
tions, as specified in subpart F of this
part.
[59 FR 56238, Nov. 10, 1994; 60 FR 50118, Sept.
28, 1995]

Subpart F—Enforcement of Com-
pliance for Long-Term Care
Facilities with Deficiencies

SOURCE: 59 FR 56243, Nov. 10, 1994, unless
otherwise noted.

§ 488.400 Statutory basis.

Sections 1819(h) and 1919(h) of the Act
specify remedies that may be used by
the Secretary or the State respectively
when a SNF or a NF is not in substan-
tial compliance with the requirements
for participation in the Medicare and
Medicaid programs. These sections also
provide for ensuring prompt compli-
ance and specify that these remedies
are in addition to any others available
under State or Federal law, and, except
for civil money penalties, are imposed
prior to the conduct of a hearing.

§ 488.401 Definitions.

As used in this subpart—
New admission means a resident who
is admitted to the facility on or after
the effective date of a denial of pay-
ment remedy and, if previously admit-
ted, has been discharged before that ef-
teffective date. Residents admitted before
the effective date of the denial of pay-
ment, and taking temporary leave, are
not considered new admissions, nor
subject to the denial of payment.
Plan of correction means a plan de-
veloped by the facility and approved by
HCFA or the survey agency that de-
scribes the actions the facility will
take to correct deficiencies and speci-
fies the date by which those defi-
ciencies will be corrected.
[59 FR 56243, Nov. 10, 1994; 60 FR 50118, Sept.
28, 1995]

§ 488.402 General provisions.

(a) Purpose of remedies. The purpose of
remedies is to ensure prompt compli-
ance with program requirements.
(b) Basis for imposition and duration of
remedies. When HCFA or the State
chooses to apply one or more remedies
specified in § 488.406, the remedies are
applied on the basis of noncompliance
found during surveys conducted by
HCFA or by the survey agency.
(c) Number of remedies. HCFA or the
State may apply one or more remedies
for each deficiency constituting non-
compliance or for all deficiencies con-
stituting noncompliance.
(d) Plan of correction requirement. (1) Except as specified in paragraph (d)(2)
of this section, regardless of which
remedy is applied, each facility that
has deficiencies with respect to pro-
gram requirements must submit a plan
of correction for approval by HCFA or
the survey agency.
(2) Isolated deficiencies. A facility is
not required to submit a plan of correc-
tion when it has deficiencies that are
isolated and have a potential for mini-
mal harm, but no actual harm has oc-
curred.
(e) Disagreement regarding remedies. If the State and HCFA disagree on the decision to impose a remedy, the disagreement is resolved in accordance with §488.452.

(f) Notification requirements—(1) Except when the State is taking action against a non-State operated NF, HCFA or the State (as authorized by HCFA) gives the provider notice of the remedy, including the—
   (i) Nature of the noncompliance;
   (ii) Which remedy is imposed;
   (iii) Effective date of the remedy; and
   (iv) Right to appeal the determination leading to the remedy.

(2) When a State is taking action against a non-State operated NF, the State's notice must include the same information required by HCFA in paragraph (f)(1) of this section.

(3) Immediate jeopardy—2 day notice. Except for civil money penalties and State monitoring imposed when there is immediate jeopardy, for all remedies specified in §488.406 imposed when there is immediate jeopardy, the notice must be given at least 2 calendar days before the effective date of the enforcement action.

(4) No immediate jeopardy—15 day notice. Except for civil money penalties and State monitoring, notice must be given at least 15 calendar days before the effective date of the enforcement action in situations in which there is no immediate jeopardy.

(5) Date of enforcement action. The 2- and 15-day notice periods begin when the facility receives the notice.

(6) Civil money penalties. For civil money penalties, the notices must be given in accordance with the provisions of §§488.434 and 488.440.

(7) State monitoring. For State monitoring, no prior notice is required.

§488.404 Factors to be considered in selecting remedies.

(a) Initial assessment. In order to select the appropriate remedy, if any, to apply to a facility with deficiencies, HCFA and the State determine the seriousness of the deficiencies.

(b) Determining seriousness of deficiencies. To determine the seriousness of the deficiency, HCFA considers and the State must consider at least the following factors:

(1) Whether a facility's deficiencies constitute—
   (i) No actual harm with a potential for minimal harm;
   (ii) No actual harm with a potential for more than minimal harm, but not immediate jeopardy;
   (iii) Actual harm that is not immediate jeopardy; or
   (iv) Immediate jeopardy to resident health or safety.

(2) Whether the deficiencies—
   (i) Are isolated;
   (ii) Constitute a pattern; or
   (iii) Are widespread.

(b) Other factors which may be considered in choosing a remedy within a remedy category. Following the initial assessment, HCFA and the State may consider other factors, which may include, but are not limited to the following:

(1) The relationship of the one deficiency to other deficiencies resulting in noncompliance.

(2) The facility's prior history of noncompliance in general and specifically with reference to the cited deficiencies.

§488.406 Available remedies.

(a) General. In addition to the remedy of termination of the provider agreement, the following remedies are available:

(1) Temporary management.

(2) Denial of payment including—
   (i) Denial of payment for all individuals, imposed by HCFA, to a—
      (A) Skilled nursing facility, for Medicare;
      (B) State, for Medicaid; or
   (ii) Denial of payment for all new admissions.

(3) Civil money penalties.

(4) State monitoring.

(5) Transfer of residents.

(6) Closure of the facility and transfer of residents.

(7) Directed plan of correction.

(8) Directed in-service training.

(9) Alternative or additional State remedies approved by HCFA.

(b) Remedies that must be established. At a minimum, and in addition to termination of the provider agreement, the State must establish the following...
§ 488.408 Selection of remedies.

(a) Categories of remedies. In this section, the remedies specified in §488.406(a) are grouped into categories and applied to deficiencies according to how serious the noncompliance is.

(b) Application of remedies. After considering the factors specified in §488.404, as applicable, if HCFA and the State choose to impose remedies, as provided in paragraphs (c)(1), (d)(1) and (e)(1) of this section, for facility noncompliance, instead of, or in addition to, termination of the provider agreement, HCFA does and the State must follow the criteria set forth in paragraphs (c)(2), (d)(2), and (e)(2) of this section, as applicable.

(c) Category 1. (1) Category 1 remedies include the following:

(i) Directed plan of correction.

(ii) State monitoring.

(iii) Directed in-service training.

(2) HCFA does or the State must apply one or more of the remedies in Category 1 when there—

(i) Are isolated deficiencies that constitute no actual harm with a potential for more than minimal harm but not immediate jeopardy; or

(ii) Is a pattern of deficiencies that constitutes no actual harm with a potential for more than minimal harm but not immediate jeopardy.

(3) Except when the facility is in substantial compliance, HCFA or the State may apply one or more of the remedies in Category 1 to any deficiency.

(d) Category 2. (1) Category 2 remedies include the following:

(i) Denial of payment for new admissions.

(ii) Denial of payment for all individuals imposed only by HCFA.

(iii) Civil money penalties of $50-$3,000 per day.

(iv) Civil money penalty of $1,000-$10,000 per instance of noncompliance.

(2) HCFA applies one or more of the remedies in Category 2, or, except for denial of payment for all individuals, the State must apply one or more of the remedies in Category 2 when there are—

(i) Widespread deficiencies that constitute no actual harm with a potential for more than minimal harm but not immediate jeopardy; or

(ii) One or more deficiencies that constitute actual harm that is not immediate jeopardy.

(3) HCFA or the State may apply one or more of the remedies in Category 2 to any deficiency except when—

(i) The facility is in substantial compliance; or

(ii) HCFA or the State imposes a civil money penalty for a deficiency that constitutes immediate jeopardy, the penalty must be in the upper range of penalty amounts, as specified in §488.438(a).

(e) Category 3. (1) Category 3 remedies include the following:

(i) Temporary management.

(ii) Immediate termination.

(iii) Civil money penalties of $3,050-$10,000 per day.

(iv) Civil money penalty of $1,000-$10,000 per instance of noncompliance.
§ 488.410 Action when there is immediate jeopardy.

(a) If there is immediate jeopardy to resident health or safety, the State must either terminate the provider agreement within 23 calendar days of the last date of the survey or appoint a temporary manager to remove the immediate jeopardy. The rules for appointment of a temporary manager in an immediate jeopardy situation are as follows:

(1) HCFA does and the State must notify the facility that a temporary manager is being appointed.

(2) If the facility fails to relinquish control to the temporary manager, HCFA does and the State must terminate the provider agreement within 23 calendar days of the last day of the survey, if the immediate jeopardy is not removed. In these cases, State monitoring may be imposed pending termination.

(3) If the facility relinquishes control to the temporary manager, the State must (and HCFA does) notify the facility that, unless it removes the immediate jeopardy, its provider agreement will be terminated within 23 calendar days of the last day of the survey.

(b) HCFA or the State may also impose other remedies, as appropriate.

(c) (1) In a NF or dually participating facility, if either HCFA or the State finds that a facility's noncompliance poses immediate jeopardy to resident health or safety, HCFA or the State must notify the other of such a finding.

(2) HCFA will (or the State must) do one or both of the following:

(i) Take immediate action to remove the jeopardy and correct the noncompliance through temporary management.

(ii) Terminate the facility's participation under the State plan. If this is done, HCFA will also terminate the facility's participation in Medicare if it is a dually participating facility.

(d) The State must provide for the safe and orderly transfer of residents when the facility is terminated.

(e) If the immediate jeopardy is also substandard quality of care, the State survey agency must notify attending physicians and the State board responsible for licensing the facility administrator of the finding of substandard quality of care, as specified in § 488.325(h).
§ 488.412 Action when there is no immediate jeopardy.

(a) If a facility's deficiencies do not pose immediate jeopardy to residents' health or safety, and the facility is not in substantial compliance, HCFA or the State may terminate the facility's provider agreement or may allow the facility to continue to participate for no longer than 6 months from the last day of the survey if—

(1) The State survey agency finds that it is more appropriate to impose alternative remedies than to terminate the facility's provider agreement;

(2) The State has submitted a plan and timetable for corrective action approved by HCFA; and

(3) The facility in the case of a Medicare SNF or the State in the case of a Medicaid NF agrees to repay to the Federal government payments received after the last day of the survey that first identified the deficiencies if corrective action is not taken in accordance with the approved plan of correction.

(b) If a facility does not meet the criteria for continuation of payment under paragraph (a) of this section, HCFA will and the State must terminate the facility's provider agreement.

(c) HCFA does and the State survey agency must impose State monitoring, as specified in § 488.422, until the facility has demonstrated to the satisfaction of HCFA or the State, that it is in substantial compliance with all requirements and will remain in substantial compliance with all requirements.

(2) The State must impose denial of payment for all new admissions, as specified in § 488.417; and

(3) HCFA does and the State survey agency must impose State monitoring, as specified in § 488.422, until the facility has demonstrated to the satisfaction of HCFA or the State, that it is in substantial compliance with all requirements and will remain in substantial compliance with all requirements.

(b) Repeated noncompliance. For purposes of this section, repeated noncompliance is based on the repeated finding of substandard quality of care and not on the basis that the substance of the deficiency or the exact tag number for the deficiency was repeated.

(c) Standard surveys to which this provision applies. Standard surveys completed by the State survey agency on or after October 1, 1990, are used to determine whether the threshold of three consecutive standard surveys is met.

(d) Program participation. (1) The determination that a certified facility has repeated instances of substandard quality of care is made without regard to any variances in the facility's program participation (that is, any standard survey completed for Medicare, Medicaid or both programs will be considered).

(2) Termination would allow the count of repeated substandard quality of care surveys to start over.

(3) Change of ownership. (i) A facility may not avoid a remedy on the basis that it underwent a change of ownership.

(ii) In a facility that has undergone a change of ownership, HCFA does not and the State may not restart the count of repeated substandard quality of care surveys unless the new owner can demonstrate to the satisfaction of HCFA or the State that the poor past performance no longer is a factor due to the change in ownership.

(e) Facility alleges corrections or achieves compliance after repeated substandard quality of care is identified. (1) If a penalty is imposed for repeated substandard quality of care, it will continue until the facility has demonstrated to the satisfaction of HCFA or the State that it is in substantial compliance with the requirements and
that it will remain in substantial compliance with the requirements for a period of time specified by HCFA or the State.

(2) A facility will not avoid the imposition of remedies or the obligation to demonstrate that it will remain in compliance when it—

(i) Alleges correction of the deficiencies cited in the most recent standard survey; or

(ii) Achieves compliance before the effective date of the remedies.

§ 488.415 Temporary management.

(a) Definition. Temporary management means the temporary appointment by HCFA or the State of a substitute facility manager or administrator with authority to hire, terminate or reassign staff, obligate facility funds, alter facility procedures, and manage the facility to correct deficiencies identified in the facility’s operation.

(b) Qualifications. The temporary manager must—

(1) Be qualified to oversee correction of deficiencies on the basis of experience and education, as determined by the State;

(2) Not have been found guilty of misconduct by any licensing board or professional society in any State;

(3) Have, or a member of his or her immediate family have, no financial ownership interest in the facility; and

(4) Not currently serve or, within the past 2 years, have served as a member of the staff of the facility.

(c) Payment of salary. The temporary manager’s salary—

(1) Is paid directly by the facility while the temporary manager is assigned to that facility; and

(2) Must be at least equivalent to the sum of the following—

(i) The prevailing salary paid by providers for positions of this type in what the State considers to be the facility’s geographic area;

(ii) Additional costs that would have reasonably been incurred by the provider if such person had been in an employment relationship; and

(iii) Any other costs incurred by such a person in furnishing services under such an arrangement or as otherwise set by the State.

(3) May exceed the amount specified in paragraph (c)(2) of this section if the State is otherwise unable to attract a qualified temporary manager.

(d) Failure to relinquish authority to temporary management—(1) Termination of provider agreement. If a facility fails to relinquish authority to the temporary manager as described in this section, HCFA will or the State must terminate the provider agreement in accordance with §488.456.

(2) Failure to pay salary of temporary manager. A facility’s failure to pay the salary of the temporary manager is considered a failure to relinquish authority to temporary management.

(e) Duration of temporary management. Temporary management ends when the facility meets any of the conditions specified in §488.454(c).

§ 488.417 Denial of payment for all new admissions.

(a) Optional denial of payment. Except as specified in paragraph (b) of this section, HCFA or the State may deny payment for all new admissions when a facility is not in substantial compliance with the requirements, as defined in §488.401, as follows:

(1) Medicare facilities. In the case of Medicare facilities, HCFA may deny payment to the facility.

(2) Medicaid facilities. In the case of Medicaid facilities—

(i) The State may deny payment to the facility; and

(ii) HCFA may deny payment to the State for all new Medicaid admissions to the facility.

(b) Required denial of payment. HCFA does or the State must deny payment for all new admissions when—

(1) The facility is not in substantial compliance, as defined in §488.401, 3 months after the last day of the survey identifying the noncompliance; or

(2) The State survey agency has cited a facility with substandard quality of care on the last three consecutive standard surveys.

(c) Resumption of payments: Repeated instances of substandard quality of care. When a facility has repeated instances of substandard quality of care, payments to the facility or, under Medicaid, HCFA payments to the State on
behalf of the facility, resume on the date that—
(1) The facility achieves substantial compliance as indicated by a revisit or written credible evidence acceptable to HCFA (for all facilities except non-State operated NFs against which HCFA is imposing no remedies) or the State (for non-State operated NFs against which HCFA is imposing no remedies); and
(2) HCFA (for all facilities except non-State operated NFs against which HCFA is imposing no remedies) or the State (for non-State operated NFs against which HCFA is imposing no remedies) believes that the facility is capable of remaining in substantial compliance.
(d) Resumption of payments: No repeated instances of substandard quality of care. When a facility does not have repeated instances of substandard quality of care, payments to the facility or, under Medicaid, HCFA payments to the State on behalf of the facility, resume prospectively on the date that the facility achieves substantial compliance, as indicated by a revisit or written credible evidence acceptable to HCFA (under Medicare) or the State (under Medicaid).
(e) Restriction. No payments to a facility or, under Medicaid, HCFA payments to the State on behalf of the facility, are made for the period between the date that—
(1) Denial of payment remedy is imposed; and
(2) Facility achieves substantial compliance, as determined by HCFA or the State.

§ 488.422 State monitoring.
(a) A State monitor—
(1) Oversees the correction of deficiencies specified by HCFA or the State survey agency at the facility site and protects the facility's residents from harm;
(2) Is an employee or a contractor of the survey agency;
(3) Is identified by the State as an appropriate professional to monitor cited deficiencies;
(4) Is not an employee of the facility;
(5) Does not function as a consultant to the facility; and
§ 488.432 Civil money penalties: When a penalty is collected.

(a) When facility requests a hearing. (1) A facility must request a hearing on the determination of the noncompliance that is the basis for imposition of the civil money penalty within the time specified in one of the following sections:

(i) Section 498.40 of this chapter for a (A) SNF; (B) Dually participating facility; (C) State-operated NF; or (D) Non-State operated NF against which HCFA is imposing remedies.

(ii) Section 431.153 of this chapter for a non-State operated NF that is not subject to imposition of remedies by HCFA.

(ii) If a facility requests a hearing within the time specified in paragraph (a)(1) of this section, for a civil money penalty imposed per day, HCFA or the
§ 488.434 Civil money penalties: Notice of penalty.

(a) HCFA notice of penalty. (1) HCFA sends a written notice of the penalty to the facility for all facilities except non-State operated NFs when the State is imposing the penalty.

(b) State notice of penalty. (1) The State must notify the facility in accordance with State procedures for all non-State operated NFs when the State takes the action.

(c) When a facility waives a hearing. (1) If a facility waives, in writing, its right to a hearing as specified in §488.436, for a civil money penalty imposed per day, HCFA or the State initiates collection of the penalty when the facility—

(i) Achieves substantial compliance; or

(ii) Is terminated.

(2) If a facility waives, in writing, its right to a hearing as specified in §488.436, for a civil money penalty imposed per instance of noncompliance, HCFA or the State initiates collection of the penalty upon receipt of the facility's notification.

(d) Accrual and computation of penalties for a facility that—

(1) Requests a hearing or does not request a hearing are specified in §488.440;

(2) Waives its right to a hearing in writing, are specified in §§488.436(b) and 488.440.

(e) The collection of civil money penalties is made as provided in §488.442.

[59 FR 56243, Nov. 10, 1994; 60 FR 50119, Sept. 28, 1995, as amended at 64 FR 13360, Mar. 18, 1999]

§ 488.434 Civil money penalties: Notice of penalty.

(a) HCFA notice of penalty. (1) HCFA sends a written notice of the penalty to the facility for all facilities except non-State operated NFs when the State is imposing the penalty.

(b) Content of notice. The notice that HCFA sends includes—

(i) The nature of the noncompliance;

(ii) The statutory basis for the penalty;

(iii) The amount of penalty per day of noncompliance or the amount of the penalty per instance of noncompliance;

(iv) Any factors specified in §488.438(f) that were considered when determining the amount of the penalty;

(v) The date of the instance of noncompliance or the date on which the penalty begins to accrue;

(vi) When the penalty stops accruing, if applicable;

(vii) When the penalty is collected; and

(viii) Instructions for responding to the notice, including a statement of the facility's right to a hearing, and the implication of waiving a hearing, as provided in §488.436.

(b) State notice of penalty. (1) The State must notify the facility in accordance with State procedures for all non-State operated NFs when the State takes the action.

(2) The State's notice must—

(i) Be in writing; and

(ii) Include, at a minimum, the information specified in paragraph (a)(2) of this section.

[59 FR 56243, Nov. 10, 1994; 60 FR 50119, Sept. 28, 1995, as amended at 64 FR 13360, Mar. 18, 1999]

§ 488.436 Civil money penalties: Waiver of hearing, reduction of penalty amount.

(a) Waiver of a hearing. The facility may waive the right to a hearing, in writing, within 60 days from the date of the notice imposing the civil money penalty.
§ 488.440 Civil money penalties: Effective date and duration of penalty.

(a) (1) The per day civil money penalty may start accruing as early as the date that the facility was first out of compliance, as determined by HCFA or the State.

(2) A civil money penalty for each instance of noncompliance is imposed in a specific amount for that particular deficiency.

(b) The per day civil money penalty is computed and collectible, as specified in §§ 488.432 and 488.442, for the
number of days of noncompliance until the date the facility achieves substantial compliance, or, if applicable, the date of termination when—

(1) HCFA's or the State's decision of noncompliance is upheld after a final administrative decision;

(2) The facility waives its right to a hearing in accordance with §488.436; or

(3) The time for requesting a hearing has expired and HCFA or the State has not received a hearing request from the facility.

c. The entire penalty, whether imposed on a per day or per instance basis, is due and collectible as specified in the notice sent to the provider under paragraphs (d) and (e) of this section.

d. (1) When a civil money penalty is imposed on a per day basis and the facility achieves substantial compliance, HCFA does or the State must send a separate notice to the facility containing the following information:

(i) The amount of penalty per day.

(ii) The number of days involved.

(iii) The total amount due.

(iv) The due date of the penalty.

(v) The rate of interest assessed on the unpaid balance beginning on the due date, as provided in §488.442.

(2) When a civil money penalty is imposed for an instance of noncompliance, HCFA does or the State must send a separate notice to the facility containing the following information:

(i) The amount of the penalty.

(ii) The total amount due.

(iii) The due date of the penalty.

(iv) The rate of interest assessed on the unpaid balance beginning on the due date, as provided in §488.442.

e. In the case of a facility for which the provider agreement has been terminated and on which a civil money penalty was imposed on a per day basis, HCFA does or the State must send this penalty information after the—

(1) Final administrative decision is made;

(2) Facility has waived its right to a hearing in accordance with §488.436; or

(3) Time for requesting a hearing has expired and HCFA or the State has not received a hearing request from the facility.

f. Accrual of penalties when there is no immediate jeopardy. (1) In the case of noncompliance that does not pose immediate jeopardy, the daily accrual of per day civil money penalties is imposed for the days of noncompliance prior to the notice specified in §488.434 and an additional period of no longer than 6 months following the last day of the survey.

(2) After the period specified in paragraph (f)(1) of this section, if the facility has not achieved substantial compliance, HCFA terminates the provider agreement and the State may terminate the provider agreement.

g. (1) In a case when per day civil money penalties are imposed, when a facility has deficiencies that pose immediate jeopardy, HCFA does or the State must terminate the provider agreement within 23 calendar days after the last day of the survey if the immediate jeopardy remains.

(2) The accrual of the civil money penalty imposed on a per day basis stops on the day the provider agreement is terminated.

(h) (1) If an on-site revisit is necessary to confirm substantial compliance and the provider can supply documentation acceptable to HCFA or the State agency that substantial compliance was achieved on a date preceding the revisit, penalties imposed on a per day basis only accrue until that date of correction for which there is written credible evidence.

(2) If an on-site revisit is not necessary to confirm substantial compliance, penalties imposed on a per day basis only accrue until the date of correction for which HCFA or the State receives and accepts written credible evidence.

[59 FR 56243, Nov. 10, 1994, as amended at 64 FR 13361, Mar. 18, 1999]

§488.442 Civil money penalties: Due date for payment of penalty.

(a) When payments are due for a civil money penalty imposed on a per day basis—(1) After a final administrative decision. A civil money penalty payment is due 15 days after a final administrative decision is made when—

(i) The facility achieves substantial compliance before the final administrative decision; or

(ii) The effective date of termination occurs before the final administrative decision.

(b) (1) After a final administrative decision. A civil money penalty payment is due 15 days after a final administrative decision is made when—

(i) The facility achieves substantial compliance before the final administrative decision; or

(ii) The effective date of termination occurs before the final administrative decision.

§488.442 Civil money penalties: Due date for payment of penalty.

(a) When payments are due for a civil money penalty imposed on a per day basis—(1) After a final administrative decision. A civil money penalty payment is due 15 days after a final administrative decision is made when—

(i) The facility achieves substantial compliance before the final administrative decision; or

(ii) The effective date of termination occurs before the final administrative decision.

§488.442 Civil money penalties: Due date for payment of penalty.

(a) When payments are due for a civil money penalty imposed on a per day basis—(1) After a final administrative decision. A civil money penalty payment is due 15 days after a final administrative decision is made when—

(i) The facility achieves substantial compliance before the final administrative decision; or

(ii) The effective date of termination occurs before the final administrative decision.

§488.442 Civil money penalties: Due date for payment of penalty.

(a) When payments are due for a civil money penalty imposed on a per day basis—(1) After a final administrative decision. A civil money penalty payment is due 15 days after a final administrative decision is made when—

(i) The facility achieves substantial compliance before the final administrative decision; or

(ii) The effective date of termination occurs before the final administrative decision.
(2) When no hearing was requested. A civil money penalty payment is due 15 days after the time period for requesting a hearing has expired and a hearing request was not received when—
   (i) The facility achieved substantial compliance before the hearing request was due; or
   (ii) The effective date of termination occurs before the hearing request was due.
(3) After a request to waive a hearing. A civil money penalty payment is due 15 days after receipt of the written request to waive a hearing when—
   (i) The facility achieved substantial compliance before HCFA or the State received the written waiver of hearing; or
   (ii) The effective date of termination occurs before HCFA or the State received the written waiver of hearing.
(4) After substantial compliance is achieved. A civil money penalty payment is due 15 days after substantial compliance is achieved when—
   (i) The final administrative decision is made before the facility came into substantial compliance;
   (ii) The facility did not file a timely hearing request before it came into substantial compliance; or
   (iii) The facility waived its right to a hearing before it came into substantial compliance;
(5) After the effective date of termination. A civil money penalty payment is due 15 days after the effective date of termination, if before the effective date of termination—
   (i) The final administrative decision was made;
   (ii) The time for requesting a hearing has expired and the facility did not request a hearing; or
   (iii) The facility waived its right to a hearing.
(6) In the cases specified in paragraph (a)(4) of this section, the period of noncompliance may not extend beyond 6 months from the last day of the survey.

(b) When payments are due for a civil money penalty imposed for an instance of noncompliance. Payment of a civil money penalty is due 15 days after one of the following dates:
   (1) The final administrative decision is made;
   (2) The time for requesting a hearing has expired and the facility did not request a hearing; or
   (3) The facility waived its right to a hearing.

(c) Deduction of penalty from amount owed. The amount of the penalty, when determined, may be deducted from any sum then or later owing by HCFA or the State to the facility.

(d) Interest—(1) Assessment. Interest is assessed on the unpaid balance of the penalty, beginning on the due date.
   (2) Medicare interest. Medicare rate of interest is the higher of—
      (i) The rate fixed by the Secretary of the Treasury after taking into consideration private consumer rates of interest prevailing on the date of the notice of the penalty amount due (published quarterly in the FEDERAL REGISTER by HHS under 45 CFR 30.13(a)); or
      (ii) The current value of funds (published annually in the FEDERAL REGISTER by the Secretary of the Treasury, subject to quarterly revisions).
   (3) Medicaid interest. The interest rate for Medicaid is determined by the State.

(e) Penalties collected by HCFA. Civil money penalties and corresponding interest collected by HCFA from—
   (1) Medicare-participating facilities are deposited as miscellaneous receipts of the United States Treasury; and
   (2) Medicaid-participating facilities are returned to the State.

(f) Collection from dually participating facilities. Civil money penalties collected from dually participating facilities are deposited as miscellaneous receipts of the United States Treasury and returned to the State in proportion commensurate with the relative proportions of Medicare and Medicaid beds at the facility actually in use by residents covered by the respective programs on the date the civil money penalty begins to accrue.

(g) Penalties collected by the State. Civil money penalties collected by the State must be applied to the protection of the health or property of residents of facilities that the State or HCFA finds noncompliant, such as—
   (1) Payment for the cost of relocating residents to other facilities;
(2) State costs related to the operation of a facility pending correction of deficiencies or closure; and
(3) Reimbursement of residents for personal funds or property lost at a facility as a result of actions by the facility or by individuals used by the facility to provide services to residents.

[59 FR 56243, Nov. 10, 1994; 60 FR 50119, Sept. 28, 1995, as amended at 64 FR 13361, Mar. 18, 1999]

§ 488.444 Civil money penalties: Settlement of penalties.
(a) HCFA has authority to settle cases at any time prior to a final administrative decision for Medicare-only SNFs, State-operated facilities, or other facilities for which HCFA’s enforcement action prevails, in accordance with § 488.330.
(b) The State has the authority to settle cases at any time prior to the evidentiary hearing decision for all cases in which the State’s enforcement action prevails.

§ 488.450 Continuation of payments to a facility with deficiencies.
(a) Criteria. (1) HCFA may continue payments to a facility not in substantial compliance for the periods specified in paragraph (c) of this section if the following criteria are met:
(i) The State survey agency finds that it is more appropriate to impose alternative remedies than to terminate the facility;
(ii) The State has submitted a plan and timetable for corrective action approved by HCFA; and
(iii) The facility, in the case of a Medicare SNF, or the State, in the case of a Medicaid NF, agrees to repay the Federal government payments received under this provision if corrective action is not taken in accordance with the approved plan and timetable for corrective action.
(2) HCFA or the State may terminate the SNF or NF agreement before the end of the correction period if the criteria in paragraph (a)(1) of this section are not met.
(b) Cessation of payments. If termination is not sought, either by itself or along with another remedy or remedies, or any of the criteria set forth in paragraph (a)(1) of this section are not met or agreed to by either the facility or the State, the facility or State will receive no Medicare or Federal Medicaid payments, as applicable, from the last day of the survey.
(c) Period of continued payments. If the conditions in paragraph (a)(1) of this section are met, HCFA may continue payments to a Medicare facility or to the State for a Medicaid facility with noncompliance that does not constitute immediate jeopardy for up to 6 months from the last day of the survey.
(d) Failure to achieve substantial compliance. If the facility does not achieve substantial compliance by the end of the period specified in paragraph (c) of this section,
(1) HCFA will—
(i) Terminate the provider agreement of the Medicare SNF in accordance with § 488.456; or
(ii) Discontinue Federal funding to the SNF for Medicare; and
(iii) Discontinue FFP to the State for the Medicaid NF.
(2) The State may terminate the provider agreement for the NF.

[59 FR 56243, Nov. 10, 1994; 60 FR 50119, Sept. 28, 1995]

§ 488.452 State and Federal disagreements involving findings not in agreement in non-State operated NFs and dually participating facilities when there is no immediate jeopardy.
The following rules apply when HCFA and the State disagree over findings of noncompliance or application of remedies in a non-State operated NF or dually participating facility:
(a) Disagreement over whether facility has met requirements. (1) The State’s finding of noncompliance takes precedence when—
(i) HCFA finds that a NF or a dually participating facility is in substantial compliance with the participation requirements; and
(ii) The State finds that a NF or dually participating facility has not achieved substantial compliance.
(2) HCFA’s findings of noncompliance take precedence when—
(i) HCFA finds that a NF or a dually participating facility has not achieved substantial compliance; and

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(ii) The State finds that a NF or a dually participating facility is in substantial compliance with the participation requirements.

(3) When HCFA's survey findings take precedence, HCFA may—

(i) Impose any of the alternative remedies specified in §488.406;

(ii) Terminate the provider agreement subject to the applicable conditions of §488.450; and

(iii) Stop FFP to the State for a NF.

(b) Disagreement over decision to terminate. (1) HCFA’s decision to terminate the participation of a facility takes precedence when—

(i) Both HCFA and the State find that the facility has not achieved substantial compliance; and

(ii) HCFA, but not the State, finds that the facility’s participation should be terminated. HCFA will permit continuation of payment during the period prior to the effective date of termination not to exceed 6 months, if the applicable conditions of §488.450 are met.

(2) The State’s decision to terminate a facility’s participation and the procedures for appealing such termination, as specified in §431.153(c) of this chapter, takes precedence when—

(i) The State, but not HCFA, finds that a NF’s participation should be terminated; and

(ii) The State’s effective date for the termination of the NF’s provider agreement is no later than 6 months after the last day of survey.

(c) Disagreement over timing of termination of facility. The State’s timing of termination takes precedence if it does not occur later than 6 months after the last day of the survey when both HCFA and the State find that—

(1) A facility is not in substantial compliance; and

(2) The facility’s participation should be terminated.

(d) Disagreement over remedies. (1) When HCFA or the State, but not both, establishes one or more remedies, in addition to or as an alternative to termination, the additional or alternative remedies will also apply when—

(i) Both HCFA and the State find that a facility has not achieved substantial compliance; and

(ii) Both HCFA and the State find that no immediate jeopardy exists.

(2) Overlap of remedies. When HCFA and the State establish one or more remedies, in addition to or as an alternative to termination, only the HCFA remedies apply when both HCFA and the State find that a facility has not achieved substantial compliance.

(e) Regardless of whether HCFA’s or the State’s decision controls, only one noncompliance and enforcement decision is applied to the Medicaid agreement, and for a dually participating facility, that same decision will apply to the Medicare agreement.

§ 488.454 Duration of remedies.

(a) Except as specified in paragraphs (b) and (d) of this section, alternative remedies continue until—

(1) The facility has achieved substantial compliance, as determined by HCFA or the State based upon a revisit or after an examination of credible written evidence that it can verify without an on-site visit; or

(2) HCFA or the State terminates the provider agreement.

(b) In the cases of State monitoring and denial of payment imposed for repeated substandard quality of care, remedies continue until—

(1) HCFA or the State determines that the facility has achieved substantial compliance and is capable of remaining in substantial compliance; or

(2) HCFA or the State terminates the provider agreement.

(c) In the case of temporary management, the remedy continues until—

(1) HCFA or the State determines that the facility has achieved substantial compliance and is capable of remaining in substantial compliance; or

(2) HCFA or the State terminates the provider agreement; or

(3) The facility which has not achieved substantial compliance re-assumes management control. In this case, HCFA or the State initiates termination of the provider agreement and may impose additional remedies.

(d) In the case of a civil money penalty imposed for an instance of noncompliance, the remedy is the specific amount of the civil money penalty imposed for the particular deficiency.
§ 488.456 Termination of provider agreement.

(a) Effect of termination. Termination of the provider agreement ends—

(1) Payment to the facility; and

(2) Any alternative remedy.

(b) Basis for termination. (1) HCFA and the State may terminate a facility's provider agreement if a facility—

(i) Is not in substantial compliance with the requirements of participation, regardless of whether or not immediate jeopardy is present; or

(ii) Fails to submit an acceptable plan of correction within the timeframe specified by HCFA or the State.

(2) HCFA and the State terminate a facility's provider agreement if a facility—

(i) Fails to relinquish control to the temporary manager, if that remedy is imposed by HCFA or the State; or

(ii) Does not meet the eligibility criteria for continuation of payment as set forth in § 488.412(a)(1).

(c) Notice of termination. Before terminating a provider agreement, HCFA does and the State must notify the facility and the public—

(1) At least 2 calendar days before the effective date of termination for a facility with immediate jeopardy deficiencies; and

(2) At least 15 calendar days before the effective date of termination for a facility with non-immediate jeopardy deficiencies that constitute noncompliance.

(d) Procedures for termination. (1) HCFA terminates the provider agreement in accordance with procedures set forth in § 489.53 of this chapter; and

(2) The State must terminate the provider agreement of a NF in accordance with procedures specified in parts 431 and 442 of this chapter.
Subpart E—Termination of Agreement and Reinstatement After Termination

489.52 Termination by the provider.
489.53 Termination by HCFA.
489.54 Termination by the OIG.
489.55 Exceptions to effective date of termination.
489.57 Reinstatement after termination.

Subpart F—Surety Bond Requirements for HHAs

489.60 Definitions.
489.61 Basic requirement for surety bonds.
489.62 Requirement waived for Government-operated HHAs.
489.63 Parties to the bond.
489.64 Authorized Surety and exclusion of surety companies.
489.65 Amount of the bond.
489.66 Additional requirements of the surety bond.
489.67 Term and type of bond.
489.68 Effect of failure to obtain, maintain, and timely file a surety bond.
489.69 Evidence of compliance.
489.70 Effect of payment by the Surety.
489.71 Surety’s standing to appeal Medicare determinations.
489.72 Effect of review reversing HCFA’s determination.
489.73 Effect of conditions of payment.
489.74 Incorporation into existing provider agreements.

Subparts G–H [Reserved]

Subpart I—Advance Directives

489.100 Definition.
489.102 Requirements for providers.
489.104 Effective dates.

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

Source: 45 FR 22937, Apr. 4, 1980, unless otherwise noted.

Subpart A—General Provisions

§ 489.1 Statutory basis.

This part implements section 1866 of the Social Security Act. Section 1866 specifies the terms of provider agreements, the grounds for terminating a provider agreement, the circumstances under which payment for new admissions may be denied, and the circumstances under which payment may be withheld for failure to make timely utilization review. The following other sections of that Act are also pertinent:

(a) Section 1861 defines the services covered under Medicare and the providers that may be reimbursed for furnishing those services.

(b) Section 1864 provides for the use of State survey agencies to ascertain whether certain entities meet the conditions of participation.

(c) Section 1871 authorizes the Secretary to prescribe regulations for the administration of the Medicare program.

(d) Although section 1866 of the Act speaks only to providers and provider agreements, the effective date rules in this part are made applicable also to the approval of suppliers that meet the requirements specified in § 489.13.

(e) Section 1861(o)(7) of the Act requires each HHA to provide HCFA with a surety bond.


§ 489.2 Scope of part.

(a) Subpart A of this part sets forth the basic requirements for submittal and acceptance of a provider agreement under Medicare. Subpart B of this part specifies the basic commitments and limitations that the provider must agree to as part of an agreement to provide services. Subpart C specifies the limitations on allowable charges to beneficiaries for deductibles, coinsurance, copayments, blood, and services that must be part of the provider agreement. Subpart D of this part specifies how incorrect collections are to be handled. Subpart F sets forth the circumstances and procedures for denial of payments for new admissions and for withholding of payment as an alternative to termination of a provider agreement.

(b) The following providers are subject to the provisions of this part:

(1) Hospitals.
(2) Skilled nursing facilities (SNFs).
(3) Home health agencies (HHAs).
(4) Clinics, rehabilitation agencies, and public health agencies.
(5) Comprehensive outpatient rehabilitation facilities (CORFs).
(6) Hospices.
(7) Critical access hospital (CAHs).
(8) Community mental health centers (CMHCs).
(c)(1) Clinics, rehabilitation agencies, and public health agencies may enter
§ 489.3 Definitions.

For purposes of this part—

Immediate jeopardy means a situation in which the provider’s noncompliance with one or more requirements of participation has caused, or is likely to cause, serious injury, harm, impairment, or death to a resident.

Provider agreement means an agreement between HCFA and one of the providers specified in § 489.2(b) to provide services to Medicare beneficiaries and to comply with the requirements of section 1866 of the Act.

§ 489.10 Basic requirements.

(a) Any of the providers specified in § 489.2 may request participation in Medicare. In order to be accepted, it must meet the conditions of participation or requirements (for SNFs) set forth in this section and elsewhere in this chapter.

(b) In order to participate in the Medicare program, the provider must meet the applicable civil rights requirements of:

(1) Title VI of the Civil Rights Act of 1964, as implemented by 45 CFR part 80, which provides that no person in the United States shall, on the ground of race, color, or national origin, be excluded from participation in, be denied the benefits of, or be subject to discrimination under, any program or activity receiving Federal financial assistance;

(2) The Age Discrimination Act of 1975, as implemented by 45 CFR part 90, which is designed to prohibit discrimination on the basis of age in programs or activities receiving Federal financial assistance. The Age Discrimination Act also permits federally assisted programs and activities, and recipients of Federal funds, to continue to use certain age distinctions, and factors other than age, that meet the requirements of the Age Discrimination Act and 45 CFR part 90; and

(4) Other pertinent requirements of the Office of Civil Rights of HHS.

(c) In order for a hospital, SNF, HHA, or hospice to be accepted, it must also meet the advance directives requirements specified in subpart I of this part.

(d) The State survey agency will ascertain whether the provider meets the conditions of participation or requirements (for SNFs) and make its recommendations to HCFA.

(e) In order for a home health agency to be accepted, it must also meet the surety bond requirements specified in subpart F of this part.

(f) In order for a home health agency to be accepted as a new provider, it must also meet the capitalization requirements specified in subpart B of this part.

§ 489.11 Acceptance of a provider as a participant.

(a) Action by HCFA. If HCFA determines that the provider meets the requirements, it will send the provider—

(1) Written notice of that determination; and

(2) Two copies of the provider agreement.

(b) Action by provider. If the provider wishes to participate, it must return both copies of the agreement, duly signed by an authorized official, to HCFA, together with a written statement indicating whether it has been adjudged insolvent or bankrupt in any State or Federal court, or whether any insololvency or bankruptcy actions are pending.
§ 489.13 Effective date of agreement or approval.

(a) Applicability—(1) General rule. Except as provided in paragraph (a)(2) of this section, this section applies to Medicare provider agreements with, and supplier approval of, entities that, as a basis for participation in Medicare—

(i) Are subject to survey and certification by HCFA or the State survey agency; or

(ii) Are deemed to meet Federal requirements on the basis of accreditation by an accrediting organization whose program has HCFA approval at the time of accreditation survey and accreditation decision.

(2) Exceptions. (i) For an agreement with a community mental health center (CMHC) or a Federally qualified health center (FQHC), the effective date is the date on which HCFA accepts a signed agreement which assures that the CMHC or FQHC meets all Federal requirements.

(ii) A Medicare supplier approval of a laboratory is effective only while the laboratory has in effect a valid CLIA certificate issued under part 493 of this chapter, and only for the specialty and subspecialty tests it is authorized to perform.

(b) All Federal requirements are met on the date of survey. The agreement or approval is effective on the date the survey (including the Life Safety Code survey, if applicable) is completed, if on that date the provider or supplier meets all applicable Federal requirements as set forth in this chapter. (If the agreement or approval is time-limited, the new agreement or approval is effective on the day following expiration of the current agreement or approval.)

(c) All Federal requirements are not met on the date of survey. If on the date the survey is completed the provider or supplier fails to meet any of the requirements specified in paragraph (b) of this section, the following rules apply:

(1) For an agreement with an SNF, the effective date is the date on which—

(i) The SNF is in substantial compliance (as defined in § 488.301 of this chapter) with the requirements for participation; and

(ii) HCFA or the State survey agency receives from the SNF, if applicable, an approvable waiver request.

(2) For an agreement with, or an approval of, any other provider or supplier, (except those specified in paragraph (a)(2) of this section), the effective date is the earlier of the following:

(i) The date on which the provider or supplier meets all requirements.

(ii) The date on which a provider or supplier is found to meet all conditions of participation or coverage, but has lower level deficiencies, and HCFA or the State survey agency receives an acceptable plan of correction for the
§ 489.18 Change of ownership or leasing: Effect on provider agreement.

(a) What constitutes change of ownership—(1) Partnership. In the case of a partnership, the removal, addition, or substitution of a partner, unless the partners expressly agree otherwise, as permitted by applicable State law, constitutes change of ownership.

(2) Unincorporated sole proprietorship. Transfer of title and property to another party constitutes change of ownership.

(3) Corporation. The merger of the provider corporation into another corporation, or the consolidation of two or more corporations, resulting in the creation of a new corporation constitutes change of ownership. Transfer of corporate stock or the merger of another corporation into the provider corporation does not constitute change of ownership.

(4) Leasing. The lease of all or part of a provider facility constitutes change of ownership of the leased portion.

(b) Notice to HCFA. A provider who is contemplating or negotiating a 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 existing provider agreement will automatically be assigned to the new owner.

(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) Any existing plan of correction.
(2) Compliance with applicable health and safety standards.
(3) Compliance with the ownership and financial interest disclosure requirements of part 420, subpart C, of this chapter.
(4) Compliance with civil rights requirements set forth in 45 CFR Parts 80, 84, and 90.

(e) Effect of leasing. The provider agreement will be assigned to the lessee only to the extent of the leased portion of the facility.

Subpart B—Essentials of Provider Agreements

§ 489.20 Basic commitments.

The provider agrees to the following:

(a) To limit its charges to beneficiaries and to other individuals on their behalf, in accordance with provisions of subpart C of this part.
(b) To comply with the requirements of subpart D of this part for the return or other disposition of any amounts incorrectly collected from a beneficiary or any other person in his or her behalf.
(c) To comply with the requirements of §420.203 of this chapter when it hires certain former employees of intermediaries.
(d) In the case of a hospital or a CAH that furnishes services to Medicare beneficiaries, either to furnish directly or to make arrangements (as defined in §409.3 of this chapter) for all Medicare-covered services to inpatients and outpatients of a hospital or a CAH except the following:
   (1) Physicians’ services that meet the criteria of §415.102(a) of this chapter for payment on a reasonable charge basis.
   (2) Physician assistant services, as defined in section 1861(s)(2)(K)(i) of the Act, that are furnished after December 31, 1990.
   (3) Nurse practitioner and clinical nurse specialist services, as defined in section 1861(s)(2)(K)(ii) of the Act.
   (4) Certified nurse-midwife services, as defined in section 1861(ff) of the Act, that are furnished after December 31, 1990.
   (5) Qualified psychologist services, as defined in section 1861(ii) of the Act, that are furnished after December 31, 1990.
   (6) Services of an anesthetist, as defined in §410.69 of this chapter.
   (e) In the case of a hospital or CAH that furnishes inpatient hospital services or inpatient CAH services for which payment may be made under Medicare, to maintain an agreement with a PRO for that organization to review the admissions, quality, appropriateness, and diagnostic information related to those inpatient services. The requirement of this paragraph (e) applies only if, for the area in which the hospital or CAH is located, there is a PRO that has a contract with HCFA under part B of title XI of the Act.
   (f) To maintain a system that, during the admission process, identifies any primary payers other than Medicare, so that incorrect billing and Medicare overpayments can be prevented.
   (g) To bill other primary payers before billing Medicare except when the primary payer is a liability insurer and except as provided in paragraph (j) of this section.
   (h) If the provider receives payment for the same services from Medicare and another payer that is primary to Medicare, to reimburse Medicare any overpaid amount within 60 days.
   (i) If the provider receives, from a payer that is primary to Medicare, a payment that is reduced because the provider failed to file a proper claim—
      (1) To bill Medicare for an amount no greater than would have been payable as secondary payment if the primary insurer’s payment had been based on a proper claim; and
      (2) To charge the beneficiary only: (i) The amount it would have been entitled to charge if it had filed a proper claim and received payment based on such a claim; and
      (ii) An amount equal to any third party payment reduction attributable to failure to file a proper claim, but only if the provider can show that—
         (A) It failed to file a proper claim solely because the beneficiary, for any reason other than mental or physical incapacity, failed to give the provider the necessary information; or
         (B) The beneficiary, who was responsible for filing a proper claim, failed to do so for any reason other than mental or physical incapacity.
   (j) In the State of Oregon, because of a court decision, and in the absence of a reversal on appeal or a statutory clarification overturning the decision, hospitals may bill liability insurers first. However, if the liability insurer does not pay “promptly”, as defined in §411.50 of this chapter, the hospital must withdraw its claim or lien and bill Medicare for covered services.
   (k) In the case of home health agencies that provide home health services to Medicare beneficiaries under subpart E of part 409 and subpart C of part 410 of this chapter, to offer to furnish catheters, catheter supplies, ostomy bags, and supplies related to ostomy care to any individual who requires them as part of their furnishing of home health services.
   (l) In the case of a hospital as defined in §489.24(b) to comply with §489.24.
   (m) In the case of a hospital as defined in §489.24(b), to report to HCFA
or the State survey agency any time it has reason to believe it may have received an individual who has been transferred in an unstable emergency medical condition from another hospital in violation of the requirements of §489.24(d).

(n) In the case of inpatient hospital services, to participate in any health plan contracted for under 10 U.S.C. 1079 or 1086 or 38 U.S.C. 613, in accordance with §489.25.

(o) In the case of inpatient hospital services, to admit veterans whose admission has been authorized under 38 U.S.C. 603, in accordance with §489.26.

(p) In the case of a hospital as defined in §489.24(b)—

(1) To post conspicuously in any emergency department or in a place or places likely to be noticed by all individuals entering the emergency department, as well as those individuals waiting for examination and treatment in areas other than traditional emergency departments (that is, entrance, admitting area, waiting room, treatment area), a sign (in a form specified by the Secretary) specifying rights of individuals under Section 1867 of the Act with respect to examination and treatment for emergency medical conditions and women in labor; and

(2) To post conspicuously (in a form specified by the Secretary) information indicating whether or not the hospital or rural primary care hospital participates in the Medicare program under title XIX.

(q) In the case of a hospital as defined in §489.24(b)—

(1) To post conspicuously in any emergency department or in a place or places likely to be noticed by all individuals entering the emergency department, as well as those individuals waiting for examination and treatment in areas other than traditional emergency departments (that is, entrance, admitting area, waiting room, treatment area), a sign (in a form specified by the Secretary) specifying rights of individuals under Section 1867 of the Act with respect to examination and treatment for emergency medical conditions and women in labor; and

(2) To post conspicuously (in a form specified by the Secretary) information indicating whether or not the hospital or rural primary care hospital participates in the Medicaid program under a State plan approved under Title XIX.

(r) In the case of a hospital as defined in §489.24(b) (including both the transferring and receiving hospitals), to maintain—

(1) Medical and other records related to individuals transferred to or from the hospital for a period of 5 years from the date of the transfer;

(2) A list of physicians who are on call for duty after the initial examination to provide treatment necessary to stabilize an individual with an emergency medical condition; and

(3) A central log on each individual who comes to the emergency department, as defined in §489.24(b), seeking assistance and whether he or she refused treatment, was refused treatment, or whether he or she was transferred, admitted and treated, stabilized and transferred, or discharged.

(s) In the case of an SNF, either to furnish directly or make arrangements (as defined in §409.3 of this chapter) for all Medicare-covered services furnished to a resident (as defined in §411.15(p)(3) of this chapter) of the SNF, except the following:

(1) Physicians’ services that meet the criteria of §415.102(a) of this chapter for payment on a fee schedule basis.

(2) Services performed under a physician’s supervision by a physician assistant who meets the applicable definition in section 1861(aa)(5) of the Act.

(3) 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.

(4) Services performed by a certified nurse-midwife, as defined in section 1861(gg) of the Act.

(5) Services performed by a qualified psychologist, as defined in section 1861(ii) of the Act.

(6) Services performed by a certified registered nurse anesthetist, as defined in section 1861(bb) of the Act.

(7) 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.

(8) Erythropoietin (EPO) for dialysis patients, as defined in section 1861(s)(2)(O) of the Act.

(9) Hospice care, as defined in section 1861(dd) of the Act.

(10) 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 §411.15(p)(3)(i) through (p)(3)(iv) of this chapter as ending the individual’s status as an SNF resident.

(11) The transportation costs of electrocardiogram equipment (HCPCS code R0076), but only with respect to those individuals...
§ 489.21 Specific limitations on charges.

Except as specified in subpart C of this part, the provider agrees not to charge a beneficiary for any of the following:

(a) Services for which the beneficiary is entitled to have payment made under Medicare.

(b) Services for which the beneficiary would be entitled to have payment made if the provider—
   (1) Had in its files the required certification and recertification by a physician relating to the services furnished to the beneficiary;
   (2) Had furnished the information required by the intermediary in order to determine the amount due the provider on behalf of the individual for the period with respect to which payment is to be made or any prior period;
   (3) Had complied with the provisions requiring timely utilization review of long stay cases so that a limitation on days of service has not been imposed under section 1866(d) of the Act (see subpart K of part 405 and part 482 of this chapter for utilization review requirements); and
   (4) Had obtained, from the beneficiary or a person acting on his or her behalf, a written request for payment to be made to the provider, and had properly filed that request. (If the beneficiary or person on his or her behalf refuses to execute a written request, the provider may charge the beneficiary for all services furnished to him or her.)

(c) Inpatient hospital services furnished to a beneficiary who exhausted his or her Part A benefits, if HCFA reimburses the provider for those services.

(d) Custodial care and services not reasonable and necessary for the diagnosis or treatment of illness or injury, if—
   (1) The beneficiary was without fault in incurring the expenses; and
   (2) The determination that payment was incorrect was not made until after the third year following the year in which the payment notice was sent to the beneficiary.

(e) Reimbursement of hospital services furnished to a beneficiary who exhausted his or her Part A benefits, if HCFA reimburses the provider for those services.

(f) Items and services furnished to a hospital inpatient (other than physicians’ services as described in §415.102(a) of this chapter or the services of an anesthetist as described in §405.553(b)(4) of this chapter) for which Medicare payment would be made if furnished by the hospital or by other providers or suppliers under arrangements made with them by the hospital. For this purpose, a charge by another provider or supplier for such an item or service is treated as a charge by the hospital for the item or service, and is also prohibited.
(g) Items and services furnished in connection with the implantation of cardiac pacemakers or pacemaker leads when HCFA denies payment for those devices under §409.19 or §410.64 of this chapter.

(h) Items and services (other than those described in §489.20(s)(1) through (15)) furnished to a resident (as defined in §411.15(p)(3) of this chapter) of an SNF for which Medicare payment would be made if furnished by the SNF or by other providers or suppliers under arrangements made with them by the SNF. For this purpose, a charge by another provider or supplier for such an item or service is treated as a charge by the SNF for the item or service, and is also prohibited.


§ 489.22 Special provisions applicable to prepayment requirements.

(a) A provider may not require an individual entitled to hospital insurance benefits to prepay in part or in whole for inpatient services as a condition of admittance as an inpatient, except where it is clear upon admission that payment under Medicare, Part A cannot be made.

(b) A provider may not deny covered inpatient services to an individual entitled to have payment made for those services on the ground of inability or failure to pay a requested amount at or before admission.

(c) A provider may not evict, or threaten to evict, an individual for inability to pay a deductible or a coinsurance amount required under Medicare.

(d) A provider may not charge an individual for (1) its agreement to admit or readmit the individual on some specified future date for covered inpatient services; or (2) for failure to remain an inpatient for any agreed-upon length of time or for failure to give advance notice of departure from the provider’s facilities.

§ 489.23 Specific limitation on charges for services provided to certain enrollees of fee-for-service FEHB plans.

A provider that furnishes inpatient hospital services to a retired Federal worker age 65 or older who is enrolled in a fee-for-service FEHB plan and who is not covered under Medicare Part A, must accept, as payment in full, an amount that approximates as closely as possible the Medicare inpatient hospital prospective payment system (PPS) rate established under part 412. The payment to the provider is composed of a payment from the FEHB plan and a payment from the enrollee. This combined payment approximates the Medicare PPS rate. The payment from the FEHB plan approximates, as closely as possible, the Medicare PPS rate minus any applicable enrollee deductible, coinsurance, or copayment amount. The payment from the enrollee is equal to the applicable deductible, coinsurance, or copayment amount.


§ 489.24 Special responsibilities of Medicare hospitals in emergency cases.

(a) General. In the case of a hospital that has an emergency department, if any individual (whether or not eligible for Medicare benefits and regardless of ability to pay) comes by himself or herself or with another person to the emergency department and a request is made on the individual's behalf for examination or treatment of a medical condition by qualified medical personnel (as determined by the hospital in its rules and regulations), the hospital must provide for an appropriate medical screening examination within the capability of the hospital’s emergency department, including ancillary services routinely available to the emergency department, to determine whether or not an emergency medical condition exists. The examinations must be conducted by individuals determined qualified by hospital by-laws or rules and regulations and who meet the requirements of §482.55 concerning
emergency services personnel and direction.

(b) Definitions. As used in this subpart—

Capacity means the ability of the hospital to accommodate the individual requesting examination or treatment of the transferred individual. Capacity encompasses such things as numbers and availability of qualified staff, beds and equipment and the hospital’s past practices of accommodating additional patients in excess of its occupancy limits.

Comes to the emergency department means, with respect to an individual requesting examination or treatment, that the individual is on the hospital property. For purposes of this section, “property” means the entire main hospital campus as defined in §413.65(b) of this chapter, including the parking lot, sidewalk, and driveway, as well as any facility or organization that is located off the main hospital campus but has been determined under §413.65 of this chapter to be a department of the hospital. Property also includes ambulances owned and operated by the hospital even if the ambulance is not on hospital grounds. An individual in a nonhospital-owned ambulance on hospital property is considered to have come to the hospital’s emergency department. An individual in a nonhospital-owned ambulance off hospital property is not considered to have come to the hospital’s emergency department even if a member of the ambulance staff contacts the hospital by telephone or telemetry communications and informs the hospital that they want to transport the individual to the hospital for examination and treatment. In these situations, the hospital may deny access if it is in “diversionary status,” that is, it does not have the staff or facilities to accept any additional emergency patients. If, however, the ambulance staff disregards the hospital’s instructions and transports the individual to hospital property, the individual is considered to have come to the emergency department.

Emergency medical condition means—

(i) A medical condition manifesting itself by acute symptoms of sufficient severity (including severe pain, psychiatric disturbances and/or symptoms of substance abuse) such that the absence of immediate medical attention could reasonably be expected to result in—

(A) Placing the health of the individual (or, with respect to a pregnant woman, the health of the woman or her unborn child) in serious jeopardy;

(B) Serious impairment to bodily functions; or

(C) Serious dysfunction of any bodily organ or part; or

(ii) With respect to a pregnant woman who is having contractions—

(A) That there is inadequate time to effect a safe transfer to another hospital before delivery; or

(B) That transfer may pose a threat to the health or safety of the woman or the unborn child.

Hospital includes a critical access hospital as defined in section 1861(mm)(1) of the Act.

Hospital with an emergency department means a hospital that offers services for emergency medical conditions (as defined in this paragraph) within its capability to do so.

Labor means the process of childbirth beginning with the latent or early phase of labor and continuing through the delivery of the placenta. A woman experiencing contractions is in true labor unless a physician certifies that, after a reasonable time of observation, the woman is in false labor.

Participating hospital means (i) a hospital or (ii) a critical access hospital as defined in section 1861(mm)(1) of the Act that has entered into a Medicare provider agreement under section 1866 of the Act.

Stabilized means, with respect to an “emergency medical condition” as defined in this section under paragraph (i) of that definition, that no material deterioration of the condition is likely, within reasonable medical probability, to result from or occur during the transfer of the individual from a facility or, with respect to an “emergency medical condition” as defined in this section under paragraph (ii) of that definition, that the woman has delivered the child and the placenta.
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To stabilize means, with respect to an “emergency medical condition” as defined in this section under paragraph (i) of that definition, to provide such medical treatment of the condition necessary to assure, within reasonable medical probability, that no material deterioration of the condition is likely to result from or occur during the transfer of the individual from a facility or that, with respect to an “emergency medical condition” as defined in this section under paragraph (ii) of that definition, the woman has delivered the child and the placenta.

Transfer means the movement (including the discharge) of an individual outside a hospital's facilities at the direction of any person employed by (or affiliated or associated, directly or indirectly, with) the hospital, but does not include such a movement of an individual who (i) has been declared dead, or (ii) leaves the facility without the permission of any such person.

(c) Necessary stabilizing treatment for emergency medical conditions—(1) General. If any individual (whether or not eligible for Medicare benefits) comes to a hospital and the hospital determines that the individual has an emergency medical condition, the hospital must provide either—

(i) Within the capabilities of the staff and facilities available at the hospital, for further medical examination and treatment as required to stabilize the medical condition; or

(ii) For transfer of the individual to another medical facility in accordance with paragraph (d) of this section.

(2) Refusal to consent to treatment. A hospital meets the requirements of paragraph (c)(1)(i) of this section with respect to an individual if the hospital offers the individual the further medical examination and treatment described in that paragraph and informs the individual (or a person acting on his or her behalf) of the risks and benefits to the individual of the proposed transfer that was refused by or on behalf of the individual.

(d) Restricting transfer until the individual is stabilized—(1) General. If an individual at a hospital has an emergency medical condition that has not been stabilized (as defined in paragraph (b) of this section), the hospital may not transfer the individual unless—

(i) The transfer is an appropriate transfer (within the meaning of paragraph (d)(2) of this section); and

(ii)(A) The individual (or a legally responsible person acting on the individual's behalf) requests the transfer, after being informed of the hospital's obligations under this section and of the risk of transfer. The request must be in writing and indicate the reasons for the request as well as indicate that
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he or she is aware of the risks and benefits of the transfer;

(B) A physician (within the meaning of section 1861(r)(1) of the Act) has signed a certification that, based upon the information available at the time of transfer, the medical benefits reasonably expected from the provision of appropriate medical treatment at another medical facility outweigh the increased risks to the individual or, in the case of a woman in labor, to the woman or the unborn child, from being transferred. The certification must contain a summary of the risks and benefits upon which it is based; or

(C) If a physician is not physically present in the emergency department at the time an individual is transferred, a qualified medical person (as determined by the hospital in its bylaws or rules and regulations) has signed a certification described in paragraph (d)(1)(ii)(B) of this section after a physician (as defined in section 1861(r)(1) of the Act) in consultation with the qualified medical person, agrees with the certification and subsequently countersigns the certification. The certification must contain a summary of the risks and benefits upon which it is based.

(2) A transfer to another medical facility will be appropriate only in those cases in which—

(i) The transferring hospital provides medical treatment within its capacity that minimizes the risks to the individual’s health and, in the case of a woman in labor, the health of the unborn child;

(ii) The receiving facility—

(A) Has available space and qualified personnel for the treatment of the individual; and

(B) Has agreed to accept transfer of the individual and to provide appropriate medical treatment;

(iii) The transferring hospital sends to the receiving facility all medical records (or copies thereof) related to the emergency condition which the individual has presented that are available at the time of the transfer, including available history, records related to the individual’s emergency medical condition, observations of signs or symptoms, preliminary diagnosis, results of diagnostic studies or telephone reports of the studies, treatment provided, results of any tests and the informed written consent or certification (or copy thereof) required under paragraph (d)(1)(ii) of this section, and the name and address of any on-call physician (described in paragraph (f) of this section) who has refused or failed to appear within a reasonable time to provide necessary stabilizing treatment. Other records (e.g., test results not yet available or historical records not readily available from the hospital’s files) must be sent as soon as practicable after transfer; and

(iv) The transfer is effected through qualified personnel and transportation equipment, as required, including the use of necessary and medically appropriate life support measures during the transfer.

(3) A participating hospital may not penalize or take adverse action against a physician or a qualified medical person described in paragraph (d)(1)(ii)(C) of this section because the physician or qualified medical person refuses to authorize the transfer of an individual with an emergency medical condition that has not been stabilized, or against any hospital employee because the employee reports a violation of a requirement of this section.

(e) Recipient hospital responsibilities. A participating hospital that has specialized capabilities or facilities (including, but not limited to, facilities such as burn units, shock-trauma units, neonatal intensive care units, or (with respect to rural areas) regional referral centers) may not refuse to accept from a referring hospital within the boundaries of the United States an appropriate transfer of an individual who requires such specialized capabilities or facilities if the receiving hospital has the capacity to treat the individual.

(f) Termination of provider agreement. If a hospital fails to meet the requirements of paragraph (a) through (e) of this section, HCFA may terminate the provider agreement in accordance with § 489.53.

(g) Consultation with Peer Review Organizations (PROs)—(1) General. Except as provided in paragraph (g)(3) of this
section, in cases where a medical opinion is necessary to determine a physician’s or hospital’s liability under section 1867(d)(1) of the Act, HCFA requests the appropriate PRO (with a contract under Part B of title XI of the Act) to review the alleged section 1867(d) violation and provide a report on its findings in accordance with paragraph (g)(2)(iv) and (v) of this section. HCFA provides to the PRO all information relevant to the case and within its possession or control. HCFA, in consultation with the OIG, also provides to the PRO a list of relevant questions to which the PRO must respond in its report.

(2) Notice of review and opportunity for discussion and additional information. The PRO shall provide the physician and hospital reasonable notice of its review, a reasonable opportunity for discussion, and an opportunity for the physician and hospital to submit additional information before issuing its report. When a PRO receives a request for consultation under paragraph (g)(1) of this section, the following provisions apply—

(i) The PRO reviews the case before the 15th calendar day and makes its tentative findings.

(ii) Within 15 calendar days of receiving the case, the PRO gives written notice, sent by certified mail, return receipt requested, to the physician or the hospital (or both if applicable).

(iii)(A) The written notice must contain the following information:

(1) The name of each individual who may have been the subject of the alleged violation.

(2) The date on which each alleged violation occurred.

(3) An invitation to meet, either by telephone or in person, to discuss the case with the PRO, and to submit additional information to the PRO within 30 calendar days of receipt of the notice, and a statement that these rights will be waived if the invitation is not accepted. The PRO must receive the information and hold the meeting within the 30-day period.

(4) A copy of the regulations at 42 CFR 489.24.

(B) For purposes of paragraph (g)(2)(iii)(A) of this section, the date of receipt is presumed to be 5 days after the certified mail date on the notice, unless there is a reasonable showing to the contrary.

(iv) The physician or hospital (or both where applicable) may request a meeting with the PRO. This meeting is not designed to be a formal adversarial hearing or a mechanism for discovery by the physician or hospital. The meeting is intended to afford the physician and/or the hospital a full and fair opportunity to present the views of the physician and/or hospital regarding the case. The following provisions apply to that meeting:

(A) The physician and/or hospital has the right to have legal counsel present during that meeting. However, the PRO may control the scope, extent, and manner of any questioning or any other presentation by the attorney. The PRO may also have legal counsel present.

(B) The PRO makes arrangements so that, if requested by HCFA or the OIG, a verbatim transcript of the meeting may be generated. If HCFA or OIG requests a transcript, the affected physician and/or the affected hospital may request that HCFA provide a copy of the transcript.

(C) The PRO affords the physician and/or the hospital an opportunity to present, with the assistance of counsel, expert testimony in either oral or written form on the medical issues presented. However, the PRO may reasonably limit the number of witnesses and length of such testimony if such testimony is irrelevant or repetitive. The physician and/or hospital, directly or through counsel, may disclose patient records to potential expert witnesses without violating any non-disclosure requirements set forth in part 476 of this chapter.

(D) The PRO is not obligated to consider any additional information provided by the physician and/or the hospital after the meeting, unless, before the end of the meeting, the PRO requests that the physician and/or hospital submit additional information to support the claims. The PRO then allows the physician and/or the hospital an additional period of time, not to exceed 5 calendar days from the meeting, to submit the relevant information to the PRO.
(v) Within 60 calendar days of receiving the case, the PRO must submit to HCFA a report on the PRO’s findings. HCFA provides copies to the OIG and to the affected physician and/or the affected hospital. The report must contain the name of the physician and/or the hospital, the name of the individual, and the dates and times the individual arrived at and was transferred (or discharged) from the hospital. The report provides expert medical opinion regarding whether the individual involved had an emergency medical condition, whether the individual’s emergency medical condition was stabilized, whether the individual was transferred appropriately, and whether there were any medical utilization or quality of care issues involved in the case.

(vi) The report required under paragraph (g)(2)(v) of this section should not state an opinion or conclusion as to whether section 1867 of the Act or §489.24 has been violated.

(3) If a delay would jeopardize the health or safety of individuals or when there was no screening examination, the PRO review described in this section is not required before the OIG may impose civil monetary penalties or an exclusion in accordance with section 1867(d)(1) of the Act and 42 CFR part 1003 of this title.

(4) If the PRO determines after a preliminary review that there was an appropriate medical screening examination and the individual did not have an emergency medical condition, as defined by paragraph (b) of this section, then the PRO may, at its discretion, return the case to HCFA and not meet the requirements of paragraph (g) except for those in paragraph (g)(2)(v).

(h) Release of PRO assessments. Upon request, HCFA may release a PRO assessment to the physician and/or hospital, or the affected individual, or his or her representative. The PRO physician’s identity is confidential unless he or she consents to its release. (See §§476.132 and 476.133 of this chapter.)

(i) Off-campus departments. If an individual comes to a facility or organization that is located off the main hospital campus but has been determined under §416.35 of this chapter to be a department of the hospital and a request is made on the individual’s behalf for examination or treatment of a potential emergency medical condition as otherwise described in paragraph (a) of this section, the hospital is obligated in accordance with the rules in this paragraph to provide the individual with an appropriate medical screening examination and any necessary stabilizing treatment or an appropriate transfer.

(1) Capability of the hospital. The capability of the hospital includes that of the hospital as a whole, not just the capability of the off-campus department. Except for cases described in paragraph (i)(3)(ii) of this section, the obligation of a hospital under this section must be discharged within the hospital as a whole. However, the hospital is not required to locate additional personnel or staff to off-campus departments to be on standby for possible emergencies.

(2) Protocols for off-campus departments. The hospital must establish protocols for the handling of individuals with potential emergency conditions at off-campus departments. These protocols must provide for direct contact between personnel at the off-campus department and emergency personnel at the main hospital campus and may provide for dispatch of practitioners, when appropriate, from the main hospital campus to the off-campus department to provide screening or stabilization services.

(i) If the off-campus department is an urgent care center, primary care center, or other facility that is routinely staffed by physicians, RNs, or LPNs, these department personnel must be trained, and given appropriate protocols, for the handling of emergency cases. At least one individual on duty at the off-campus department during its regular hours of operation must be designated as a qualified medical person as described in paragraph (d) of this section. The qualified medical person must initiate screening of individuals who come to the off-campus department with a potential emergency medical condition, and may be able to complete the screening and provide any necessary stabilizing treatment at the off-campus department, or to arrange an appropriate transfer.

(ii) If the off-campus department is a physical therapy, radiology, or other
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Facility not routinely staffed with physicians, RNs, or LPNs, the department’s personnel must be given protocols that direct them to contact emergency personnel at the main hospital campus for direction. Under this direction, and in accordance with protocols established in advance by the hospital, the personnel at the off-campus department must describe patient appearance and report symptoms and, if appropriate, either arrange transportation of the individual to the main hospital campus in accordance with paragraph (i)(3)(i) of this section or assist in an appropriate transfer as described in paragraphs (i)(3)(ii) and (d)(2) of this section.

(3) Movement or appropriate transfer from off-campus departments—(i) If the main hospital campus has the capability required by the individual and movement of the individual to the main campus would not significantly jeopardize the life or health of the individual, the personnel at the off-campus department must assist in arranging this movement. Movement of the individual to the main campus of the hospital is not considered a transfer under this section, since the individual is simply being moved from one department of a hospital to another department or facility of the same hospital.

(ii) If transfer of an individual with a potential emergency condition to a medical facility other than the main hospital campus is warranted, either because the main hospital campus does not have the specialized capability or facilities required by the individual, or because the individual’s condition is deteriorating so rapidly that taking the time needed to move the individual to the main hospital campus would significantly jeopardize the life or health of the individual, personnel at the off-campus department must, in accordance with protocols established in advance by the hospital, assist in arranging an appropriate transfer of the individual to a medical facility other than the main hospital. The protocols must include procedures and agreements established in advance with other hospitals or medical facilities in the area of the off-campus department to facilitate these appropriate transfers. Such a transfer would require—

(A) That there be either a request by or on behalf of the individual as described in paragraph (d)(1)(ii)(A) of this section or a certification by a physician or a qualified medical person as described in paragraph (d)(1)(ii)(B) or (d)(1)(ii)(C) of this section; and

(B) That the transfer comply with the requirements described in paragraph (d)(2) of this section.

(iii) If the individual is being appropriately transferred to another medical facility from the off-campus department, the requirement for the provision of medical treatment in paragraph (d)(2)(ii) of this section would be met by provision of medical treatment within the capability of the transferring off-campus department.


EFFECTIVE DATE NOTES: 1. At 59 FR 32120, June 22, 1994, §489.24 was added. Paragraphs (d) and (g) contain information collection and recordkeeping requirements and will not become effective until approved by the Office of Management and Budget.

2. At 65 FR 18548, Apr. 7, 2000, §489.24 was amended by revising in paragraph (b) the definition of Comes to the emergency department and adding paragraph (i), effective Oct. 10, 2000. At 65 FR 58919, Oct. 3, 2000, the effective date was delayed until Jan. 10, 2001. For the convenience of the user, the superseded text is set forth as follows:

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* * * * *

(b) * * *

Comes to the emergency department means, with respect to an individual requesting examination or treatment, that the individual is on the hospital property (property includes ambulances owned and operated by the hospital, even if the ambulance is not on hospital grounds). An individual in a nonhospital-owned ambulance on hospital property is not considered to have come to the hospital’s emergency department. An individual in a nonhospital-owned ambulance off hospital property is considered to have come to the hospital’s emergency department even if a member of the ambulance staff contacts the hospital by telephone or telemetry communications and informs the hospital that they want to transport the individual to the hospital for examination and treatment. In such situations, the hospital may deny access if it is in “diversionary status,” that is,
§ 489.25 Special requirements concerning CHAMPUS and CHAMPVA programs.

For inpatient services, a hospital that participates in the Medicare program must participate in any health plan contracted under 10 U.S.C. 1079 or 1086 (Civilian Health and Medical Program of the Uniformed Services) and under 38 U.S.C. 613 (Civilian Health and Medical Program of the Veterans Administration) and accept the CHAMPUS/CHAMPVA-determined allowable amount as payment in full, less applicable deductible, patient cost-share, and noncovered items. Hospitals must meet the requirements of 32 CFR part 199 concerning program benefits under the Department of Defense. This section applies to inpatient services furnished to beneficiaries admitted on or after January 1, 1987.

[59 FR 32123, June 22, 1994]

§ 489.26 Special requirements concerning veterans.

For inpatient services, a hospital that participates in the Medicare program must admit any veteran whose admission is authorized by the Department of Veterans Affairs under 38 U.S.C. 603 and must meet the requirements of 32 CFR part 17 concerning admissions practices and payment methodology and amounts. This section applies to services furnished to veterans admitted on and after July 1, 1987.

[59 FR 32123, June 22, 1994]

§ 489.27 Beneficiary notice of discharge rights.

A hospital that participates in the Medicare program must furnish each Medicare beneficiary, or an individual acting on his or her behalf, the notice of discharge rights HCFA supplies to the hospital to implement section 1866(a)(1)(M) of the Act. The hospital must provide timely notice during the course of the hospital stay. For purposes of this paragraph, the course of the hospital stay may begin with the provision of a package of information regarding scheduled preadmission testing and registration for a planned hospital admission. The hospital must be able to demonstrate compliance with this requirement.


§ 489.28 Special capitalization requirements for HHAs.

(a) Basic rule. An HHA entering the Medicare program on or after January 1, 1998, including a new HHA as a result of a change of ownership, if the change of ownership results in a new provider number being issued, must have available sufficient funds, which we term "initial reserve operating funds," to operate the HHA for the three month period after its Medicare provider agreement becomes effective, exclusive of actual or projected accounts receivable from Medicare or other health care insurers.

(b) Standard. Initial reserve operating funds are sufficient to meet the requirement of this section if the total amount of such funds is equal to or greater than the product of the actual average cost per visit of three or more similarly situated HHAs in their first year of operation (selected by HCFA for comparative purposes) multiplied by the number of visits reported by the comparison HHAs—whichever is greater.

(c) Method. HCFA, through the intermediary, will determine the amount of the initial reserve operating funds using reported cost and visit data from submitted cost reports for the first full year of operation from at least three HHAs that the intermediary serves that are comparable to the HHA that is seeking to enter the Medicare program, considering such factors as geographic location and urban/rural status, number of visits, provider-based versus free-standing, and proprietary versus non-proprietary status. The determination of the adequacy of the required initial reserve operating funds is based...
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on the average cost per visit of the comparable HHAs, by dividing the sum of total reported costs of the HHAs in their first year of operation by the sum of the HHAs' total reported visits. The resulting average cost per visit is then multiplied by the projected visits for the first three months of operation of the HHA seeking to enter the program, but not less than 90 percent of average visits for a three month period for the HHAs used in determining the average cost per visit.

(d) Required proof of availability of initial reserve operating funds. The HHA must provide HCFA with adequate proof of the availability of initial reserve operating funds. Such proof, at a minimum, will include a copy of the statement(s) of the HHA's savings, checking, or other account(s) that contains the funds, accompanied by an attestation from an officer of the bank or other financial institution that the funds are in the account(s) and that the funds are immediately available to the HHA. In some cases, an HHA may have all or part of the initial reserve operating funds in cash equivalents. For the purpose of this section, cash equivalents are short-term, highly liquid investments that are readily convertible to known amounts of cash and that present insignificant risk of changes in value. A cash equivalent that is not readily convertible to a known amount of cash as needed during the initial three month period for which the initial reserve operating funds are required does not qualify in meeting the initial reserve operating funds requirement. Examples of cash equivalents for the purpose of this section are Treasury bills, commercial paper, and money market funds. As with funds in a checking, savings, or other account, the HHA also must be able to document the availability of any cash equivalents. HCFA later may require the HHA to furnish another attestation from the financial institution that the funds remain available, or, if applicable, documentation from the HHA that any cash equivalents remain available, until a date when the HHA will have been surveyed by the State agency or by an approved accrediting organization. The officer of the HHA who will be certifying the accuracy of the information on the HHA's cost report must certify what portion of the required initial reserve operating funds is non-borrowed funds, including funds invested in the business by the owner. That amount must be at least 50 percent of the required initial reserve operating funds. The remainder of the reserve operating funds may be secured through borrowing or line of credit from an unrelated lender.

(e) Borrowed funds. If borrowed funds are not in the same account(s) as the HHA's own non-borrowed funds, the HHA also must provide proof that the borrowed funds are available for use in operating the HHA, by providing, at a minimum, a copy of the statement(s) of the HHA's savings, checking, or other account(s) containing the borrowed funds, accompanied by an attestation from an officer of the bank or other financial institution that the funds are in the account(s) and are immediately available to the HHA. As with the HHA's own (that is, non-borrowed) funds, HCFA later may require the HHA to establish the current availability of such borrowed funds, including furnishing an attestation from a financial institution or other source, as may be appropriate, and to establish that such funds will remain available until a date when the HHA will have been surveyed by the State agency or by an approved accrediting organization.

(f) Line of credit. If the HHA chooses to support the availability of a portion of the initial reserve operating funds with a line of credit, it must provide HCFA with a letter of credit from the lender. HCFA later may require the HHA to furnish another attestation from the lender that the HHA, upon its certification into the Medicare program, continues to be approved to borrow the amount specified in the letter of credit.

(g) Provider agreement. HCFA does not enter into a provider agreement with an HHA unless the HHA meets the initial reserve operating funds requirement of this section.

[63 FR 312, Jan. 5, 1998]
Subpart C—Allowable Charges

§ 489.30 Allowable charges: Deductibles and coinsurance.
(a) Part A deductible and coinsurance. The provider may charge the beneficiary or other person on his or her behalf:
(1) The amount of the inpatient hospital deductible or, if less, the actual charges for the services;
(2) The amount of inpatient hospital coinsurance applicable for each day the individual is furnished inpatient hospital services after the 60th day, during a benefit period; and
(3) The posthospital SNF care coinsurance amount.
(4) In the case of durable medical equipment (DME) furnished as a home health service, 20 percent of the customary charge for the service.
(b) Part B deductible and coinsurance.
(1) The basic allowable charges are the $75 deductible and 20 percent of the customary (insofar as reasonable) charges in excess of that deductible.
(2) For hospital outpatient services, the allowable deductible charges depend on whether the hospital can determine the beneficiary’s deductible status.
(i) If the hospital is unable to determine the deductible status, it may charge the beneficiary its full customary charges up to $75.
(ii) If the beneficiary provides official information as to deductible status, the hospital may charge only the unmet portion of the deductible.
(3) In either of the cases discussed in paragraph (b)(2) of this section, the hospital is required to file with the intermediary, on a form prescribed by HCFA, information as to the services, charges, and amounts collected.
(4) The intermediary must reimburse the beneficiary if reimbursement is authorized and the expenses to the beneficiary’s deductible if the deductible has not yet been met.
(5) In the case of DME furnished as a home health service under Medicare Part B, the coinsurance is 20 percent of the customary (insofar as reasonable) charge for the services, with the following exception: If the DME is purchased by or on behalf of the beneficiary at a price at least 25 percent less than the reasonable charge for comparable new equipment, no coinsurance is required.

§ 489.31 Allowable charges: Blood.
(a) Limitations on charges. (1) A provider may charge the beneficiary (or other person on his or her behalf) only for the first three pints of blood or units of packed red cells furnished under Medicare Part A during a calendar year, or furnished under Medicare Part B during a calendar year.
(2) The charges may not exceed the provider’s customary charges.
(3) The provider may not charge for any whole blood or packed red cells in any of the circumstances specified in §409.87(c)(2) of this chapter.
(b) Offset for excessive charges. If the charge exceeds the cost to the provider, that excess will be deducted from any Medicare payments due the provider.

§ 489.32 Allowable charges: Non-covered and partially covered services.
(a) Services requested by beneficiary. If services furnished at the request of a beneficiary (or his or her representative) are more expensive than, or in excess of, services covered under Medicare—
(1) A provider may charge the beneficiary an amount that does not exceed the difference between—
(i) The provider’s customary charges for the services furnished; and
(ii) The provider’s customary charges for the kinds and amounts of services that are covered under Medicare.
(2) A provider may not charge for the services unless they have been requested by the beneficiary (or his or her representative) nor require a beneficiary to request services as a condition of admission.
(3) To avoid misunderstanding and disputes, a provider must inform any beneficiary who requests a service for which a charge will be made that there will be a specified charge for that service.
(b) Services not requested by the beneficiary. For special provisions that
apply when a provider customarily furnishes more expensive services, see §413.35 of this chapter.


§ 489.34 Allowable charges: Hospitals participating in State reimbursement control systems or demonstration projects.

A hospital receiving payment for a covered hospital stay under either a State reimbursement control system approved under §1886(c) of the Act or a demonstration project authorized under section 402(a) of Pub. L. 90–248 (42 U.S.C. 1395b–1) or section 222(a) of Pub. L. 92–603 (42 U.S.C. 1395b–1 (note)) and that would otherwise be subject to the prospective payment system set forth in part 412 of this chapter may charge a beneficiary for noncovered services as follows:

(a) For the custodial care and medically unnecessary services described in §412.42(c) of this chapter, after the conditions of §412.42(c)(1) through (c)(4) are met; and

(b) For all other services in accordance with the applicable rules of this subpart C.

[54 FR 41747, Oct. 11, 1989]

§ 489.35 Notice to intermediary.

The provider must inform its intermediary of any amounts collected from a beneficiary (or someone on his or her behalf) that are not authorized under subpart C of this part.

§ 489.40 Definition of incorrect collection.

(a) As used in this subpart, “incorrect collections” means any amounts collected from a beneficiary (or someone on his or her behalf) that are not authorized under subpart C of this part.

(b) A payment properly made to a provider by an individual not considered entitled to Medicare benefits will be deemed to be an “incorrect collection” when the individual is found to be retroactively entitled to benefits.

§ 489.41 Timing and methods of handling.

(a) Refund. Prompt refund to the beneficiary or other person is the preferred method of handling incorrect collections.

(b) Setting aside. If the provider cannot refund within 60 days from the date on the notice of incorrect collection, it must set aside an amount, equal to the amount incorrectly collected, in a separate account identified as to the individual to whom the payment is due. This amount incorrectly collected must be carried on the provider’s records in this manner until final disposition is made in accordance with the applicable State law.

(c) Notice to, and action by, intermediary. (1) The provider must notify the intermediary of the refund or setting aside required under paragraphs (a) and (b) of this section.

(2) If the provider fails to refund or set aside the required amounts, they may be offset against amounts otherwise due the provider.

§ 489.42 Payment of offset amounts to beneficiary or other person.

(a) In order to carry out the commitment to refund amounts incorrectly collected, HCFA may determine that amounts offset in accordance with §489.41 are to be paid directly to the beneficiary or other person from whom the provider received the incorrect collection, if:

(1) HCFA finds that the provider has failed, following written request, to refund the amount of the incorrect collection to the beneficiary or other person; and

(2) The provider agreement has been terminated in accordance with the provisions of subpart E of this part.

(b) Before making a determination to make payment under paragraph (a) of this section, HCFA will give written notice to the provider (1) explaining that an incorrect collection was made and the amount; (2) requesting the provider to refund the incorrect collection to the beneficiary or other person; and (3) advising of HCFA’s intention to make a determination under paragraph (a) of this section.
§ 489.53 Termination by HCFA.

(a) Basis for termination of agreement with any provider. HCFA may terminate the agreement with any provider if HCFA finds that any of the following failings is attributable to that provider:

(1) It is not complying with the provisions of title XVIII and the applicable regulations of this chapter or with the provisions of the agreement.

(2) It places restrictions on the persons it will accept for treatment and it fails either to exempt Medicare beneficiaries from those restrictions or to apply them to Medicare beneficiaries the same as to all other persons seeking care.

(3) It no longer meets the appropriate conditions of participation or requirements (for SNFs and NFs) set forth elsewhere in this chapter.

(4) It fails to furnish information that HCFA finds necessary for a determination as to whether payments are or were due under Medicare and the amounts due.

(5) It refuses to permit examination of its fiscal or other records by, or on behalf of HCFA, as necessary for verification of information furnished as a basis for payment under Medicare.

(6) It failed to furnish information on business transactions as required in § 420.205 of this chapter.

(7) It failed at the time the agreement was entered into or renewed to disclose information on convicted individuals as required in § 420.204 of this chapter.

(8) It failed to furnish ownership information as required in § 420.206 of this chapter.

(9) It failed to comply with civil rights requirements set forth in 45 CFR parts 80, 84, and 90.

(10) In the case of a hospital or a critical access hospital as defined in section 1861(mm)(1) of the Act that has reason to believe it may have received an individual transferred by another hospital in violation of § 489.24(d), the hospital failed to report the incident to HCFA or the State survey agency.

(11) In the case of a hospital requested to furnish inpatient services to
§ 489.53 CHAMPUS or CHAMPVA beneficiaries or to veterans, it failed to comply with § 489.25 or § 489.26, respectively.

(12) It failed to furnish the notice of discharge rights as required by § 489.27.

(13) It refuses to permit photocopying of any records or other information by, or on behalf of HCFA, as necessary to determine or verify compliance with participation requirements.

(14) The hospital knowingly and willfully fails to accept, on a repeated basis, an amount that approximates the Medicare rate established under the inpatient hospital prospective payment system, minus any enrollee deductibles or copayments, as payment in full from a fee-for-service FEHB plan for inpatient hospital services provided to a retired Federal enrollee of a fee-for-service FEHB plan, age 65 or older, who does not have Medicare Part A benefits.

(a) Termination of agreements with certain hospitals.

In the case of a hospital with an emergency department, as defined in § 489.24(b), HCFA may terminate the provider agreement if—

(1) The hospital fails to comply with the requirements of § 489.24(a) through (e), which require the hospital to examine, treat, or transfer emergency medical condition cases appropriately, and require that hospitals with specialized capabilities or facilities accept an appropriate transfer; or

(2) The hospital fails to comply with § 489.20(m), (q), and (r), which require the hospital to report suspected violations of § 489.24(d), to post conspicuously in emergency departments or in a place or places likely to be noticed by all individuals entering the emergency departments, as well as those individuals waiting for examination and treatment in areas other than traditional emergency departments, (that is, entrance, admitting area, waiting room, treatment area), signs specifying rights of individuals under this subpart, to post conspicuously information indicating whether or not the hospital participates in the Medicaid program, and to maintain medical and other records related to transferred individuals for a period of 5 years, a list of on-call physicians for individuals with emergency medical conditions, and a central log on each individual who comes to the emergency department seeking assistance.

(c) Notice of termination—(1) Timing: Basic rule. Except as provided in paragraph (c)(2) of this section, HCFA gives the provider notice of termination at least 15 days before the effective date of termination of the provider agreement.

(2) Timing exceptions: Immediate jeopardy situations—(i) Hospital with emergency department. If HCFA finds that a hospital with an emergency department is in violation of § 489.24, paragraphs (a) through (e), and HCFA determines that the violation poses immediate jeopardy to the health or safety of individuals who present themselves to the hospital for emergency services, HCFA—

(A) Gives the hospital a preliminary notice indicating that its provider agreement will be terminated in 23 days if it does not correct the identified deficiencies or refute the finding; and

(B) Gives a final notice of termination, and concurrent notice to the public, at least 2, but not more than 4, days before the effective date of termination of the provider agreement.

(ii) Skilled nursing facilities (SNFs). For an SNF with deficiencies that pose immediate jeopardy to the health or safety of residents, HCFA gives notice at least 2 days before the effective date of termination of the provider agreement.

(3) Content of notice. The notice states the reasons for, and the effective date of, the termination, and explains the extent to which services may continue after that date, in accordance with § 489.55.

(4) Notice to public. HCFA concurrently gives notice of the termination to the public.

(d) Appeal by the provider. A provider may appeal the termination of its provider agreement by HCFA in accordance with part 498 of this chapter.

§ 489.53 CHAMPUS or CHAMPVA beneficiaries or to veterans, it failed to comply with § 489.25 or § 489.26, respectively.
§ 489.60 Definitions.

(a) Inpatient hospital services (including inpatient psychiatric hospital services) and posthospital extended care services furnished to a beneficiary who was admitted before the effective date of termination; and

(b) Home health services and hospice care furnished under a plan established before the effective date of termination.¹

[50 FR 37376, Sept. 13, 1985]

§ 489.57 Reinstatement after termination.

When a provider agreement has been terminated by HCFA under §489.53, or by the OIG under §489.54, a new agreement with that provider will not be accepted unless HCFA or the OIG, as appropriate, finds—

(a) That the reason for termination of the previous agreement has been removed and there is reasonable assurance that it will not recur; and

(b) That the provider has fulfilled, or has made satisfactory arrangements to fulfill, all of the statutory and regulatory responsibilities of its previous agreement.

[51 FR 24493, July 3, 1986]

Subpart F—Surety Bond
Requirements for HHAs

SOURCE: 63 FR 313, Jan. 5, 1998, unless otherwise noted.

§ 489.60 Definitions.

As used in this subpart unless the context indicates otherwise—

Assessment means a sum certain that HCFA may assess against an HHA in lieu of damages under Titles XI, XVIII, or XXI of the Social Security Act or under regulations in this chapter.

Assets includes but is not limited to any listing that identifies Medicare beneficiaries to whom home health services were furnished by a participating or formerly participating HHA.

Civil money penalty means a sum certain that HCFA has the authority to impose on an HHA as a penalty under Titles XI, XVIII, or XXI of the Social Security Act.

¹For termination before July 18, 1984, payment was available through the calendar year in which the termination was effective.
§ 489.61 Basic requirement for surety bonds.

Except as provided in §489.62, each HHA that is a Medicare participating HHA, or that seeks to become a Medicare participating HHA, must obtain a surety bond (and furnish to HCFA a copy of such surety bond) that meets the requirements of this subpart F and HCFA's instructions.

§ 489.62 Requirement waived for Government-operated HHAs.

An HHA operated by a Federal, State, local, or tribal government agency is deemed to have provided HCFA with a comparable surety bond under State law, and HCFA therefore waives the requirements of this subpart with respect to such an HHA if, during the preceding 5 years the HHA has—

(a) Not had any unpaid claims or unpaid civil money penalties or assessments; and

(b) Not had any of its claims referred by HCFA to the Department of Justice or the General Accounting Office in accordance with part 401 of this chapter.

§ 489.63 Parties to the bond.

The surety bond must name the HHA as Principal, HCFA as Obligee, and the surety company (and its heirs, executors, administrators, successors and assigns, jointly and severally) as Surety.

§ 489.64 Authorized Surety and exclusion of surety companies.

(a) An HHA may obtain a surety bond required under §489.61 only from an authorized Surety.

(b) An authorized Surety is a surety company that—

1. Has been issued a Certificate of Authority by the U.S. Department of the Treasury in accordance with 31 U.S.C. 9304 to 9308 and 31 CFR parts 223, 224, and 225 as an acceptable surety on Federal bonds and the Certificate has neither expired nor been revoked; and

2. Has not been determined by HCFA to be an unauthorized Surety for the purpose of an HHA obtaining a surety bond under this section.
(c) HCFA determines that a surety company is an unauthorized Surety under this section—
(1) If, upon request by HCFA, the surety company fails to furnish timely confirmation of the issuance of, and the validity and accuracy of information appearing on, a surety bond an HHA presents to HCFA that shows the surety company as Surety on the bond;
(2) If, upon presentation by HCFA to the surety company of a request for payment on a surety bond and of sufficient evidence to establish the surety company’s liability on the bond, the surety company fails to timely pay HCFA in full the amount requested, up to the face amount of the bond; or
(3) For other good cause.
(d) Any determination HCFA makes under paragraph (c) of this section is effective immediately when notice of the determination is published in the FEDERAL REGISTER and remains in effect until a notice of reinstatement is published in the FEDERAL REGISTER.
(e) Any determination HCFA makes under paragraph (c) of this section does not affect the Surety’s liability under any surety bond issued by a surety company to an HHA before notice of such determination is published in accordance with paragraph (d) of this section.
(f) A determination by HCFA that a surety company is an unauthorized Surety under this section is not a barment, suspension, or exclusion for the purposes of Executive Order No. 12549 (3 CFR, 1986 comp., p. 189).
§ 489.65 Amount of the bond.
(a) Basic rule. The amount of the surety bond must be $50,000 or 15 percent of the Medicare payments made by HCFA to the HHA in the HHA’s most recent fiscal year for which a cost report has been accepted by HCFA, whichever is greater.
(b) Computation of the 15 percent: Participating HHA.
The 15 percent is computed as follows:
(1) For the initial bond—on the basis of Medicare payments made by HCFA to the HHA in the HHA’s most recent fiscal year as shown in the HHA’s most recent cost report that has been accepted by HCFA. If the initial bond will cover less than a full fiscal year, the computation of the 15 percent will be based on the number of months of the fiscal year that the bond will cover.
(2) For subsequent bonds—on the basis of Medicare payments made by HCFA in the most recent fiscal year for which a cost report has been accepted. However, if payments in the first six months of the current fiscal year differ from such an amount by more than 25 percent, then the amount of the bond is 15 percent of such payments projected on an annualized basis.
(c) Computation of 15 percent: An HHA that seeks to become a participating HHA by obtaining assets or ownership interest.
For an HHA that seeks to become a participating HHA by purchasing the assets or the ownership interest of a participating or formerly participating HHA, the 15 percent is computed on the basis of Medicare payments made by HCFA to the participating or formerly participating HHA in the most recent fiscal year that a cost report has been accepted.
(d) Change of ownership.
For an HHA that undergoes a change of ownership the 15 percent is computed on the basis of Medicare payments made by HCFA to the HHA for the most recently accepted cost report.
(e) An HHA that seeks to become a participating HHA without obtaining assets or ownership interest.
For an HHA that seeks to become a participating HHA without purchasing the assets or the ownership interest of a participating or formerly participating HHA, the 15 percent computation does not apply.
(f) Exception to the basic rule.
If an HHA’s overpayment in the most recently accepted cost report exceeds 15 percent of annual payments, HCFA may require the HHA to secure a bond in an amount up to or equal to the amount of overpayment, provided the amount of the bond is not less than $50,000.
(g) Expiration of the 15 percent provision.
For an annual surety bond, or for a rider on a continuous surety bond, that is required to be submitted on or after June 1, 2005, notwithstanding any reference in this subpart to 15 percent as a basis for determining the amount of the bond, the amount of the bond or rider, as applicable, must be $50,000 or
§ 489.66 Additional requirements of the surety bond.

The surety bond that an HHA obtains under this subpart must meet the following additional requirements:

(a) The bond must guarantee that within 30 days of receiving written notice from HCFA of an unpaid claim or unpaid civil money penalty or assessment, which notice contains sufficient evidence to establish the Surety’s liability under the bond, the Surety will pay HCFA, up to the stated amount of the bond—

(1) The full amount of any unpaid claim, plus accrued interest, for which the HHA is responsible; and

(2) The full amount of any unpaid civil money penalty or assessment imposed by HCFA on the HHA, plus accrued interest.

(b) The bond must provide the following:

(1) The Surety is liable for unpaid claims, unpaid civil money penalties, and unpaid assessments that are discovered when the surety bond is in effect, regardless of when the payment, overpayment, or other event giving rise to the claim, civil money penalty, or assessment occurred, provided HCFA makes a written demand for payment from the Surety during, or within 90 days after, the term of the bond.

(2) If the HHA fails to furnish a bond meeting the requirements of this subpart F for the year following expiration of the term of an annual bond, or if the HHA fails to submit a rider when a rider is required to be submitted under this subpart, or if the HHA’s provider agreement is terminated, the last bond or rider, as applicable, submitted by the HHA to HCFA, which bond or applicable rider meets the requirements of this subpart, remains effective and the Surety remains liable for unpaid claims, civil money penalties, and assessments that—

(i) HCFA determines or imposes on or asserts against the HHA based on overpayments or other events that took place during or prior to the term of the last bond or rider; and

(ii) Were determined or imposed during the 2 years following the date the HHA failed to submit a bond or required rider or the date the HHA’s provider agreement is terminated, whichever is later.

(c) The bond must provide that the Surety’s liability to HCFA under the bond is not extinguished by any action of the HHA, the Surety, or HCFA, including but not necessarily limited to any of the following actions:

(1) Action by the HHA or the Surety to terminate or limit the scope or term of the bond. The Surety’s liability may be extinguished, however, when—

(i) The Surety furnishes HCFA with notice of such action not later than 10 days after receiving notice from the HHA of action by the HHA to terminate or limit the scope of the bond, or not later than 60 days before the effective date of such action by the Surety; or

(ii) The HHA furnishes HCFA with a new bond that meets the requirements of this subpart.

(2) The Surety’s failure to continue to meet the requirements of § 489.64(a) or HCFA’s determination that the surety company is an unauthorized Surety under § 489.64(b).

(3) Termination of the HHA’s provider agreement.

(4) Any action by HCFA to suspend, offset, or otherwise recover payments to the HHA.

(5) Any action by the HHA to—

(i) Cease operation;

(ii) Sell or transfer any asset or ownership interest;

(iii) File for bankruptcy; or

(iv) Fail to pay the Surety.

(6) Any fraud, misrepresentation, or negligence by the HHA in obtaining the surety bond or by the Surety (or by the Surety’s agent, if any) in issuing the surety bond, except that any fraud, misrepresentation, or negligence by the HHA in identifying to the Surety (or to the Surety’s agent) the amount of Medicare payments upon which the amount of the surety bond is determined will not cause the Surety’s liability to HCFA to exceed the amount of the bond.
§ 489.68 Effect of failure to obtain, maintain, and timely file a surety bond.

(a) The failure of a participating HHA to obtain, file timely, and maintain a surety bond in accordance with this subpart F and HCFA’s instructions is sufficient under §489.53(a)(1) for HCFA to terminate the HHA’s provider agreement.

(b) The failure of an HHA seeking to become a participating HHA to obtain and file timely a surety bond in accordance with this Subpart F and HCFA’s instructions is sufficient under §489.12(a)(3) for HCFA to refuse to enter into a provider agreement with the HHA.
§ 489.69 Evidence of compliance.
(a) HCFA may at any time require an HHA to make a specific showing of being in compliance with the requirements of this Subpart F and may require the HHA to submit such additional evidence as HCFA considers sufficient to demonstrate the HHA’s compliance.
(b) If requested by HCFA to do so, the failure of an HHA to timely furnish sufficient evidence to HCFA to demonstrate compliance with the requirements of this Subpart F is sufficient for HCFA to terminate the HHA’s provider agreement under § 489.53(a)(1) or to refuse to enter into a provider agreement with the HHA under § 489.12(a)(3), as applicable.

§ 489.70 Effect of payment by the Surety.
A Surety’s payment to HCFA under a bond for an unpaid claim or an unpaid civil money penalty or assessment, constitutes—
(a) Collection of the unpaid claim or unpaid civil money penalty or assessment (to the extent the Surety’s payment on the bond covers such unpaid claim, civil money penalty, or assessment); and
(b) A basis for termination of the HHA’s provider agreement under § 489.53(a)(1).

§ 489.71 Surety’s standing to appeal Medicare determinations.
A Surety has standing to appeal any matter that the HHA could appeal, provided the Surety satisfies all jurisdictional and procedural requirements that would otherwise have applied to the HHA, and provided the HHA is not, itself, actively pursuing its appeal rights under this chapter, and provided further that, with respect to unpaid claims, the Surety has paid HCFA all amounts owed to HCFA by the HHA on such unpaid claims, up to the amount of the bond.

§ 489.72 Effect of review reversing determination.
In the event a Surety has paid HCFA on the basis of liability incurred under a bond obtained by an HHA under this subpart F, and to the extent the HHA that obtained such bond (or the Surety under § 489.71) is subsequently successful in appealing the determination that was the basis of the unpaid claim or unpaid civil money penalty or assessment that caused the Surety to pay HCFA under the bond, HCFA will refund to the Surety the amount the Surety paid to HCFA to the extent such amount relates to the matter that was successfully appealed by the HHA (or by the Surety), provided all review, including judicial review, has been completed on such matter. Any additional amounts owing as a result of the appeal will be paid to the HHA.

§ 489.73 Effect of conditions of payment.
If a Surety has paid an amount to HCFA on the basis of liability incurred under a bond obtained by an HHA under this subpart F, and HCFA subsequently collects from the HHA, in whole or in part, on such unpaid claim, civil money penalty, or assessment that was the basis for the Surety’s liability, HCFA reimburses the Surety such amount as HCFA collected from the HHA, up to the amount paid by the Surety to HCFA, provided the Surety has no other liability to HCFA under the bond.

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

[63 FR 29656, June 1, 1998]

§ 489.74 Incorporation into existing provider agreements.
The requirements of this subpart F are deemed to be incorporated into existing HHA provider agreements effective January 1, 1998.

[63 FR 315, Jan. 5, 1998. Redesignated at 63 FR 29656, June 1, 1998]

Subparts G–H [Reserved]

Subpart I—Advance Directives

Source: 57 FR 8203, Mar. 6, 1992, unless otherwise noted.

§ 489.100 Definition.
For purposes of this part, advance directive means a written instruction,
such as a living will or durable power of attorney for health care, recognized under State law (whether statutory or as recognized by the courts of the State), relating to the provision of health care when the individual is incapacitated.

§ 489.102 Requirements for providers.

(a) Hospitals, critical access hospitals, skilled nursing facilities, nursing facilities, home health agencies, providers of home health care (and for Medicaid purposes, providers of personal care services), and hospices must maintain written policies and procedures concerning advance directives with respect to all adult individuals receiving medical care by or through the provider and are required to:

1. Provide written information to such individuals concerning—
   (i) An individual’s rights under State law (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. Providers are to update and disseminate amended information as soon as possible, but no later than 90 days from the effective date of the changes to State law; and
   (ii) The written policies of the provider or organization respecting the implementation of such rights, including a clear and precise statement of limitation if the provider cannot implement an advance directive on the basis of conscience. At a minimum, a provider’s statement of limitation should:
      (A) Clarify any differences between institution-wide conscience objections and those that may be raised by individual physicians;
      (B) Identify the state legal authority permitting such objection; and
      (C) Describe the range of medical conditions or procedures affected by the conscience objection.

2. Document in a prominent part of the individual’s current medical record, or patient care record in the case of an individual in a religious nonmedical health care institution, whether or not the individual has executed an advance directive;

3. Not condition the provision of care or otherwise discriminate against an individual based on whether or not the individual has executed an advance directive;

4. Ensure compliance with requirements of State law (whether statutory or recognized by the courts of the State) regarding advance directives. The provider must inform individuals that complaints concerning the advance directive requirements may be filed with the State survey and certification agency;

5. Provide for education of staff concerning its policies and procedures on advance directives; and

6. Provide for community education regarding issues concerning advance directives that may include material required in paragraph (a)(1) of this section, either directly or in concert with other providers and organizations. Separate community education materials may be developed and used, at the discretion of providers. The same written materials do not have to be provided in 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. A provider must be able to document its community education efforts.

(b) The information specified in paragraph (a) of this section is furnished:

1. In the case of a hospital, at the time of the individual’s admission as an inpatient.

2. In the case of a skilled nursing facility at the time of the individual’s admission as a resident.

3. In the case of a home health agency, in advance of the individual coming under the care of the agency. The HHA may furnish advance directives information to a patient at the time of the first home visit, as long as the information is furnished before care is provided.
(ii) In the case of personal care services, in advance of the individual coming under the care of the personal care services provider. The personal care provider may furnish advance directives information to a patient at the time of the first home visit, as long as the information is furnished before care is provided.

(4) In the case of a hospice program, at the time of initial receipt of hospice care by the individual from the program.

(c) The providers listed in paragraph (a) of this section—

(1) Are not required to provide care that conflicts with an advance directive.

(2) Are not required to implement an advance directive if, as a matter of conscience, the provider cannot implement an advance directive and State law allows any health care provider or any agent of such provider to conscientiously object.

(d) Prepaid or eligible organizations (as specified in sections 1833(a)(1)(A) and 1876(b) of the Act) must meet the requirements specified in §417.436 of this chapter.

(e) If an adult individual is incapacitated at the time of admission or at the start of care and is unable to receive information (due to the incapacitating conditions or a mental disorder) or articulate whether or not he or she has executed an advance directive, then the provider may give advance directive information to the individual’s family or surrogate in the same manner that it issues other materials about policies and procedures to the family of the incapacitated individual or to a surrogate or other concerned persons in accordance with State law. The provider is not relieved of its obligation to provide this information to the individual once he or she is no longer incapacitated or unable to receive such information. Follow-up procedures must be in place to provide the information to the individual directly at the appropriate time.

§ 489.104 Effective dates.

These provisions apply to services furnished on or after December 1, 1991, payments made under section 1833(a)(1)(A) of the Act on or after December 1, 1991, and contracts effective on or after December 1, 1991.

PART 491—CERTIFICATION OF CERTAIN HEALTH FACILITIES

Subpart A—Rural Health Clinics: Conditions for Certification; and FQHCs Conditions for Coverage

Sec.
491.1 Purpose and scope.
491.2 Definitions.
491.3 Certification procedures.
491.4 Compliance with Federal, State and local laws.
491.5 Location of clinic.
491.6 Physical plant and environment.
491.7 Organizational structure.
491.8 Staffing and staff responsibilities.
491.9 Provision of services.
491.10 Patient health records.
491.11 Program evaluation.

AUTHORITY: Sec. 1102 of the Social Security Act (42 U.S.C. 1302); and sec. 353 of the Public Health Service Act (42 U.S.C. 263a).

EDITORIAL NOTE: Nomenclature changes to part 491 appear at 61 FR 14658, Apr. 3, 1996.

Subpart A—Rural Health Clinics: Conditions for Certification; and FQHCs Conditions for Coverage

§ 491.1 Purpose and scope.

This subpart sets forth the conditions that rural health clinics or FQHCs must meet in order to qualify for reimbursement under Medicare (title XVIII of the Social Security Act) and that rural health clinics must meet in order to qualify for reimbursement under Medicaid (title XIX of the Act).

[57 FR 24082, June 12, 1992]

§ 491.2 Definitions.

As used in this subpart, unless the context indicates otherwise:

Direct services means services provided by the clinic’s staff.

FQHC means an entity as defined in §405.24(b).
Nurse practitioner means a registered professional nurse who is currently licensed to practice in the State, who meets the State’s requirements governing the qualifications of nurse practitioners, and who meets one of the following conditions:

1. Is currently certified as a primary care nurse practitioner by the American Nurses’ Association or by the National Board of Pediatric Nurse Practitioners and Associates; or

2. Has satisfactorily completed a formal 1 academic year educational program that:
   (i) Prepares registered nurses to perform an expanded role in the delivery of primary care;
   (ii) Includes at least 4 months (in the aggregate) of classroom instruction and a component of supervised clinical practice; and
   (iii) Awards a degree, diploma, or certificate to persons who successfully complete the program; or

3. Has successfully completed a formal educational program (for preparing registered nurses to perform an expanded role in the delivery of primary care) that does not meet the requirements of paragraph (b)(2) of this section and has been performing an expanded role in the delivery of primary care for a total of 12 months during the 18-month period immediately preceding the effective date of this subpart.

Physician assistant means a person who meets the applicable State requirements governing the qualifications for assistants to primary care physicians, and who meets at least one of the following conditions:

1. Is currently certified by the National Commission on Certification of Physician Assistants to assist primary care physicians; or

2. Has satisfactorily completed a program for preparing physician’s assistants that:
   (i) Was at least 1 academic year in length;
   (ii) Consisted of supervised clinical practice and at least 4 months (in the aggregate) of classroom instruction directed toward preparing students to deliver health care; and
   (iii) Was accredited by the American Medical Association’s Committee on Allied Health Education and Accreditation; or

3. Has satisfactorily completed a formal educational program (for preparing physician assistants) that does not meet the requirements of paragraph (d)(2) of this section and assisted primary care physicians for a total of 12 months during the 18-month period that ended on December 31, 1986.

Rural area means an area that is not delineated as an urbanized area by the Bureau of the Census.

Rural health clinic or clinic means a clinic that is located in a rural area designated as a shortage area, is not a rehabilitation agency or a facility primarily for the care and treatment of mental diseases, and meets all other requirements of this subpart.

Shortage area means a defined geographic area designated by the Department as having either a shortage of personal health services (under section 1302(7) of the Public Health Service Act) or a shortage of primary medical care manpower (under section 332 of that Act).

Secretary means the Secretary of Health and Human Services, or any official to whom he has delegated the pertinent authority.

§491.3 Certification procedures.

A rural health clinic will be certified for participation in Medicare in accordance with subpart S of 42 CFR part 405. The Secretary will notify the State Medicaid agency whenever he has certified or denied certification under Medicare for a prospective rural health clinic in that State. A clinic certified under Medicare will be deemed to meet the standards for certification under Medicaid.
§ 491.4 Compliance with Federal, State and local laws.

The rural health clinic or FQHC and its staff are in compliance with applicable Federal, State and local laws and regulations.

(a) Licensure of clinic or center. The clinic or center is licensed pursuant to applicable State and local law.

(b) Licensure, certification or registration of personnel. Staff of the clinic or center are licensed, certified or registered in accordance with applicable State and local laws.

§ 491.5 Location of clinic.

(a) Basic requirements. (1) An RHC is located in a rural area that is designated as a shortage area.

(2) An FQHC is located in a rural or urban area that is designated as either a shortage area or an area that has a medically underserved population.

(3) Both the RHC and the FQHC may be permanent or mobile units.

(i) Permanent unit. The objects, equipment, and supplies necessary for the provision of the services furnished directly by the clinic or center are housed in a permanent structure.

(ii) Mobile unit. The objects, equipment, and supplies necessary for the provision of the services furnished directly by the clinic or center are housed in a mobile structure, which has fixed, scheduled location(s).

(iii) Permanent unit in more than one location. If clinic or center services are furnished at permanent units in more than one location, each unit is independently considered for approval as a rural health clinic or for approval as an FQHC.

(b) Exceptions. (1) HCFA does not disqualify an RHC approved under this subpart if the area in which it is located subsequently fails to meet the definition of a rural, shortage area.

(2) A private, nonprofit facility that meets all other conditions of this subpart except for location in a shortage area will be certified if, on July 1, 1977, it was operating in a rural area that is determined by the Secretary (on the basis of the ratio of primary care physicians to the general population) to have an insufficient supply of physicians to meet the needs of the area served.

(3) Determinations on these exceptions will be made by the Secretary upon application by the facility.

(c) Criteria for designation of rural areas. (1) Rural areas are areas not delineated as urbanized areas in the last census conducted by the Census Bureau.

(2) Excluded from the rural area classification are:

(i) Central cities of 50,000 inhabitants or more;

(ii) Cities with at least 25,000 inhabitants which, together with contiguous areas having stipulated population density, have combined populations of 50,000 and constitute, for general economic and social purposes, single communities;

(iii) Closely settled territories surrounding cities and specifically designated by the Census Bureau as urban.

(3) Included in the rural area classification are those portions of extended cities that the Census Bureau has determined to be rural.

(d) Criteria for designation of shortage areas. (1) The criteria for determination of shortage of personal health services (under section 1302(7) of the Public Health Services Act), are:

(i) The ratio of primary care physicians practicing within the area to the resident population;

(ii) The infant mortality rate;

(iii) The percent of the population 65 years of age or older; and

(iv) The percent of the population with a family income below the poverty level.

(2) The criteria for determination of shortage of primary medical care manpower (under section 332(a)(3)(A) of the Public Health Services Act) are:

(i) The area served is a rational area for the delivery of primary medical care services;

(ii) The ratio of primary care physicians practicing within the area to the resident population; and

(iii) The primary medical care manpower in contiguous areas is overutilized, excessively distant, or inaccessible to the population in this area.

(e) Medically underserved population. A medically underserved population includes the following:
§ 491.8 Staffing and staff responsibilities.

(a) Staffing. (1) The clinic or center has a health care staff that includes one or more physicians. Rural health clinic staffs must also include one or more physician's assistants or nurse practitioners.

(2) The physician member of the staff may be the owner of the rural health clinic, an employee of the clinic or center, or under agreement with the clinic or center to carry out the responsibilities required under this section.

(3) The physician assistant, nurse practitioner, nurse-midwife, clinical social worker, or clinical psychologist member of the staff may be the owner or an employee of the clinic or center, or may furnish services under contract to the center.

(4) The staff may also include ancillary personnel who are supervised by the professional staff.

(5) The staff is sufficient to provide the services essential to the operation of the clinic or center.

(b) Physician responsibilities. (1) In a physician, nurse practitioner, physician assistant, nurse-midwife, clinical social worker, or clinical psychologist is available to furnish patient care services at all times the clinic or center operates. In addition, for rural health clinics, a nurse practitioner or a physician assistant is available to furnish patient care services at least 60 percent of the time the clinic operates.

(2) The physician:

(i) Except for services furnished by a clinical psychologist in an FQHC, which State law permits to be provided...
§ 491.9 Provision of services.

(a) Basic requirements. (1) All services offered by the clinic or center are furnished in accordance with applicable Federal, State, and local laws; and

(2) The clinic or center is primarily engaged in providing outpatient health services and meets all other conditions of this subpart.

(b) Patient care policies. (1) The clinic's or center's health care services are furnished in accordance with appropriate written policies which are consistent with applicable State law.

(2) The policies are developed with the advice of a group of professional personnel that includes one or more physicians and one or more physician assistants or nurse practitioners. At least one member is not a member of the clinic or center staff.

(3) The policies include:

(i) A description of the services the clinic or center furnishes directly and those furnished through agreement or arrangement.

(ii) Guidelines for the medical management of health problems which include the conditions requiring medical consultation and/or patient referral, the maintenance of health care records, and procedures for the periodic review and evaluation of the services furnished by the clinic or center.

(iii) Rules for the storage, handling, and administration of drugs and biologicals.

(4) These policies are reviewed at least annually by the group of professional personnel required under paragraph (b)(2) of this section and reviewed as necessary by the clinic or center.

(c) Direct services—(1) General. The clinic or center staff furnishes those diagnostic and therapeutic services and supplies that are commonly furnished in a physician's office or at the entry point into the health care delivery system. These include medical history, physical examination, assessment of health status, and treatment for a variety of medical conditions.

(2) Laboratory. These requirements apply to RHCs but not to FQHCs. The

[57 FR 24083, June 12, 1992, as amended at 61 FR 14658, Apr. 3, 1996]
RHC provides laboratory services in accordance with part 493 of this chapter, which implements the provisions of section 353 of the Public Health Service Act. The RHC provides basic laboratory services essential to the immediate diagnosis and treatment of the patient, including:

(i) Chemical examinations of urine by stick or tablet method or both (including urine ketones);
(ii) Hemoglobin or hematocrit;
(iii) Blood glucose;
(iv) Examination of stool specimens for occult blood;
(v) Pregnancy tests; and
(vi) Primary culturing for transmittal to a certified laboratory.

(3) Emergency. The clinic or center provides medical emergency procedures as a first response to common life-threatening injuries and acute illness and has available the drugs and biologicals commonly used in life saving procedures, such as analgesics, anesthetics (local), antibiotics, anticonvulsants, antidotes and emetics, serums and toxoids.

(d) Services provided through agreements or arrangements. (1) The clinic or center has agreements or arrangements with one or more providers or suppliers participating under Medicare or Medicaid to furnish other services to its patients, including:

(i) Inpatient hospital care;
(ii) Physician(s) services (whether furnished in the hospital, the office, the patient’s home, a skilled nursing facility, or elsewhere); and
(iii) Additional and specialized diagnostic and laboratory services that are not available at the clinic or center.

(2) If the agreements are not in writing, there is evidence that patients referred by the clinic or center are being accepted and treated.


§ 491.10 Patient health records.

(a) Records system. (1) The clinic or center maintains a clinical record system in accordance with written policies and procedures.

(2) A designated member of the professional staff is responsible for maintaining the records and for insuring that they are completely and accurately documented, readily accessible, and systematically organized.

(3) For each patient receiving health care services, the clinic or center maintains a record that includes, as applicable:

(i) Identification and social data, evidence of consent forms, pertinent medical history, assessment of the health status and health care needs of the patient, and a brief summary of the episode, disposition, and instructions to the patient;
(ii) Reports of physical examinations, diagnostic and laboratory test results, and consultative findings;
(iii) All physician’s orders, reports of treatments and medications, and other pertinent information necessary to monitor the patient’s progress;
(iv) Signatures of the physician or other health care professional.

(b) Protection of record information. (1) The clinic or center maintains the confidentiality of record information and provides safeguards against loss, destruction or unauthorized use.

(2) Written policies and procedures govern the use and removal of records from the clinic or center and the conditions for release of information.

(3) The patient’s written consent is required for release of information not authorized to be released without such consent.

(c) Retention of records. The records are retained for at least 6 years from date of last entry, and longer if required by State statute.

(57 FR 24984, June 12, 1992)

§ 491.11 Program evaluation.

(a) The clinic or center carries out, or arranges for, an annual evaluation of its total program.

(b) The evaluation includes review of:

(1) The utilization of clinic or center services, including at least the number of patients served and the volume of services;
(2) A representative sample of both active and closed clinical records; and
The purpose of the evaluation is to determine whether:

(1) The utilization of services was appropriate;
(2) The established policies were followed; and
(3) Any changes are needed.

(d) The clinic or center staff considers the findings of the evaluation and takes corrective action if necessary.

[57 FR 24984, June 12, 1992]
Health Care Financing Administration, HHS

493.640 Methodology for determining fee amount.

Subpart G [Reserved]

Subpart H—Participation in Proficiency Testing for Laboratories Performing Tests of Moderate Complexity (Including the Subcategory), High Complexity, or Any Combination of These Tests

493.801 Condition: Enrollment and testing of samples.
493.803 Condition: Successful participation.
493.807 Condition: Reinstatement of laboratories performing tests of moderate complexity (including the subcategory), high complexity, or any combination of these tests, after failure to participate successfully.

Proficiency Testing by Specialty and Subspecialty for Laboratories Performing Tests of Moderate Complexity (Including the Subcategory), High Complexity, or Any Combination of These Tests

493.821 Condition: Microbiology.
493.823 Standard; Bacteriology.
493.825 Standard; Mycobacteriology.
493.827 Standard; Mycology.
493.829 Standard; Parasitology.
493.831 Standard; Virology.
493.833 Condition: Diagnostic immunology.
493.835 Standard; Syphilis serology.
493.837 Standard; General immunology.
493.839 Condition: Chemistry.
493.841 Standard; Routine chemistry.
493.843 Standard; Endocrinology.
493.845 Standard; Toxicology.
493.847 Condition: Hematology.
493.851 Standard; Hematology.
493.853 Condition: Pathology.
493.855 Standard; Cytology; gynecologic examinations.
493.857 Condition: Immunohematology.
493.859 Standard; ABO group and D (Rh) typing.
493.861 Standard; Unexpected antibody detection.
493.863 Standard; Compatibility testing.
493.865 Standard; Antibody identification.

Subpart I—Proficiency Testing Programs for Tests of Moderate Complexity (Including the Subcategory), High Complexity, or Any Combination of These Tests

493.901 Approval of proficiency testing programs.
493.903 Administrative responsibilities.
493.905 Nonapproved proficiency testing programs.

Proficiency Testing Programs by Specialty and Subspecialty

493.909 Microbiology.
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493.1225 Condition: Microbiology.
493.1227 Condition: Bacteriology.
493.1229 Condition: Mycobacteriology.
493.1231 Condition: Mycology.
493.1233 Condition: Parasitology.
493.1235 Condition: Virology.
493.1237 Condition: Diagnostic immunology.
493.1239 Condition: Syphilis serology.
493.1241 Condition: General immunology.
493.1243 Condition: Chemistry.
493.1245 Condition: Routine chemistry.
493.1247 Condition: Endocrinology.
493.1249 Condition: Toxicology.
493.1251 Condition: Urinalysis.
493.1253 Condition: Hematology.
493.1255 Condition: Pathology.
493.1257 Condition: Cytology.
493.1259 Condition: Histopathology.
493.1261 Condition: Oral pathology.
493.1263 Condition: Radiobioassay.
493.1265 Condition: Histocompatibility.
493.1267 Condition: Clinical cytogenetics.
493.1269 Condition: Immunohematology.
493.1271 Condition: Transfusion services and bloodbanking.
493.1273 Standard; Immunohematological collection, processing, dating periods, labeling and distribution of blood and blood products.
493.1275 Standard; Blood and blood products storage facilities.
493.1277 Standard; Arrangement for services.
493.1279 Standard; Provision of testing.
493.1283 Standard; Retention of samples of transfused blood.
493.1285 Standard; Investigation of transfusion reactions.

Subpart L [Reserved]

Subpart M—Personnel for Moderate Complexity (Including the Subcategory) and High Complexity Testing

493.1351 General.

LABORATORIES PERFORMING PROVIDER-PERFORMED MICROSCOPY (PPM) PROCEDURES

493.1353 Scope.
493.1355 Condition: Laboratories performing PPM procedures; laboratory director.
493.1357 Standard; laboratory director qualifications.
493.1359 Standard; PPM laboratory director responsibilities.
493.1361 Condition: Laboratories performing PPM procedures; testing personnel.
493.1363 Standard; PPM testing personnel qualifications.
493.1365 Standard; PPM testing personnel responsibilities.

LABORATORIES PERFORMING MODERATE COMPLEXITY TESTING

493.1403 Condition; Laboratories performing moderate complexity testing; laboratory director.
493.1405 Standard; Laboratory director qualifications.
493.1406 Standard; Laboratory director qualifications on or before February 28, 1992.
493.1407 Standard; Laboratory director responsibilities.
493.1409 Condition; Laboratories performing moderate complexity testing; technical consultant.
493.1411 Standard; Technical consultant qualifications.
493.1413 Standard; Technical consultant responsibilities.
493.1415 Condition; Laboratories performing moderate complexity testing; clinical consultant.
493.1417 Standard; Clinical consultant qualifications.
493.1419 Standard; Clinical consultant responsibilities.
493.1421 Condition; Laboratories performing moderate complexity testing; clinical consultant.
493.1423 Standard; Testing personnel qualifications.
493.1425 Standard; Testing personnel responsibilities.

LABORATORIES PERFORMING HIGH COMPLEXITY TESTING

493.1441 Condition; Laboratories performing high complexity testing; laboratory director.
493.1443 Standard; Laboratory director qualifications.
493.1445 Standard; Laboratory director responsibilities.
493.1447 Condition; Laboratories performing high complexity testing; technical supervisor.
493.1449 Standard; Technical supervisor qualifications.
493.1451 Standard; Technical supervisor responsibilities.
493.1453 Condition; Laboratories performing high complexity testing; clinical consultant.
493.1455 Standard; Clinical consultant qualifications.
493.1457 Standard; Clinical consultant responsibilities.
493.1459 Condition; Laboratories performing high complexity testing; general supervisor.
493.1461 Standard; General supervisor qualifications.
493.1462 General supervisor qualifications on or before February 28, 1992.
493.1463 Standard; General supervisor responsibilities.
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493.1467 Condition: Laboratories performing high complexity testing; cytology general supervisor.
493.1469 Standard; Cytology general supervisor qualifications.
493.1471 Standard; Cytology general supervisor responsibilities.
493.1481 Condition: Laboratories performing high complexity testing; cytotechnologist.
493.1483 Standard; Cytotechnologist qualifications.
493.1485 Standard; Cytotechnologist responsibilities.
493.1487 Condition: Laboratories performing high complexity testing; testing personnel.
493.1489 Standard; Testing personnel qualifications.
493.1491 Technologist qualifications on or before February 28, 1992.
493.1495 Standard; Testing personnel responsibilities.

Subpart N–O [Reserved]

Subpart P—Quality Assurance for Moderate Complexity (Including the Subcategory) or High Complexity Testing, or Any Combination of These Tests

493.1701 Condition: Quality assurance; moderate complexity (including the subcategory) or high complexity testing, or any combination of these tests.
493.1703 Standard; Patient test management assessment.
493.1705 Standard; Quality control assessment.
493.1707 Standard; Proficiency testing assessment.
493.1709 Standard; Comparison of test results.
493.1711 Standard; Relationship of patient information to patient test results.
493.1713 Standard; Personnel assessment.
493.1715 Standard; Communications.
493.1717 Standard; Complaint investigations.
493.1719 Standard; Quality assurance review with staff.
493.1721 Standard; Quality assurance records.

Subpart Q—Inspection

493.1771 Condition: Inspection requirements applicable to all CLIA-certified and CLIA-exempt laboratories.
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493.1775 Standard; Inspection of laboratories issued a certificate of waiver or a certificate for provider-performed microscopy procedures.
493.1777 Standard; Inspection of laboratories that have requested or have been issued a certificate of compliance.
493.1780 Standard; Inspection of CLIA-exempt laboratories or laboratories requesting or issued a certificate of accreditation.

Subpart R—Enforcement Procedures

493.1800 Basis and scope.
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Subpart S [Reserved]

Subpart T—Consultations

493.2001 Establishment and function of the Clinical Laboratory Improvement Advisory Committee.

AUTHORITY: Sec. 353 of the Public Health Service Act, secs. 1102, 1861(e), the sentence following sections 1861(s)(11) through 1861(s)(16) of the Social Security Act (42 U.S.C. 263a, 1302, 1395x(e), the sentence following 1395x(s)(11) through 1395x(s)(16)).

SOURCE: 55 FR 9576, Mar. 14, 1990, unless otherwise noted.
§ 493.1 Basis and scope.
This part sets forth the conditions that all laboratories must meet to be certified to perform testing on human specimens under the Clinical Laboratory Improvement Amendments of 1988 (CLIA). It implements sections 1861(e) and (j), the sentence following section 1861(s)(13), and 1902(a)(9) of the Social Security Act, and section 353 of the Public Health Service Act. This part applies to all laboratories as defined under “laboratory” in § 493.2 of this part. This part also applies to laboratories seeking payment under the Medicare and Medicaid programs. The requirements are the same for Medicare approval as for CLIA certification.

§ 493.2 Definitions.
As used in this part, unless the context indicates otherwise—

Accredited institution means a school or program which—

(a) Admits as regular student only persons having a certificate of graduation from a school providing secondary education, or the recognized equivalent of such certificate;

(b) Is legally authorized within the State to provide a program of education beyond secondary education;

(c) Provides an educational program for which it awards a bachelor’s degree or provides not less than a 2-year program which is acceptable toward such a degree, or provides an educational program for which it awards a master’s or doctoral degree;

(d) Is accredited by a nationally recognized accrediting agency or association.

This definition includes any foreign institution of higher education that HHS or its designee determines meets substantially equivalent requirements.

Accredited laboratory means a laboratory that has voluntarily applied for and been accredited by a private, nonprofit accreditation organization approved by HCFA in accordance with this part;

Adverse action means the imposition of a principal or alternative sanction by HCFA.

ALJ stands for Administrative Law Judge.

Alternative sanctions means sanctions that may be imposed in lieu of or in addition to principal sanctions. The term is synonymous with “intermediate sanctions” as used in section 1846 of the Act.

Analyte means a substance or constituent for which the laboratory conducts testing.

Approved accreditation organization for laboratories means a private, nonprofit accreditation organization that has formally applied for and received HCFA’s approval based on the organization’s compliance with this part.

Approved State laboratory program means a licensure or other regulatory program for laboratories in a State, the requirements of which are imposed under State law, and the State laboratory program has received HCFA approval based on the State’s compliance with this part.

Authorized person means an individual authorized under State law to order tests or receive test results, or both.

Challenge means, for quantitative tests, an assessment of the amount of substance or analyte present or measured in a sample. For qualitative tests, a challenge means the determination of the presence or the absence of an analyte, organism, or substance in a sample.

CLIA means the Clinical Laboratory Improvement Amendments of 1988.

CLIA certificate means any of the following types of certificates issued by HCFA or its agent:

(1) Certificate of compliance means a certificate issued to a laboratory after an inspection that finds the laboratory to be in compliance with all applicable condition level requirements, or reissued before the expiration date, pending an appeal, in accordance with § 493.49, when an inspection has found the laboratory to be out of compliance with one or more condition level requirements.

(2) Certificate for provider-performed microscopy (PPM) procedures means a certificate issued or reissued before the
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expiration date, pending an appeal, in accordance with §493.47, to a laboratory in which a physician, midlevel practitioner or dentist performs no tests other than PPM procedures and, if desired, waived tests listed in §493.15(c).

(3) Certificate of accreditation means a certificate issued on the basis of the laboratory's accreditation by an accreditation organization approved by HCFA (indicating that the laboratory is deemed to meet applicable CLIA requirements) or reissued before the expiration date, pending an appeal, in accordance with §493.61, when a validation or complaint survey has found the laboratory to be noncompliant with one or more CLIA conditions.

(4) Certificate of registration or registration certificate means a certificate issued or reissued before the expiration date, pending an appeal, in accordance with §493.45, that enables the entity to conduct moderate or high complexity laboratory testing or both until the entity is determined to be in compliance through a survey by HCFA or its agent; or in accordance with §493.57 to an entity that is accredited by an approved accreditation organization.

(5) Certificate of waiver means a certificate issued or reissued before the expiration date, pending an appeal, in accordance with §493.37, to a laboratory to perform only the waived tests listed at §493.15(c).

CLIA-exempt laboratory means a laboratory that has been licensed or approved by a State where HCFA has determined that the State has enacted laws relating to laboratory requirements that are equal to or more stringent than CLIA requirements and the State licensure program has been approved by HCFA in accordance with subpart E of this part.

Condition level deficiency means noncompliance with one or more condition level requirements.

Condition level requirements means any of the requirements identified as “conditions” in subparts G through Q of this part.

Credible allegation of compliance means a statement or documentation that—

(1) Is made by a representative of a laboratory that has a history of having maintained a commitment to compliance and of taking corrective action when required;

(2) Is realistic in terms of its being possible to accomplish the required corrective action between the date of the exit conference and the date of the allegation; and

(3) Indicates that the problem has been resolved.

Dentist means a doctor of dental medicine or doctor of dental surgery licensed by the State to practice dentistry within the State in which the laboratory is located.

Equivalency means that an accreditation organization's or a State laboratory program's requirements, taken as a whole, are equal to or more stringent than the CLIA requirements established by HCFA, taken as a whole. It is acceptable for an accreditation organization's or State laboratory program's requirements to be organized differently or otherwise vary from the CLIA requirements, as long as (1) all of the requirements taken as a whole would provide at least the same protection as the CLIA requirements taken as a whole; and (2) a finding of noncompliance with respect to CLIA requirements taken as a whole would be matched by a finding of noncompliance with the accreditation or State requirements taken as a whole.

HCFA agent means an entity with which HCFA arranges to inspect laboratories and assess laboratory activities against CLIA requirements and may be a State survey agency, a private, nonprofit organization other than an approved accreditation organization, a component of HHS, or any other governmental component HCFA approves for this purpose. In those instances where all of the laboratories in a State are exempt from CLIA requirements, based on the approval of a State's exemption request, the State survey agency is not the HCFA agent.

HHS means the Department of Health and Human Services, or its designee.

Immediate jeopardy means a situation in which immediate corrective action is necessary because the laboratory's noncompliance with one or more condition level requirements has already caused, is causing, or is likely to cause, at any time, serious injury or harm, or
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death, to individuals served by the laboratory or to the health or safety of the general public. This term is synonymous with imminent and serious risk to human health and significant hazard to the public health.

Intentional violation means knowing and willful noncompliance with any CLIA condition.

Kit means all components of a test that are packaged together.

Laboratory means a facility for 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. Facilities only collecting or preparing specimens (or both) or only serving as a mailing service and not performing testing are not considered laboratories.

Midlevel practitioner means a nurse midwife, nurse practitioner, or physician assistant, licensed by the State within which the individual practices, if such licensing is required in the State in which the laboratory is located.

Operator means the individual or group of individuals who oversee all facets of the operation of a laboratory and who bear primary responsibility for the safety and reliability of the results of all specimen testing performed in that laboratory. The term includes—

1. A director of the laboratory if he or she meets the stated criteria; and

2. The members of the board of directors and the officers of a laboratory that is a small corporation under subchapter S of the Internal Revenue Code.

Owner means any person who owns any interest in a laboratory except for an interest in a laboratory whose stock and/or securities are publicly traded. (That is e.g., the purchase of shares of stock or securities on the New York Stock Exchange in a corporation owning a laboratory would not make a person an owner for the purpose of this regulation.)

Party means a laboratory affected by any of the enforcement procedures set forth in this subpart, by HCFA or the OIG, as appropriate.

Performance characteristic means a property of a test that is used to describe its quality, e.g., accuracy, precision, analytical sensitivity, analytical specificity, reportable range, reference range, etc.

Performance specification means a value or range of values for a performance characteristic, established or verified by the laboratory, that is used to describe the quality of patient test results.

Physician means an individual with a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine degree who is licensed by the State to practice medicine, osteopathy, or podiatry within the State in which the laboratory is located.

Principal sanction means the suspension, limitation, or revocation of any type of CLIA certificate or the cancellation of the laboratory's approval to receive Medicare payment for its services.

Prospective laboratory means a laboratory that is operating under a registration certificate or is seeking any of the three other types of CLIA certificates.

Rate of disparity means the percentage of sample validation inspections for a specific accreditation organization or State where HCFA, the State survey agency or other HCFA agent finds noncompliance with one or more condition level requirements but no comparable deficiencies were cited by the accreditation organization or State, and it is reasonable to conclude that the deficiencies were present at the time of the most recent accreditation organization or State licensure inspection.

EXAMPLE: Assume the State survey agency, HCFA or other HCFA agent performs 200 sample validation inspections for laboratories accredited by a single accreditation organization or licensed in an exempt State during a validation review period and finds that 60 of the 200 laboratories had one or more condition level requirements out of compliance. HCFA reviews the validation and accreditation organization's or State's
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inspections of the validated laboratories and determines that the State or accreditation organization found comparable deficiencies in 22 of the 60 laboratories and it is reasonable to conclude that deficiencies were present in the remaining 38 laboratories at the time of the accreditation organization’s or State’s inspection. Thirty-eight divided by 200 equals a 19 percent rate of disparity.

Referee laboratory means a laboratory currently in compliance with applicable CLIA requirements, that has had a record of satisfactory proficiency testing performance for all testing events for at least one year for a specific test, analyte, subspecialty, or specialty and has been designated by an HHS approved proficiency testing program as a referee laboratory for analyzing proficiency testing specimens for the purpose of determining the correct response for the specimens in a testing event for that specific test, analyte, subspecialty, or specialty.

Reference range means the range of test values expected for a designated population of individuals, e.g., 95 percent of individuals that are presumed to be healthy (or normal).

Sample in proficiency testing means the material contained in a vial, on a slide, or other unit that contains material to be tested by proficiency testing program participants. When possible, samples are of human origin.

State includes, for purposes of this part, each of the 50 States, the District of Columbia, the Commonwealth of Puerto Rico, the Virgin Islands and a political subdivision of a State where the State, acting pursuant to State law, has expressly delegated powers to the political subdivision sufficient to authorize the political subdivision to act for the State in enforcing requirements equal to or more stringent than CLIA requirements.

State licensure means the issuance of a license to, or the approval of, a laboratory by a State laboratory program as meeting standards for licensing or approval established under State law.

State licensure program means a State laboratory licensure or approval program.

State survey agency means the State health agency or other appropriate State or local agency that has an agreement under section 1864 of the Social Security Act and is used by HCF A to perform surveys and inspections.

Substantial allegation of noncompliance means a complaint from any of a variety of sources (including complaints submitted in person, by telephone, through written correspondence, or in newspaper or magazine articles) that, if substantiated, would have an impact on the health and safety of the general public or of individuals served by a laboratory and raises doubts as to a laboratory’s compliance with any condition level requirement.

Target value for quantitative tests means either the mean of all participant responses after removal of outliers (those responses greater than 3 standard deviations from the original mean) or the mean established by definitive or reference methods acceptable for use in the National Reference System for the Clinical Laboratory (NRSCl) by the National Committee for the Clinical Laboratory Standards (NCCLS). In instances where definitive or reference methods are not available or a specific method’s results demonstrate bias that is not observed with actual patient specimens, as determined by a defensible scientific protocol, a comparative method or a method group (“peer” group) may be used. If the method group is less than 10 participants, “target value” means the overall mean after outlier removal (as defined above) unless acceptable scientific reasons are available to indicate that such an evaluation is not appropriate.

Unsatisfactory proficiency testing performance means failure to attain the minimum satisfactory score for an analyte, test, subspecialty, or specialty for a testing event.

Unsuccessful participation in proficiency testing means any of the following:

1. Unsatisfactory performance for the same analyte in two consecutive or two out of three testing events.

2. Repeated unsatisfactory overall testing event scores for two consecutive or two out of three testing events for the same specialty or subspecialty.

3. An unsatisfactory testing event score for those subspecialties not graded by analyte (that is, bacteriology, mycobacteriology, virology, ...
§493.3 Applicability.

(a) Basic rule. Except as specified in paragraph (b) of this section, a laboratory will be cited as out of compliance with section 353 of the Public Health Service Act unless it—

1. Has a current, unrevoked or unsuspended certificate of waiver, registration certificate, certificate of compliance, certificate for PPM procedures, or certificate of accreditation issued by HHS applicable to the category of examinations or procedures performed by the laboratory; or

2. Is CLIA-exempt.

(b) Exception. These rules do not apply to components or functions of—

1. Any facility or component of a facility that only performs testing for forensic purposes;

2. Research laboratories that test human specimens but do not report patient specific results for the diagnosis, prevention or treatment of any disease or impairment of, or the assessment of the health of individual patients; or

3. Laboratories certified by the National Institutes on Drug Abuse (NIDA), in which drug testing is performed which meets NIDA guidelines and regulations. However, all other testing conducted by a NIDA-certified laboratory is subject to this rule.

(c) Federal laboratories. Laboratories under the jurisdiction of an agency of the Federal Government are subject to the rules of this part, except that the Secretary may modify the application of such requirements as appropriate.


§493.5 Categories of tests by complexity.

(a) Laboratory tests are categorized as one of the following:

1. Waived tests.

2. Tests of moderate complexity, including the subcategory of PPM procedures.

3. Tests of high complexity.

(b) A laboratory may perform only waived tests, only tests of moderate complexity, only PPM procedures, only tests of high complexity or any combination of these tests.

(c) Each laboratory must be either CLIA-exempt or possess one of the following CLIA certificates, as defined in §493.2:

1. Certificate of registration or registration certificate.

2. Certificate of waiver.


5. Certificate of accreditation.

[60 FR 20043, Apr. 24, 1995]

§493.15 Laboratories performing waived tests.

(a) Requirement. Tests for certificate of waiver must meet the descriptive criteria specified in paragraph (b) of this section.

(b) Criteria. Test systems are simple laboratory examinations and procedures which—

1. Are cleared by FDA for home use;

2. Employ methodologies that are so simple and accurate as to render the likelihood of erroneous results negligible; or

3. Pose no reasonable risk of harm to the patient if the test is performed incorrectly.

(c) Certificate of waiver. A laboratory may qualify for a certificate of waiver under section 535 of the PHS
§ 493.17 Test categorization.

(a) Categorization by criteria. Notices will be published in the Federal Register which list each specific test system, assay, and examination categorized by complexity. Using the seven criteria specified in this paragraph for categorizing tests of moderate or high complexity, each specific laboratory test system, assay, and examination will be graded for level of complexity by assigning scores of 1, 2, or 3 within each criteria. The score of "1" indicates the lowest level of complexity, and the score of "3" indicates the highest level. These scores will be totaled. Test systems, assays or examinations receiving scores of 12 or less will be categorized as moderate complexity, while those receiving scores above 12 will be categorized as high complexity.

NOTE: A score of "2" will be assigned to a criteria heading when the characteristics for a particular test are intermediate between the descriptions listed for scores of "1" and "3."

(1) Knowledge.

(i) Score 1. (A) Minimal scientific and technical knowledge is required to perform the test; and
(B) Knowledge required to perform the test may be obtained through on-the-job instruction.

(ii) Score 3. (A) Specialized scientific and technical knowledge is essential to perform preanalytic, analytic or postanalytic phases of the testing; or
(B) Substantial experience may be necessary for analytic test performance.

(2) Training and experience.

(i) Score 1. (A) Minimal training is required for preanalytic, analytic and postanalytic phases of the testing process; and
(B) Limited experience is required to perform the test.

(ii) Score 3. (A) Specialized training is essential to perform the preanalytic, analytic or postanalytic testing process; or
(B) Substantial experience may be necessary for analytic test performance.

(3) Reagents and materials preparation.

(i) Score 1. (A) Reagents and materials are generally stable and reliable; and
(B) Reagents and materials are prepackaged, or premeasured, or require...
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no special handling, precautions or storage conditions.

(ii) Score 3. (A) Reagents and materials may be labile and may require special handling to assure reliability; or

(B) Reagents and materials preparation may include manual steps such as gravimetric or volumetric measurements.

(4) Characteristics of operational steps.

(i) Score 1. Operational steps are either automatically executed (such as pipetting, temperature monitoring, or timing of steps), or are easily controlled.

(ii) Score 3. Operational steps in the testing process require close monitoring or control, and may require special specimen preparation, precise temperature control or timing of procedural steps, accurate pipetting, or extensive calculations.

(5) Calibration, quality control, and proficiency testing materials.

(i) Score 1. (A) Calibration materials are stable and readily available; (B) Quality control materials are stable and readily available; and (C) External proficiency testing materials, when available, are stable.

(ii) Score 3. (A) Calibration materials, if available, may be labile; (B) Quality control materials may be labile, or not available; or (C) External proficiency testing materials, if available, may be labile.

(6) Test system troubleshooting and equipment maintenance.

(i) Score 1. (A) Test system troubleshooting is automatic or self-correcting, or clearly described or requires minimal judgment; and (B) Equipment maintenance is provided by the manufacturer, is seldom needed, or can easily be performed.

(ii) Score 3. (A) Troubleshooting is not automatic and requires decision-making and direct intervention to resolve most problems; or (B) Maintenance requires special knowledge, skills, and abilities.

(7) Interpretation and judgment.

(i) Score 1. (A) Minimal interpretation and judgment are required to perform preanalytic, analytic and postanalytic processes; and (B) Resolution of problems requires limited independent interpretation and judgment; and

(ii) Score 3. (A) Extensive independent interpretation and judgment are required to perform the preanalytic, analytic or postanalytic processes; and (B) Resolution of problems requires extensive interpretation and judgment.

(b) Revisions to the criteria for categorization. The Clinical Laboratory Improvement Advisory Committee, as defined in subpart T of this part, will conduct reviews upon request of HHS and recommend to HHS revisions to the criteria for categorization of tests.

(c) Process for device/test categorization utilizing the scoring system under § 493.17(a). (1)(i) For new commercial test systems, assays, or examinations, the manufacturer, as part of its 510(k) and PMA application to FDA, will submit supporting data for device/test categorization. FDA will determine the complexity category, notify the manufacturers directly, and will simultaneously inform both HCFA and CDC of the device/test category. FDA will consult with CDC concerning test categorization in the following three situations:

(A) When categorizing previously uncategorized new technology;

(B) When FDA determines it to be necessary in cases involving a request for a change in categorization; and

(C) If a manufacturer requests review of a categorization decision by FDA in accordance with 21 CFR 10.75.

(ii) Test categorization will be effective as of the notification to the applicant.

(2) For test systems, assays, or examinations not commercially available, a laboratory or professional group may submit a written request for categorization to PHS. These requests will be forwarded to CDC for evaluation; CDC will determine complexity category and notify the applicant, HCFA, and FDA of the categorization decision. In the case of request for a change of category or for previously uncategorized new technology, PHS will receive the request application and forward it to CDC for categorization.

(3) A request for recategorization will be accepted for review if it is based on
§ 493.19 Provider-performed microscopy (PPM) procedures.

(a) Requirement. To be categorized as a PPM procedure, the procedure must meet the criteria specified in paragraph (b) of this section.

(b) Criteria. Procedures must meet the following specifications:

(1) The examination must be personally performed by one of the following practitioners:

   (i) A physician during the patient's visit on a specimen obtained from his or her own patient or from a patient of a group medical practice of which the physician is a member or an employee.

   (ii) A midlevel practitioner, under the supervision of a physician or in independent practice only if authorized by the State, during the patient's visit on a specimen obtained from his or her own patient or from a patient of a clinic, group medical practice, or other health care provider of which the midlevel practitioner is a member or an employee.

   (iii) A dentist during the patient's visit on a specimen obtained from his or her own patient or from a patient of a group dental practice of which the dentist is a member or an employee.

(2) The procedure must be categorized as moderately complex.

(3) The primary instrument for performing the test is the microscope, limited to bright-field or phase-contrast microscopy.

(4) The specimen is labile or delay in performing the test could compromise the accuracy of the test result.

(5) Control materials are not available to monitor the entire testing process.

(6) Limited specimen handling or processing is required.

(c) Provider-performed microscopy (PPM) examinations. A laboratory may qualify to perform tests under this section if it restricts PPM examinations to one or more of the following procedures (or additional procedures added to this list as provided under paragraph (d) of this section), waived tests and no others:

(1) All direct wet mount preparations for the presence or absence of bacteria, fungi, parasites, and human cellular elements.

(2) All potassium hydroxide (KOH) preparations.

(3) Pinworm examinations.

(4) Fern tests.

(5) Post-coital direct, qualitative examinations of vaginal or cervical mucus.

(6) Urine sediment examinations.

(7) Nasal smears for granulocytes.

(8) Fecal leukocyte examinations.

(9) Qualitative semen analysis (limited to the presence or absence of sperm and detection of motility).

(d) Revisions to criteria and the list of PPM procedures.

(1) The CLIAC conducts reviews upon HHS' request and recommends to HHS revisions to the criteria for categorization of procedures.

(2) HHS determines whether a laboratory procedure meets the criteria listed under paragraph (b) of this section for a PPM procedure. Revisions to the list of PPM procedures proposed by HHS are published in the Federal Register as a notice with an opportunity for public comment.

(e) Laboratory requirements. Laboratories eligible to perform PPM examinations must—

(1) Meet the applicable requirements in subpart C or subpart D, and subparts F, H, J, K, M, and P of this part.

(2) Be subject to inspection as specified under subpart Q of this part.

[60 FR 20044, Apr. 24, 1995]
§ 493.20 Laboratories performing tests of moderate complexity.

(a) A laboratory may qualify for a certificate to perform tests of moderate complexity provided that it restricts its test performance to waived tests or examinations and one or more tests or examinations meeting criteria for tests of moderate complexity including the subcategory of PPM procedures.

(b) A laboratory that performs tests or examinations of moderate complexity must meet the applicable requirements in subpart C or subpart D, and subparts F, H, J, K, M, P, and Q of this part. Under a registration certificate or certificate of compliance, laboratories also performing PPM procedures must meet the inspection requirements at §493.1777.

(c) If the laboratory also performs waived tests, compliance with subparts H, J, K, M, and P of this part is not applicable to the waived tests. However, the laboratory must comply with the requirements in §§493.15(e) and 493.1775.

[60 FR 20044, Apr. 24, 1995]

§ 493.25 Laboratories performing tests of high complexity.

(a) A laboratory must obtain a certificate for tests of high complexity if it performs one or more tests that meet the criteria for tests of high complexity as specified in §493.17(a).

(b) A laboratory performing one or more tests of high complexity must meet the applicable requirements of subpart C or subpart D, and subparts F, H, J, K, M, P, and Q of this part.

(c) If the laboratory also performs waived tests, compliance with subparts H, J, K, M, and P of this part must be met. Under a registration certificate or certificate of compliance, PPM procedures must meet the inspection requirements at §493.1777.

(d) If the laboratory also performs waived tests, the requirements of subparts H, J, K, M, and P are not applicable to the waived tests. However, the laboratory must comply with the requirements in §§493.15(e) and 493.1775.

[57 FR 7142, Feb. 28, 1992, as amended at 60 FR 20044, Apr. 24, 1995]

Subpart B—Certificate of Waiver

§ 493.35 Application for a certificate of waiver.

(a) Filing of application. Except as specified in paragraph (b) of this section, a laboratory performing only one or more waived tests listed in §493.15 must file a separate application for each laboratory location.

(b) Exceptions. (1) Laboratories that are not at a fixed location, that is, laboratories that move from testing site to testing site, such as mobile units providing laboratory testing, health screening fairs, or other temporary testing locations may be covered under the certificate of the designated primary site or home base, using its address.

(2) Not-for-profit or Federal, State, or local government laboratories that engage in limited (not more than a combination of 15 moderately complex or waived tests per certificate) public health testing may file a single application.

(3) Laboratories within a hospital that are located at contiguous buildings on the same campus and under common direction may file a single application or multiple applications for the laboratory sites within the same physical location or street address.

(c) Application format and contents. The application must—

(1) Be made to HHS or its designee on a form or forms prescribed by HHS;

(2) Be signed by an owner, or by an authorized representative of the laboratory who attests that the laboratory will be operated in accordance with requirements established by the Secretary under section 353 of the PHS Act; and

(3) Describe the characteristics of the laboratory operation and the examinations and other test procedures performed by the laboratory including—

(i) The name and the total number of test procedures and examinations performed annually (excluding tests the laboratory may run for quality control, quality assurance or proficiency testing purposes;
§ 493.37 Requirements for a certificate of waiver.

(a) HHS will issue a certificate of waiver to a laboratory only if the laboratory meets the requirements of §493.35.

(b) Laboratories issued a certificate of waiver—

(1) Are subject to the requirements of this subpart and §493.15(e) of subpart A of this part; and

(2) Must permit announced or unannounced inspections by HHS in accordance with subpart Q of this part.

(c) Laboratories must remit the certificate of waiver fee specified in subpart F of this part.

(d) In accordance with subpart R of this part, HHS will suspend or revoke or limit a laboratory’s certificate of waiver for failure to comply with the requirements of this subpart. In addition, failure to meet the requirements of this subpart will result in suspension or denial of payments under Medicare and Medicaid in accordance with subpart R of this part.

(e)(1) A certificate of waiver issued under this subpart is valid for no more than 2 years. In the event of a non-compliance determination resulting in HHS action to revoke, suspend, or limit the laboratory’s certificate of waiver, HHS will provide the laboratory with a statement of grounds on which the determination of non-compliance is based and offer an opportunity for appeal as provided in subpart R of this part.

(2) If the laboratory requests a hearing within the time specified by HHS, it retains its certificate of waiver or reissued certificate of waiver until a decision is made by an administrative law judge, as specified in subpart R of this part, except when HHS finds that conditions at the laboratory pose an imminent and serious risk to human health.

(3) For laboratories receiving payment from the Medicare or Medicaid program, such payments will be suspended on the effective date specified in the notice to the laboratory of a non-compliance determination even if there has been no appeals decision issued.

(f) A laboratory seeking to renew its certificate of waiver must—
§ 493.39 Notification requirements for laboratories issued a certificate of waiver.

Laboratories performing one or more tests listed in §493.15 and no others must notify HHS or its designee—

(a) Before performing and reporting results for any test or examination that is not specified under §493.15 for which the laboratory does not have the appropriate certificate as required in subpart C or subpart D of this part, as applicable; and

(b) Within 30 days of any change(s) in—

(1) Ownership;
(2) Name;
(3) Location; or
(4) Director.

§ 493.39 Notification requirements for laboratories issued a certificate of waiver.

(1) Complete the renewal application prescribed by HHS and return it to HHS not less than 9 months nor more than 1 year before the expiration of the certificate; and

(2) Meet the requirements of §§493.35 and 493.37.

(g) A laboratory with a certificate of waiver that wishes to perform examinations or tests not listed in the waiver test category must meet the requirements set forth in subpart C or subpart D of this part, as applicable.

§ 493.39 Notification requirements for laboratories issued a certificate of waiver.

Laboratories performing one or more tests listed in §493.15 and no others must notify HHS or its designee—

(a) Before performing and reporting results for any test or examination that is not specified under §493.15 for which the laboratory does not have the appropriate certificate as required in subpart C or subpart D of this part, as applicable; and

(b) Within 30 days of any change(s) in—

(1) Ownership;
(2) Name;
(3) Location; or
(4) Director.

§ 493.39 Notification requirements for laboratories issued a certificate of waiver.

(1) Complete the renewal application prescribed by HHS and return it to HHS not less than 9 months nor more than 1 year before the expiration of the certificate; and

(2) Meet the requirements of §§493.35 and 493.37.

(g) A laboratory with a certificate of waiver that wishes to perform examinations or tests not listed in the waiver test category must meet the requirements set forth in subpart C or subpart D of this part, as applicable.

(1) Be made to HHS or its designee on a form or forms prescribed by HHS;

(2) Be signed by an owner, or by an authorized representative of the laboratory who attests that the laboratory will be operated in accordance with the requirements established by the Secretary under section 353 of the Public Health Service Act; and

(3) Describe the characteristics of the laboratory operation and the examinations and other test procedures performed by the laboratory including—

(i) The name and total number of test procedures and examinations performed annually (excluding waived tests or tests for quality control, quality assurance or proficiency testing purposes);

(ii) The methodologies for each laboratory test procedure or examination performed, or both;

(iii) The qualifications (educational background, training, and experience) of the personnel directing and supervising the laboratory and performing the examinations and test procedures.

(d) Access and reporting requirements. All laboratories must make records available and submit reports to HHS as
§ 493.45 Requirements for a registration certificate.

Laboratories performing only waived tests, PPM procedures, or any combination of these tests, are not required to obtain a registration certificate.

(a) A registration certificate is required—(1) Initially for all laboratories performing test procedures of moderate complexity (other than the subcategory of PPM procedures) or high complexity, or both; and

(2) For all laboratories that have been issued a certificate of waiver or certificate for PPM procedures that intend to perform tests of moderate or high complexity, or both, in addition to those tests listed in §493.15(c) or specified as PPM procedures.

(b) HHS will issue a registration certificate if the laboratory—

(1) Complies with the requirements of §493.43;

(2) Agrees to notify HHS or its designee within 30 days of any changes in ownership, name, location, director or technical supervisor (laboratories performing high complexity testing only);

(3) Agrees to treat proficiency testing samples in the same manner as it treats patient specimens; and

(4) Remits the fee for the registration certificate, as specified in subpart F of this part.

(c) Prior to the expiration of the registration certificate, a laboratory must—

(1) Remit the certificate fee specified in subpart F of this part;

(2) Be inspected by HHS as specified in subpart Q of this part; and

(3) Demonstrate compliance with the applicable requirements of this subpart and subparts H, J, K, M, P, and Q of this part.

(d) In accordance with subpart R of this part, HHS will initiate suspension or revocation of a laboratory's registration certificate and will deny the laboratory's application for a certificate of compliance for failure to comply with the requirements set forth in this subpart. HHS may also impose certain alternative sanctions. In addition, failure to meet the requirements of this subpart will result in suspension of payments under Medicare and Medicaid as specified in subpart R of this part.

(e) A registration certificate is—

(1) Valid for a period of no more than two years or until such time as an inspection to determine program compliance can be conducted, whichever is shorter; and

(2) Not renewable; however, the registration certificate may be reissued if compliance has not been determined by HHS prior to the expiration date of the registration certificate.

(f) In the event of a noncompliance determination resulting in an HHS denial of a laboratory's certificate of compliance application, HHS will provide the laboratory with a statement of grounds on which the noncompliance determination is based and offer an opportunity for appeal as provided in subpart R.

(g) If the laboratory requests a hearing within the time specified by HHS, it retains its registration certificate or reissued registration certificate until a decision is made by an administrative law judge as provided in subpart R of this part, except when HHS finds that conditions at the laboratory pose an imminent and serious risk to human health.

(h) For laboratories receiving payment from the Medicare or Medicaid program, such payments will be suspended on the effective date specified in the notice to the laboratory of denial of the certificate application even if there has been no appeals decision issued.

§ 493.47 Requirements for a certificate for provider-performed microscopy (PPM) procedures.

(a) A certificate for PPM procedures is required—

(1) Initially for all laboratories performing test procedures specified as PPM procedures; and

(2) For all certificate of waiver laboratories that intend to perform only
§493.49  Requirements for a certificate of compliance.

A certificate of compliance may include any combination of tests categorized as high complexity or moderate complexity or listed in §493.15(c) as waived tests. Moderate complexity tests may include those specified as PPM procedures.

(a) HHS will issue a certificate of compliance to a laboratory only if the laboratory—

(1) Meets the requirements of §§493.43 and 493.45;

(2) Remits the certificate fee specified in subpart F of this part; and

(3) Meets the applicable requirements of this subpart and subparts H, J, K, M, P, and Q of this part.

(b) Laboratories issued a certificate of compliance—

(1) Are subject to the notification requirements of §493.51; and

(2) Must permit announced or unannounced inspections by HHS in accordance with subpart O of this part—

(i) To determine compliance with the applicable requirements of this part;

(ii) To evaluate complaints;

(iii) When HHS has substantive reason to believe that tests are being performed, or the laboratory is being operated in a manner that constitutes an imminent and serious risk to human health; and

(iv) To collect information regarding the appropriateness of tests listed in §493.15 or tests categorized as moderate complexity (including the subcategory) or high complexity.

(c) Failure to comply with the requirements of this subpart will result in—

(1) Suspension, revocation or limitation of a laboratory's certificate of compliance in accordance with subpart R of this part; and

(2) Suspension or denial of payments under Medicare and Medicaid in accordance with subpart R of this part.

(d) A certificate of compliance issued under this subpart is valid for no more than 2 years.

(e) In the event of a noncompliance determination resulting in an HHS action to revoke, suspend or limit the laboratory's certificate of compliance, HHS will—

(1) Provide the laboratory with a statement of grounds on which the determination of noncompliance is based; and

(2) Offer an opportunity for appeal as provided in subpart R of this part. If the laboratory requests a hearing within 60 days of the notice of sanction, it retains its certificate of compliance or reissued certificate of compliance until a decision is made by an administrative law judge (ALJ) as provided in subpart R of this part, except when HHS finds that conditions at the laboratory pose an imminent and serious risk to human health or when the criteria at §493.1840(a) (4) and (5) are met.

[58 FR 5223, Jan. 19, 1993, as amended at 60 FR 20045, Apr. 24, 1995]
§ 493.55 Application for registration certificate and certificate of accreditation.

(a) Filing of application. A laboratory may be issued a certificate of accreditation in lieu of the applicable certificate specified in subpart B or subpart C of this part provided the laboratory—
(1) Meets the standards of a private non-profit accreditation program approved by HHS in accordance with subpart E; and

(2) Files a separate application for each location, except as specified in paragraph (b) of this section.

(b) Exceptions. (1) Laboratories that are not at fixed locations, that is, laboratories that move from testing site to testing site, such as mobile units providing laboratory testing, health screening fairs, or other temporary testing locations may be covered under the certificate of the designated primary site or home base, using its address.

(2) Not-for-profit or Federal, State, or local government laboratories that engage in limited (not more than a combination of 15 moderately complex or waived tests per certificate) public health testing may file a single application.

(3) Laboratories within a hospital that are located at contiguous buildings on the same campus and under common direction may file a single application or multiple applications for the laboratory sites within the same physical location or street address.

(c) Application format and contents. The application must—(1) Be made to HHS on a form or forms prescribed by HHS;

(2) Be signed by an owner or authorized representative of the laboratory who attests that the laboratory will be operated in accordance with the requirements established by the Secretary under section 353 of the Public Health Service Act; and

(3) Describe the characteristics of the laboratory operation and the examinations and other test procedures performed by the laboratory including—

(i) The name and total number of tests and examinations performed annually (excluding waived tests and tests for quality control, quality assurance or proficiency testing purposes);

(ii) The methodologies for each laboratory test procedure or examination performed, or both; and

(iii) The qualifications (educational background, training, and experience) of the personnel directing and supervising the laboratory and performing the laboratory examinations and test procedures.

(d) Access and reporting requirements. All laboratories must make records available and submit reports to HHS as HHS may reasonably require to determine compliance with this section.


§ 493.57 Requirements for a registration certificate.

A registration certificate is required for all laboratories seeking a certificate of accreditation, unless the laboratory holds a valid certificate of compliance issued by HHS.

(a) HHS will issue a registration certificate if the laboratory—

(1) Complies with the requirements of §493.55;

(2) Agrees to notify HHS within 30 days of any changes in ownership, name, location, director, or supervisor (laboratories performing high complexity testing only);

(3) Agrees to treat proficiency testing samples in the same manner as it treats patient specimens; and

(4) Remits the fee for the registration certificate specified in subpart F of this part.

(b)(1) The laboratory must provide HHS with proof of accreditation by an approved accreditation program—

(i) Within 11 months of issuance of the registration certificate; or

(ii) Prior to the expiration of the certificate of compliance.

(2) If such proof of accreditation is not supplied within this timeframe, the laboratory must meet, or continue to meet, the requirements of §493.49.

(c) In accordance with subpart R of this part, HHS will initiate suspension, revocation, or limitation of a laboratory's registration certificate and will deny the laboratory's application for a certificate of accreditation for failure to comply with the requirements set forth in this subpart. In addition, failure to meet the requirements of this subpart will result in suspension or denial of payments under Medicare and Medicaid as specified in subpart R of this part.

(d) A registration certificate is valid for a period of no more than 2 years.
However, it may be reissued if the laboratory is subject to subpart C of this part, as specified in §493.57(b)(2) and compliance has not been determined by HHS before the expiration date of the registration certificate.

(e) In the event that the laboratory does not meet the requirements of this subpart, HHS will—

(1) Deny a laboratory’s request for certificate of accreditation;
(2) Notify the laboratory if it must meet the requirements for a certificate as defined in subpart C of this part;
(3) Provide the laboratory with a statement of grounds on which the application denial is based;
(4) Offer an opportunity for appeal on the application denial as provided in subpart R of this part. If the laboratory requests a hearing within the time specified by HHS, the laboratory will retain its registration certificate or reissued registration certificate until a decision is made by an administrative law judge as provided in subpart R, unless HHS finds that conditions at the laboratory pose an imminent and serious risk to human health; and
(5) For those laboratories receiving payment from the Medicare or Medicaid program, such payments will be suspended on the effective date specified in the notice to the laboratory of denial of the request even if there has been no appeals decision issued.

§ 493.61 Requirements for a certificate of accreditation.

(a) HHS will issue a certificate of accreditation to a laboratory if the laboratory—

(1) Meets the requirements of §493.57 or, if applicable, §493.49 of subpart C of this part; and
(2) Remits the certificate of accreditation fee specified in subpart F of this part.

(b) Laboratories issued a certificate of accreditation must—

(1) Treat proficiency testing samples in the same manner as patient samples;
(2) Meet the requirements of §493.63;
(3) Comply with the requirements of the approved accreditation program;
(4) Permit random sample validation and complaint inspections as required in subpart Q of this part;
(5) Permit HHS to monitor the correction of any deficiencies found through the inspections specified in paragraph (b)(4) of this section;
(6) Authorize the accreditation program to release to HHS the laboratory’s inspection findings whenever HHS conducts random sample or complaint inspections; and
(7) Authorize its accreditation program to submit to HHS the results of the laboratory’s proficiency testing.

(c) A laboratory failing to meet the requirements of this section—

(1) Will no longer meet the requirements of this part by virtue of its accreditation in an approved accreditation program;
(2) Will be subject to full determination of compliance by HHS;
(3) May be subject to suspension, revocation or limitation of the laboratory’s certificate of accreditation or certain alternative sanctions; and
(4) May be subject to suspension of payments under Medicare and Medicaid as specified in subpart R.

(d) A certificate of accreditation issued under this subpart is valid for no more than 2 years. In the event of a non-compliance determination as a result of a random sample validation or complaint inspection, a laboratory will be subject to a full review by HHS in accordance with §488.11 of this chapter.

(e) Failure to meet the applicable requirements of part 493, will result in an action by HHS to suspend, revoke or limit the certificate of accreditation. HHS will—

(1) Provide the laboratory with a statement of grounds on which the determination of noncompliance is based;
(2) Notify the laboratory if it is eligible to apply for a certificate as defined in subpart C of this part; and
(3) Offer an opportunity for appeal as provided in subpart R of this part.

(f) If the laboratory requests a hearing within the time frame specified by HHS—

(1) It retains its certificate of accreditation or reissued certificate of accreditation until a decision is made by an administrative law judge as provided in subpart R of this part, unless
§ 493.63 Notification requirements for laboratories issued a certificate of accreditation.

Laboratories issued a certificate of accreditation must:

(a) Notify HHS and the approved accreditation program within 30 days of any changes in—

(1) Ownership;

(2) Name;

(3) Location; or

(4) Director.

(b) Notify the approved accreditation program no later than 6 months after performing any test or examination within a specialty or subspecialty area that is not included in the laboratory’s accreditation, so that the accreditation organization can determine compliance and a new certificate of accreditation can be issued.

(c) Notify the accreditation program no later than 6 months after of any deletions or changes in test methodologies for any test or examination included in a specialty or subspecialty, or both, for which the laboratory has been issued a certificate of accreditation.

Subpart E—Accreditation by a Private, Nonprofit Accreditation Organization or Exemption Under an Approved State Laboratory Program

SOURCE: 63 FR 26732, May 14, 1998, unless otherwise noted.

§ 493.551 General requirements for laboratories.

(a) Applicability. HCFA may deem a laboratory to meet all applicable CLIA program requirements through accreditation by a private nonprofit accreditation program (that is, grant deemed status), or may exempt from CLIA program requirements all State licensed or approved laboratories in a State that has a State licensure program established by law, if the following conditions are met:

(1) The requirements of the accreditation organization or State licensure program are equal to, or more stringent than, the CLIA condition-level requirements specified in this part, and
the laboratory would meet the condition-level requirements if it were inspected against these requirements.

(2) The accreditation program or the State licensure program meets the requirements of this subpart and is approved by HCFA.

(3) The laboratory authorizes the approved accreditation organization or State licensure program to release to HCFA all records and information required and permits inspections as outlined in this part.

(b) Meeting CLIA requirements by accreditation. A laboratory seeking to meet CLIA requirements through accreditation by an approved accreditation organization must do the following:

(1) Obtain a certificate of accreditation as required in subpart D of this part.

(2) Pay the applicable fees as required in subpart F of this part.

(3) Meet the proficiency testing (PT) requirements in subpart H of this part.

(4) Authorize its PT organization to furnish to its accreditation organization the results of the laboratory’s participation in an approved PT program for the purpose of monitoring the laboratory’s PT and for making the annual PT results, along with explanatory information required to interpret the PT results, available on a reasonable basis, upon request of any person.

A laboratory that refuses to authorize release of its PT results is no longer deemed to meet the condition-level requirements and is subject to a full review by HCFA, in accordance with subpart Q of this part, and may be subject to the suspension or revocation of its certificate of accreditation under §493.1840.

(5) Authorize its accreditation organization to release to HCFA or a HCFA agent the laboratory’s PT results that constitute unsuccessful participation in an approved PT program, in accordance with the definition of “unsuccessful participation in an approved PT program,” as specified in §493.2 of this part, when the laboratory has failed to achieve successful participation in an approved PT program.

(6) Authorize its accreditation organization to release to HCFA a notification of the actions taken by the organization as a result of the unsuccessful participation in a PT program within 30 days of the initiation of the action. Based on this notification, HCFA may take an adverse action against a laboratory that fails to participate successfully in an approved PT program.

(c) Withdrawal of laboratory accreditation. After an accreditation organization has withdrawn or revoked its accreditation of a laboratory, the laboratory retains its certificate of accreditation for 45 days after the laboratory receives notice of the withdrawal or revocation of the accreditation, or the effective date of any action taken by HCFA, whichever is earlier.

§493.553 Approval process (application and reapplication) for accreditation organizations and State licensure programs.

(a) Information required. An accreditation organization that applies or re-applies to HCFA for deeming authority, or a State licensure program that applies or re-applies to HCFA for exemption from CLIA program requirements of licensed or approved laboratories within the State, must provide the following information:

(1) A detailed comparison of the individual accreditation, or licensure or approval requirements with the comparable condition-level requirements; that is, a crosswalk.

(2) A detailed description of the inspection process, including the following:

(i) Frequency of inspections.

(ii) Copies of inspection forms.

(iii) Instructions and guidelines.

(iv) A description of the review and decision-making process of inspections.

(v) A statement concerning whether inspections are announced or unannounced.

(vi) A description of the steps taken to monitor the correction of deficiencies.

(3) A description of the process for monitoring PT performance, including action to be taken in response to unsuccessful participation in a HCFA-approved PT program.

(4) Procedures for responding to and for the investigation of complaints against its laboratories.

(5) A list of all its current laboratories and the expiration date of their
§ 493.555 Federal review of laboratory requirements.

HCFA's review of an accreditation organization or State licensure program includes, but is not limited to, an evaluation of the following:

(a) Whether the organization's or State's requirements for laboratories are equal to, or more stringent than, the condition-level requirements for laboratories.

(b) The organization's or State's inspection process to determine the comparability of the full inspection and complaint inspection procedures and requirements to those of HCFA, including, but not limited to, inspection frequency and the ability to investigate and respond to complaints against its laboratories.

(c) The organization's or State's agreement with HCFA that requires it to do the following:

(1) Notify HCFA within 30 days of the action taken, of any laboratory that has—
   (i) Had its accreditation or licensure suspended, withdrawn, revoked, or limited;
   (ii) In any way been sanctioned; or
   (iii) Had any adverse action taken against it.

(2) Notify HCFA within 10 days of any deficiency identified in an accredited or CLIA-exempt laboratory if the deficiency poses an immediate jeopardy to the laboratory's patients or a hazard to the general public.

(3) Notify HCFA, within 30 days, of all newly—
   (i) Accredited laboratories (or laboratories whose areas of specialty/subspecialty testing have changed); or
   (ii) Licensed laboratories, including the specialty/subspecialty areas of testing.

(4) Notify each accredited or licensed laboratory within 10 days of HCFA's withdrawal of the organization's deeming authority or State's exemption.

(5) Provide HCFA with inspection schedules, as requested, for validation purposes.

§ 493.557 Additional submission requirements.

(a) Specific requirements for accreditation organizations. In addition to the information specified in §§ 493.553 and 493.555, as part of the approval and review process, an accreditation organization applying or reapplying for deeming authority must also provide the following:

(1) The specialty or subspecialty areas for which the organization is requesting deeming authority and its mechanism for monitoring compliance with all requirements equivalent to condition-level requirements within the scope of the specialty or subspecialty areas.

(2) A description of the organization's data management and analysis system.
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with respect to its inspection and accreditation decisions, including the kinds of routine reports and tables generated by the systems.

(3) Detailed information concerning the inspection process, including, but not limited to the following:

(i) The size and composition of individual accreditation inspection teams.

(ii) Qualifications, education, and experience requirements that inspectors must meet.

(iii) The content and frequency of training provided to inspection personnel, including the ability of the organization to provide continuing education and training to inspectors.

(4) Procedures for removal or withdrawal of accreditation status for laboratories that fail to meet the organization’s standards.

(5) A proposed agreement between HCFA and the accreditation organization with respect to the notification requirements specified in § 493.555(c).

(6) Procedures for monitoring laboratories found to be out of compliance with its requirements. (These monitoring procedures must be used only when the accreditation organization identifies noncompliance. If noncompliance is identified through validation inspections, HCFA or a HCFA agent monitors corrections, as authorized at § 493.565(d)).

(7) A demonstration of its ability to provide HCFA with electronic data and reports in compatible code, including the crosswalk specified in § 493.553(a)(1), that are necessary for effective validation and assessment of the organization’s inspection process.

(8) A demonstration of its ability to provide HCFA with electronic data, in compatible code, related to the adverse actions resulting from PT results constituting unsuccessful participation in PT programs as well as data related to the PT failures, within 30 days of the initiation of adverse action.

(9) A demonstration of its ability to provide HCFA with electronic data, in compatible code, for all accredited laboratories, including the area of specialty or subspecialty.

(10) Information defining the adequacy of numbers of staff and other resources.

(11) Information defining the organization’s ability to provide adequate funding for performing required inspections.

(12) Any facility-specific data, upon request by HCFA, which includes, but is not limited to, the following:

(i) PT results that constitute unsuccessful participation in a HCFA-approved PT program.

(ii) Notification of the adverse actions or corrective actions imposed by the accreditation organization as a result of unsuccessful PT participation.

(13) An agreement to provide written notification to HCFA at least 30 days in advance of the effective date of any proposed change in its requirements.

(14) An agreement to disclose any laboratory’s PT results upon reasonable request by any person.

(b) Specific requirements for a State licensure program. In addition to requirements in §§ 493.553 and 493.555, as part of the approval and review process, when a State licensure program applies or reapplies for exemption from the CLIA program, the State must do the following:

(1) Demonstrate to HCFA that it has enforcement authority and administrative structures and resources adequate to enforce its laboratory requirements.

(2) Permit HCFA or a HCFA agent to inspect laboratories in the State.

(3) Require laboratories in the State to submit to inspections by HCFA or a HCFA agent as a condition of licensure or approval.

(4) Agree to pay the cost of the validation program administered in that State as specified in §§ 493.645(a) and 493.646(b).

(5) Take appropriate enforcement action against laboratories found by HCFA not to be in compliance with requirements equivalent to CLIA requirements.

(6) Submit for Medicare and Medicaid payment purposes, a list of the specialties and subspecialties of tests performed by each laboratory.

(7) Submit a written presentation that demonstrates the agency’s ability to furnish HCFA with electronic data in compatible code, including the crosswalk specified in § 493.553(a)(1).

(8) Submit a statement acknowledging that the State will notify HCFA
through electronic transmission of the following:

(i) Any laboratory that has had its license or approval revoked or withdrawn or has been in any way sanctioned by the State within 30 days of taking the action.

(ii) Changes in licensure or inspection requirements.

(iii) Changes in specialties or subspecialties under which any licensed laboratory in the State performs testing.

(9) Provide information for the review of the State’s enforcement procedures for laboratories found to be out of compliance with the State’s requirements.

(10) Submit information that demonstrates the ability of the State to provide HCFA with the following:

(i) Electronic data and reports in compatible code with the adverse or corrective actions resulting from PT results that constitute unsuccessful participation in PT programs.

(ii) Other data that HCFA determines are necessary for validation and assessment of the State’s inspection process requirements.

(11) Agree to provide HCFA with written notification of any changes in its licensure/approval and inspection requirements.

(12) Agree to disclose any laboratory’s PT results in accordance with a State’s confidentiality requirements.

(13) Agree to take the appropriate enforcement action against laboratories found by HCFA not to be in compliance with requirements comparable to condition-level requirements and report these enforcement actions to HCFA.

(14) If approved, reapply to HCFA every 2 years to renew its exempt status and to renew its agreement to pay the cost of the HCFA-administered validation program in that State.

§ 493.559 Publication of approval of deeming authority or CLIA exemption.

(a) Notice of deeming authority or exemption. HCFA publishes a notice in the Federal Register when it grants deeming authority to an accreditation organization or exemption to a State licensure program.

(b) Contents of notice. The notice includes the following:

(1) The name of the accreditation organization or State licensure program.

(2) For an accreditation organization:

(i) The specific specialty or subspecialty areas for which it is granted deeming authority.

(ii) A description of how the accreditation organization provides reasonable assurance to HCFA that a laboratory accredited by the organization meets CLIA requirements equivalent to those in this part and would meet CLIA requirements if the laboratory had not been granted deemed status, but had been inspected against condition-level requirements.

(3) For a State licensure program, a description of how the laboratory requirements of the State are equal to, or more stringent than, those specified in this part.

(4) The basis for granting deeming authority or exemption.

(5) The term of approval, not to exceed 6 years.

§ 493.561 Denial of application or reapplication.

(a) Reconsideration of denial. (1) If HCFA denies a request for approval, an accreditation organization or State licensure program may request, within 60 days of the notification of denial, that HCFA reconsider its original application or application for renewal, in accordance with part 488, subpart D.

(2) If the accreditation organization or State licensure program requests a reconsideration of HCFA’s determination to deny its request for approval or reapproval, it may not submit a new application until HCFA issues a final reconsideration determination.

(b) Resubmittal of a request for approval— accreditation organization. An accreditation organization may resubmit a request for approval if a final reconsideration determination is not pending and the accreditation program meets the following conditions:

(1) It has revised its accreditation program to address the rationale for denial of its previous request.

(2) It demonstrates that it can provide reasonable assurance that its accredited facilities meet condition-level requirements.
(3) It resubmits the application in its entirety.

(c) Resubmittal of request for approval—State licensure program. The State licensure program may resubmit a request for approval if a final reconsideration determination is not pending and it has taken the necessary action to address the rationale for any previous denial.

§ 493.563 Validation inspections—Basis and focus.

(a) Basis for validation inspection—(1) Laboratory with a certificate of accreditation. (i) HCFA or a HCFA agent may conduct an inspection of an accredited laboratory that has been issued a certificate of accreditation on a representative sample basis or in response to a substantial allegation of noncompliance.

(ii) HCFA uses the results of these inspections to validate the accreditation organization’s accreditation process.

(2) Laboratory in a State with an approved State licensure program. (i) HCFA or a HCFA agent may conduct an inspection of any laboratory in a State with an approved State licensure program on a representative sample basis or in response to a substantial allegation of noncompliance.

(ii) The results of these inspections are used to validate the appropriateness of the exemption of that State’s licensed or approved laboratories from CLIA program requirements.

(b) Validation inspection conducted on a representative sample basis. (1) If HCFA or a HCFA agent conducts a validation inspection on a representative sample basis, the inspection is comprehensive, addressing all condition-level requirements, or it may be focused on a specific condition-level requirement.

(2) The number of laboratories sampled is sufficient to allow a reasonable estimate of the performance of the accreditation organization or State.

(c) Validation inspection conducted in response to a substantial allegation of noncompliance. (1) If HCFA or a HCFA agent conducts a validation inspection in response to a substantial allegation of noncompliance, the inspection focuses on any condition-level requirement that HCFA determines to be related to the allegation.

(2) If HCFA or a HCFA agent substantiates a deficiency and determines that the laboratory is out of compliance with any condition-level requirement, HCFA or a HCFA agent conducts a full CLIA inspection.

(d) Inspection of operations and offices. As part of the validation review process, HCFA may conduct an onsite inspection of the operations and offices to verify the following:

(1) The accreditation organization’s representations and to assess the accreditation organization’s compliance with its own policies and procedures.

(2) The State’s representations and to assess the State’s compliance with its own policies and procedures, including verification of State enforcement actions taken on the basis of validation inspections performed by HCFA or a HCFA agent.

(e) Onsite inspection of an accreditation organization. An onsite inspection of an accreditation organization may include, but is not limited to, the following:

(1) A review of documents.

(2) An audit of meetings concerning the accreditation process.

(3) Evaluation of accreditation inspection results and the accreditation decision-making process.

(4) Interviews with the accreditation organization’s staff.

(f) Onsite inspection of a State licensure program. An onsite inspection of a State licensure program office may include, but is not limited to, the following:

(1) A review of documents.

(2) An audit of meetings concerning the licensure or approval process.

(3) Evaluation of State inspection results and the licensure or approval decision-making process.

(4) Interviews with State employees.

§ 493.565 Selection for validation inspection—laboratory responsibilities.

A laboratory selected for a validation inspection must do the following:

(a) Authorize its accreditation organization or State licensure program, as applicable, to release to HCFA or a HCFA agent, on a confidential basis, a copy of the laboratory’s most recent
§ 493.567 Refusal to cooperate with validation inspection.

(a) Laboratory with a certificate of accreditation. (1) A laboratory with a certificate of accreditation that refuses to cooperate with a validation inspection by failing to comply with the requirements in § 493.565—
   (i) Is subject to full review by HCFA or a HCFA agent, in accordance with this part; and
   (ii) May be subject to suspension, revocation, or limitation of its certificate of accreditation under this part.

(b) CLIA-exempt laboratory. If a validation inspection results in a finding that a CLIA-exempt laboratory is out of compliance with one or more condition-level requirements, HCFA directs the State to take appropriate enforcement action.

§ 493.569 Consequences of a finding of noncompliance as a result of a validation inspection.

(a) Laboratory with a certificate of accreditation. If a validation inspection results in a finding that the accredited laboratory is out of compliance with one or more condition-level requirements, the laboratory is subject to—
   (1) The same requirements and survey and enforcement processes applied to laboratories that are not accredited and that are found out of compliance following an inspection under this part; and
   (2) Full review by HCFA, in accordance with this part; that is, the laboratory is subject to the principal and alternative sanctions in § 493.1806.

(b) CLIA-exempt laboratory. If a validation inspection results in a finding that a CLIA-exempt laboratory is out of compliance with one or more condition-level requirements, HCFA directs the State to take appropriate enforcement action.

§ 493.571 Disclosure of accreditation, State and HCFA validation inspection results.

(a) Accreditation organization inspection results. HCFA may disclose accreditation organization inspection results to the public only if the results are related to an enforcement action taken by the Secretary.

(b) State inspection results. Disclosure of State inspection results is the responsibility of the approved State licensure program, in accordance with State law.

(c) HCFA validation inspection results. HCFA may disclose the results of all validation inspections conducted by HCFA or its agent.

§ 493.573 Continuing Federal oversight of private nonprofit accreditation organizations and approved State licensure programs.

(a) Comparability review. In addition to the initial review for determining equivalency of specified organization or State requirements to the comparable condition-level requirements, HCFA reviews the equivalency of requirements in the following cases:
   (1) When HCFA promulgates new condition-level requirements.
(2) When HCFA identifies an accreditation organization or a State licensure program whose requirements are no longer equal to, or more stringent than, condition-level requirements.

(3) When an accreditation organization or State licensure program adopts new requirements.

(4) When an accreditation organization or State licensure program adopts changes to its inspection process, as required by §493.575(b)(1), as applicable.

(5) Every 6 years, or sooner if HCFA determines an earlier review is required.

(b) Validation review. Following the end of a validation review period, HCFA evaluates the validation inspection results for each approved accreditation organization and State licensure program.

(c) Reapplication procedures. (1) Every 6 years, or sooner, as determined by HCFA, an approved accreditation organization must reapply for continued approval of deeming authority and a State licensure program must reapply for continued approval of a CLIA exemption. HCFA provides notice of the materials that must be submitted as part of the reapplication procedure.

(2) An accreditation organization or State licensure program that does not meet the requirements of this subpart, as determined through a comparability or validation review, must furnish HCFA, upon request, with the reapplication materials HCFA requests. HCFA establishes a deadline by which the materials must be submitted.

(d) Notice. (1) HCFA provides written notice, as appropriate, to the following:

(ii) An accreditation organization indicating that its approval may be in jeopardy if a comparability or validation review reveals that it is not meeting the requirements of this subpart and HCFA is initiating a review of the accreditation organization’s deeming authority.

(ii) A State licensure program indicating that its CLIA exemption may be in jeopardy if a comparability or validation review reveals that it is not meeting the requirements of this subpart and that a review is being initiated of the CLIA exemption of the State’s laboratories.

(2) The notice contains the following information:

(i) A statement of the discrepancies that were found as well as other related documentation.

(ii) An explanation of HCFA’s review process on which the final determination is based and a description of the possible actions, as specified in §493.575, that HCFA may impose based on the findings from the comparability or validation review.

(iii) A description of the procedures available if the accreditation organization or State licensure program, as applicable, desires an opportunity to explain or justify the findings made during the comparability or validation review.

(iv) The reapplication materials that the accreditation organization or State licensure program must submit and the deadline for that submission.

§ 493.575 Removal of deeming authority or CLIA exemption and final determination review.

(a) HCFA review. HCFA conducts a review of the following:

(1) A deeming authority review of an accreditation organization’s program if the comparability or validation review produces findings, as described at §493.573. HCFA reviews, as appropriate, the criteria described in §§493.555 and 493.557(a) to reevaluate whether the accreditation organization continues to meet all these criteria.

(2) An exemption review of a State’s licensure program if the comparability or validation review produces findings, as described at §493.573. HCFA reviews, as appropriate, the criteria described in §§493.555 and 493.557(b) to reevaluate whether the licensure program continues to meet all these criteria.

(3) A review of an accreditation organization or State licensure program, at HCFA’s discretion, if validation review findings, irrespective of the rate of disparity, indicate widespread or systematic problems in the organization’s accreditation or State’s licensure process that provide evidence that the requirements, taken as a whole, are no longer equivalent to CLIA requirements, taken as a whole.

(4) A review of the accreditation organization or State licensure program
whenever validation inspection results indicate a rate of disparity of 20 percent or more between the findings of the organization or State and those of HCFA or a HCFA agent for the following periods:

(i) One year for accreditation organizations.

(ii) Two years for State licensure programs.

(b) HCFA action after review. Following the review, HCFA may take the following action:

(1) If HCFA determines that the accreditation organization or State has failed to adopt requirements equal to, or more stringent than, CLIA requirements, HCFA may give a conditional approval for a probationary period of its deeming authority to an organization 30 days following the date of HCFA's determination, or exempt status to a State within 30 days of HCFA's determination, both not to exceed 1 year, to afford the organization or State an opportunity to adopt equal or more stringent requirements.

(2) If HCFA determines that there are widespread or systematic problems in the organization's or State's inspection process, HCFA may give conditional approval during a probationary period, not to exceed 1 year, effective 30 days following the date of the determination.

(c) Final determination. HCFA makes a final determination as to whether the organization or State continues to meet the criteria described in this subpart and issues a notice that includes the reasons for the determination to the organization or State within 60 days after the end of any probationary period. This determination is based on an evaluation of any of the following:

(1) The most recent validation inspection and review findings. To continue to be approved, the organization or State must meet the criteria of this subpart.

(2) Facility-specific data, as well as other related information.

(3) The organization's or State's inspection procedures, surveyors' qualifications, ongoing education, training, and composition of inspection teams.

(4) The organization's accreditation requirements, or the State's licensure or approval requirements.

(d) Date of withdrawal of approval. HCFA may withdraw its approval of the accreditation organization or State licensure program, effective 30 days from the date of written notice to the organization or State of this proposed action, if improvements acceptable to HCFA have not been made during the probationary period.

(e) Continuation of validation inspections. The existence of any validation review, probationary status, or any other action, such as a deeming authority review, by HCFA does not affect or limit the conduct of any validation inspection.

(f) Federal Register notice. HCFA publishes a notice in the Federal Register containing a justification for removing the deeming authority from an accreditation organization, or the CLIA-exempt status of a State licensure program.

(g) Withdrawal of approval—effect on laboratory status—(1) Accredited laboratory. After HCFA withdraws approval of an accreditation organization's deeming authority, the certificate of accreditation of each affected laboratory continues in effect for 60 days after it receives notification of the withdrawal of approval.

(2) CLIA-exempt laboratory. After HCFA withdraws approval of a State licensure program, the exempt status of each licensed or approved laboratory in the State continues in effect for 60 days after a laboratory receives notification from the State of the withdrawal of HCFA's approval of the program.

(3) Extension. After HCFA withdraws approval of an accreditation organization or State licensure program, HCFA may extend the period for an additional 60 days for a laboratory if it determines that the laboratory submitted an application for accreditation to an approved accreditation organization or an application for the appropriate certificate to HCFA or a HCFA agent before the initial 60-day period ends.

(h) Immediate jeopardy to patients. (1) If at any time HCFA determines that the continued approval of a deeming authority of any accreditation organization poses immediate jeopardy to the patients of the laboratories accredited
§ 493.638 Certificate fees.

(a) Basic rule. Laboratories must pay a fee for the issuance of a registration certificate, certificate for PPM procedures, certificate of waiver, certificate of accreditation, or a certificate of compliance, as applicable. Laboratories must also pay a fee to reapply for a certificate for PPM procedures, certificate of waiver, certificate of accreditation, or a certificate of compliance. The total of fees collected by HHS under the laboratory program must be sufficient to cover the general costs of administering the laboratory certification program under section 353 of the PHS Act.

(1) For registration certificates and certificates of compliance, the costs include issuing the certificates, collecting the fees, evaluating and monitoring proficiency testing programs, evaluating which procedures, tests or examinations meet the criteria for inclusion in the appropriate complexity category, and implementing section 353 of the PHS Act.

(2) For a certificate of waiver, the costs include issuing the certificate, collecting the fees, determining if a certificate of waiver should be issued, evaluating which tests qualify for inclusion in the waived category, and other direct administrative costs.

(3) For a certificate for PPM procedures, the costs include issuing the certificate, collecting the fees, determining if a certificate for PPM procedures should be issued, evaluating which procedures meet the criteria for inclusion in the subcategory of PPM procedures, and other direct administrative costs.

(4) For a certificate of accreditation, the costs include issuing the certificate, collecting the fees, evaluating the programs of accrediting bodies, and other direct administrative costs.

SOURCE: 57 FR 7138 and 7213, Feb. 28, 1992, unless otherwise noted.

§ 493.602 Scope of subpart.

This subpart sets forth the methodology for determining the amount of the fees for issuing the appropriate certificate, and for determining compliance with the applicable standards of the Public Health Service Act (the PHS Act) and the Federal validation of accredited laboratories and of CLIA-exempt laboratories.

[60 FR 20047, Apr. 24, 1995]

§ 493.606 Applicability of subpart.

The rules of this subpart are applicable to those laboratories specified in §493.3.

[58 FR 5212, Jan. 19, 1993]
§ 493.639 Fee amount.

(b) Fee amount. The fee amount is set annually by HHS on a calendar year basis and is based on the category of test complexity, or on the category of test complexity and schedules or ranges of annual laboratory test volume (excluding waived tests and tests performed for quality control, quality assurance, and proficiency testing purposes) and specialties tested, with the amounts of the fees in each schedule being a function of the costs for all aspects of general administration of CLIA as set forth in § 493.649 (b) and (c). This fee is assessed and payable at least biennially. The methodology used to determine the amount of the fee is found in § 493.649. The amount of the fee applicable to the issuance of the registration certificate or the issuance or renewal of the certificate for PPM procedures, certificate of waiver, certificate of accreditation, or certificate of compliance is the amount in effect at the time the application is received. Upon receipt of an application for a certificate, HHS or its designee notifies the laboratory of the amount of the required fee for the requested certificate.

§ 493.643 Fee for determination of program compliance.

(a) Fee requirement. In addition to the fee required under § 493.638, a laboratory subject to routine inspections must pay a fee to cover the cost of determining program compliance. Laboratories issued a certificate for PPM procedures, certificate of waiver, or a certificate of accreditation are not subject to this fee for routine inspections.

(b) Costs included in the fee. Included in the fee for determining program compliance is the cost of evaluating qualifications of personnel; monitoring proficiency testing; conducting onsite inspections; documenting deficiencies; evaluating laboratories’ plans to correct deficiencies; and necessary administrative costs. HHS sets the fee amounts annually on a calendar year basis. Laboratories are inspected biennially; therefore, fees are assessed and payable biennially. If additional expenses are incurred to conduct follow up visits to verify correction of deficiencies, to impose sanctions, and/or for surveyor preparation for and attendance at ALJ hearings, HHS assesses an additional fee to include these costs. The additional fee is based on the actual resources and time necessary to perform the activities.

(c) Classification of laboratories that require inspection for purpose of determining amount of fee. (1) There are ten classifications (schedules) of laboratories for the purpose of determining the fee amount a laboratory is assessed. Each laboratory is placed into...
one of the ten following schedules based on the laboratory's scope and volume of testing (excluding tests performed for quality control, quality assurance, and proficiency testing purposes).

(i) (A) Schedule A Low Volume. The laboratory performs not more than 2,000 laboratory tests annually.
(B) Schedule A. The laboratory performs tests in no more than 3 specialties of service with a total annual volume of more than 2,000 but not more than 10,000 laboratory tests.

(ii) Schedule B. The laboratory performs tests in at least 4 specialties of service with a total annual volume of not more than 10,000 laboratory tests.

(iii) Schedule C. The laboratory performs tests in no more 3 specialties of service with a total annual volume of more than 10,000 but not more than 25,000 laboratory tests.

(iv) Schedule D. The laboratory performs tests in at least 4 specialties with a total annual volume of more than 10,000 but not more than 25,000 laboratory tests.

(v) Schedule E. The laboratory performs more than 25,000 but not more than 50,000 laboratory tests annually.

(vi) Schedule F. The laboratory performs more than 50,000 but not more than 75,000 laboratory tests annually.

(vii) Schedule G. The laboratory performs more than 75,000 but not more than 100,000 laboratory tests annually.

(viii) Schedule H. The laboratory performs more than 100,000 but not more than 500,000 laboratory tests annually.

(ix) Schedule I. The laboratory performs more than 500,000 but not more than 1,000,000 laboratory tests annually.

(x) Schedule J. The laboratory performs more than 1,000,000 laboratory tests annually.

(2) For purposes of determining a laboratory's classification under this section, a test is a procedure or examination for a single analyte. (Tests performed for quality control, quality assurance, and proficiency testing are excluded from the laboratory's total annual volume). Each profile (that is, group of tests) is counted as the number of separate procedures or examinations; for example, a chemistry profile consisting of 18 tests is counted as 18 separate procedures or tests.

(3) For purposes of determining a laboratory's classification under this section, the specialties and subspecialties of service for inclusion are:

(i) The specialty of Microbiology, which includes one or more of the following subspecialties:
(A) Bacteriology.
(B) Mycobacteriology.
(C) Mycology.
(D) Parasitology.
(E) Virology.

(ii) The specialty of Serology, which includes one or more of the following subspecialties:
(A) Syphilis Serology.
(B) General immunology.

(iii) The specialty of Chemistry, which includes one or more of the following subspecialties:
(A) Routine chemistry.
(B) Endocrinology.
(C) Toxicology.
(D) Urinalysis.

(iv) The specialty of Hematology.

(v) The specialty of Immunohematology, which includes one or more of the following subspecialties:
(A) ABO grouping and Rh typing.
(B) Unexpected antibody detection.
(C) Compatibility testing.
(D) Unexpected antibody identification.

(vi) The specialty of Pathology, which includes the following subspecialties:
(A) Cytology.
(B) Histopathology.
(C) Oral pathology.

(vii) The specialty of Radiobiology.

(viii) The specialty of Histocompatibility.

(ix) The specialty of Cytogenetics.

(d) Additional fees. (1) If after a certificate of compliance is issued, a laboratory adds services and requests that its certificate be upgraded, the laboratory must pay an additional fee if, in order to determine compliance with additional requirements, it is necessary to conduct an inspection, evaluate personnel, or monitor proficiency testing performance. The additional fee is based on the actual resources and time necessary to perform the activities.
§ 493.645 Payment of fees.

(a) Except for CLIA-exempt laboratories, all laboratories are notified in writing by HHS or its designee of the appropriate fee(s) and instructions for submitting the fee(s), including the due date for payment and where to make payment. The appropriate certificate is not issued until the applicable fees have been paid.

(b) For State-exempt laboratories, HHS estimates the cost of conducting validation surveys within the State for a 2-year period. HHS or its designee notifies the State by mail of the appropriate fees, including the due date for payment and the address of the United States Department of Treasury designated commercial bank to which payment must be made. In addition, if complaint investigations are conducted.
§ 493.649 Methodology for determining fee amount.

(a) General rule. The amount of the fee in each schedule for compliance determination inspections is based on the average hourly rate (which includes the costs to perform the required activities and necessary administration costs) multiplied by the average number of hours required or, if activities are performed by more than one of the entities listed in paragraph (b) of this section, the sum of the products of the applicable hourly rates multiplied by the average number of hours required by the entity to perform the activity. The fee for issuance of the registration certificate or certificate of compliance is based on the laboratory’s scope and volume of testing.

(b) Determining average hourly rates used in fee schedules. Three different entities perform activities related to the issuance or reissuance of any certificate. HHS determines the average hourly rates for the activities of each of these entities.

(1) State survey agencies. The following costs are included in determining an average hourly rate for the activities performed by State survey agencies:

(i) The costs incurred by the State survey agencies in evaluating personnel qualifications and monitoring each laboratory’s participation in an approved proficiency testing program. The cost of on-site inspections and monitoring activities is the hourly rate derived as a result of an annual budget negotiation process with each State. The hourly rate encompasses salary costs (as determined by each State’s civil service pay scale) and fringe benefit costs to support the required number of State inspectors, management and direct support staff.

(ii) Travel costs necessary to comply with each State’s administrative requirements and other direct costs such as equipment, printing, and supplies. These costs are established based on historical State requirements.

(iii) Indirect costs as negotiated by HHS.

(2) Federal agencies. The hourly rate for activities performed by Federal agencies is the most recent average hourly cost to HHS to staff and support a full time equivalent employee. Included in this cost are salary and fringe benefit costs, necessary administrative costs, such as printing, training, postage, express mail, supplies, equipment, computer system and building service charges associated with support services provided by organizational components such as a computer center, and any other oversight activities necessary to support the program.

(3) HHS contractors. The hourly rate for activities performed by HHS contractors is the average hourly rate established for contractor assistance based on an independent government cost estimate for the required workload. This rate includes the cost of contractor support to provide proficiency testing programs to laboratories that do not participate in an approved proficiency testing program, provide specialized assistance in the evaluation of laboratory performance in an approved proficiency testing program, perform assessments of cytology testing laboratories, conduct special studies, bill and collect fees, issue certificates, establish accounting, monitoring and reporting systems, and assist with necessary surveyor training.

(c) Determining number of hours. The average number of hours used to determine the overall fee in each schedule is HHS’s estimate, based on historical experience, of the average time needed by each entity to perform the activities for which it is responsible.

[57 FR 7138 and 7213, Feb. 28, 1992, as amended at 60 FR 20048, Apr. 24, 1995]
§ 493.801

Subpart H—Participation in Proficiency Testing for Laboratories Performing Tests of Moderate Complexity (Including the Subcategory), High Complexity, or Any Combination of These Tests

SOURCE: 57 FR 7146, Feb. 28, 1992, unless otherwise noted.

§ 493.801 Condition: Enrollment and testing of samples.

Each laboratory must enroll in a proficiency testing (PT) program that meets the criteria in subpart I of this part and is approved by HHS. The laboratory must enroll in an approved program or programs for each of the specialties and subspecialties for which it seeks certification. The laboratory must test the samples in the same manner as patients' specimens. For laboratories subject to 42 CFR part 493 published on March 14, 1990 (55 FR 9538) prior to September 1, 1992, the rules of this subpart are effective on September 1, 1992. For all other laboratories, the rules of this subpart are effective January 1, 1994.

(a) Standard; Enrollment. The laboratory must—

(1) Notify HHS of the approved program or programs in which it chooses to participate to meet proficiency testing requirements of this subpart.

(2)(i) Designate the program(s) to be used for each specialty, subspecialty, and analyte or test to determine compliance with this subpart if the laboratory participates in more than one proficiency testing program approved by HCFA; and

(ii) For those tests performed by the laboratory that are not included in subpart I of this part, a laboratory must establish and maintain the accuracy of its testing procedures, in accordance with § 493.1709.

(b) Standard; Testing of proficiency testing samples. The laboratory must examine or test, as applicable, the proficiency testing samples it receives from the proficiency testing program in the same manner as it tests patient specimens.

(1) The samples must be examined or tested with the laboratory's regular patient workload by personnel who routinely perform the testing in the laboratory, using the laboratory's routine methods. The individual testing or examining the samples and the laboratory director must attest to the routine integration of the samples into the patient workload using the laboratory's routine methods.

(2) The laboratory must test samples the same number of times that it routinely tests patient samples.

(3) Laboratories that perform tests on proficiency testing samples must not engage in any inter-laboratory communications pertaining to the results of proficiency testing sample(s) until after the date by which the laboratory must report proficiency testing results to the program for the testing event in which the samples were sent. Laboratories with multiple testing sites or separate locations must not participate in any communications or discussions across sites/locations concerning proficiency testing sample results until after the date by which the laboratory must report proficiency testing results to the program.

(4) The laboratory must not send PT samples or portions of samples to another laboratory for any analysis which it is certified to perform in its own laboratory. Any laboratory that HCFA determines intentionally referred its proficiency testing samples to another laboratory for analysis will have its certification revoked for at least one year. Any laboratory that receives proficiency testing samples from another laboratory for testing must...
§ 493.807 Condition: Reinstatement of laboratories performing tests of moderate complexity (including the subcategory), high complexity, or any combination of these tests, after failure to participate successfully.

(a) If a laboratory’s certificate is suspended or limited or its Medicare or Medicaid approval is cancelled or its Medicare or Medicaid payments are suspended because it fails to participate successfully in proficiency testing for one or more specialties, subspecialties, analyte or test, or voluntarily withdraws its certification under CLIA for the failed specialty, subspecialty, or analyte, the laboratory must then demonstrate sustained satisfactory performance on two consecutive proficiency testing events, one of which may be on site, before HCFA will consider it for reinstatement for certification and Medicare or Medicaid approval in that specialty, subspecialty, analyte or test.

(b) The cancellation period for Medicare and Medicaid approval or period for suspension of Medicare or Medicaid payments or suspension or limitation of certification under CLIA for the failed specialty, subspecialty, or analyte or test is for a period of not less than six months from the date of cancellation, limitation or suspension of the CLIA certificate.

[58 FR 5228, Jan. 19, 1993, as amended at 60 FR 20048, Apr. 24, 1995]
§ 493.821 Condition: Microbiology.

The specialty of microbiology includes, for purposes of proficiency testing, the subspecialties of bacteriology, mycobacteriology, mycology, parasitology and virology.

§ 493.823 Standard; Bacteriology.

(a) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.

(b) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

(3) The laboratory participated in the previous two proficiency testing events.

(c) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(d)(1) For any unsatisfactory testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

(2) Remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(3) The laboratory participated in the previous two proficiency testing events.

(e) Failure to achieve an overall testing event score of satisfactory performance for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.

§ 493.825 Standard; Mycobacteriology.

(a) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.

(b) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

(3) The laboratory participated in the previous two proficiency testing events.

(d)(1) For any unsatisfactory testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

(2) Remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(e) Failure to achieve an overall testing event score of satisfactory performance for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.
§ 493.827 Standard; Mycology.

(a) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.

(b) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

1. Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

2. The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

3. The laboratory participated in the previous two proficiency testing events.

(c) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(d)(1) For any unsatisfactory testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

2. Remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(e) Failure to achieve an overall testing event score of satisfactory performance for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.

§ 493.831 Standard; Virology.

(a) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.

(b) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

1. Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

2. The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

3. The laboratory participated in the previous two proficiency testing events.

(c) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(d)(1) For any unsatisfactory testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

2. Remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(e) Failure to achieve an overall testing event score of satisfactory performance for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.

§ 493.829 Standard; Parasitology.

(a) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.

(b) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

1. Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

2. The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

3. The laboratory participated in the previous two proficiency testing events.

(c) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(d)(1) For any unsatisfactory testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

2. Remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(e) Failure to achieve an overall testing event score of satisfactory performance for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.
§ 493.833 Condition: Diagnostic immunology.

The specialty of diagnostic immunology includes for purposes of proficiency testing the subspecialties of syphilis serology and general immunology.

§ 493.835 Standard: Syphilis serology.

(a) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.

(b) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

(3) The laboratory participated in the previous two proficiency testing events.

(c) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(d)(1) For any unsatisfactory testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

(2) For any unsatisfactory testing events, remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(e) Failure to achieve an overall testing event score of satisfactory performance for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.

§ 493.837 Standard: General immunology.

(a) Failure to attain a score of at least 80 percent of acceptable responses for each analyte in each testing event is unsatisfactory analyte performance for the testing event.

(b) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.

(c) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

(3) The laboratory participated in the previous two proficiency testing events.

(c) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(d)(1) For any unsatisfactory testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

(2) For any unacceptable testing event score, remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(e) Failure to achieve an overall testing event score of satisfactory performance for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.
§ 493.843 Standard; Endocrinology.

(a) Failure to attain a score of at least 80 percent of acceptable responses for each analyte in each testing event is unsatisfactory analyte performance for the testing event.

(b) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.

(c) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

(3) The laboratory participated in the previous two proficiency testing events.

(d) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(e)(1) For any unsatisfactory analyte or test performance or testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

(2) For any unacceptable analyte or testing event score, remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(f) Failure to achieve satisfactory performance for the same analyte or test in two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.

(g) Failure to achieve an overall testing event score of satisfactory performance for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.

§ 493.839 Condition: Chemistry.

The specialty of chemistry includes for the purposes of proficiency testing the subspecialties of routine chemistry, endocrinology, and toxicology.

§ 493.841 Standard; Routine chemistry.

(a) Failure to attain a score of at least 80 percent of acceptable responses for each analyte in each testing event is unsatisfactory analyte performance for the testing event.

(b) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.

(c) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

(3) The laboratory participated in the previous two proficiency testing events.

(d) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(e)(1) For any unsatisfactory analyte or test performance or testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

(2) For any unacceptable analyte or testing event score, remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(f) Failure to achieve satisfactory performance for the same analyte or test in two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.

(g) Failure to achieve an overall testing event score of satisfactory performance for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.
(c) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—
(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;
(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and
(3) The laboratory participated in the previous two proficiency testing events.
(d) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.
(e)(1) For any unsatisfactory analyte or test performance or testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.
(2) For any unacceptable analyte or testing event score, remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.
(f) Failure to achieve satisfactory performance for the same analyte or test in two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.
(g) Failure to achieve an overall testing event score of satisfactory performance for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.
§ 493.845 Standard; Toxicology.
(a) Failure to attain a score of at least 80 percent of acceptable responses for each analyte in each testing event is unsatisfactory analyte performance for the testing event.
(b) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.
(c) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—
(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;
(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and
(3) The laboratory participated in the previous two proficiency testing events.
(d) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.
(e)(1) For any unsatisfactory analyte or test performance or testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.
(2) For any unacceptable analyte or testing event score, remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.
(f) Failure to achieve satisfactory performance for the same analyte or test in two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.
(g) Failure to achieve an overall testing event score of satisfactory performance for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.
§ 493.849 Condition: Hematology.
The specialty of hematology, for the purpose of proficiency testing, is not
subdivided into subspecialties of testing.

§ 493.851 Standard; Hematology.

(a) Failure to attain a score of at least 80 percent of acceptable responses for each analyte in each testing event is unsatisfactory analytic performance for the testing event.

(b) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.

(c) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

(3) The laboratory participated in the previous two proficiency testing events.

(d) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(e)(1) For any unsatisfactory analytic or test performance or testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

(2) For any unacceptable analytic or testing event score, remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(f) Failure to achieve satisfactory performance for the same analytic in two consecutive events or two out of three consecutive testing events is unsuccessful performance.

(g) Failure to achieve an overall testing event score of satisfactory performance for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.

§ 493.853 Condition; Pathology.

The specialty of pathology includes, for purposes of proficiency testing, the subspecialty of cytology limited to gynecologic examinations.

§ 493.855 Standard; Cytology; gynecologic examinations.

To participate successfully in a cytology proficiency testing program for gynecologic examinations (Pap smears), the laboratory must meet the requirements of paragraphs (a) through (c) of this section.

(a) The laboratory must ensure that each individual engaged in the examination of gynecologic preparations is enrolled in a proficiency testing program approved by HCFA by January 1, 1995, if available in the State in which he or she is employed. The laboratory must ensure that each individual is tested at least once per year and obtains a passing score. To ensure this annual testing of individuals, an announced or unannounced testing event will be conducted on-site in each laboratory at least once each year. Laboratories will be notified of the time of each announced on-site testing event at least 30 days prior to each event. Additional testing events will be conducted as necessary in each State or region for the purpose of testing individuals who miss the on-site testing event and for retesting individuals as described in paragraph (b) of this section.

(b) The laboratory must ensure that each individual participates in an annual testing event that involves the examination of a 10-slide test set as described in §493.845. Individuals who fail this testing event are retested with another 10-slide test set as described in paragraphs (b)(1) and (b)(2) of this section. Individuals who fail this second test are subsequently retested with a 20-slide test set as described in paragraphs (b)(2) and (b)(3) of this section. Individuals are given not more than 2 hours to complete a 10-slide test and not more than 4 hours to complete a 20-
slide test. Unexcused failure to appear by an individual for a retest will result in test failure with resulting remediation and limitations on slide examinations as specified in (b)(1), (b)(2), and (b)(3) of this section.

1. An individual is determined to have failed the annual testing event if he or she scores less than 90 percent on a 10-slide test set. For an individual who fails an annual proficiency testing event, the laboratory must schedule a retesting event which must take place not more than 45 days after receipt of the notification of failure.

2. An individual is determined to have failed the second testing event if he or she scores less than 90 percent on a 10-slide test set. For an individual who fails a second testing event, the laboratory must provide him or her with documented, remedial training and education in the area of failure, and must assure that all gynecologic slides evaluated subsequent to the notice of failure are reexamined until the individual is again retested with a 20-slide test set and scores at least 90 percent. Reexamination of slides must be documented.

3. An individual is determined to have failed the third testing event if he or she scores less than 90 percent on a 20-slide test set. An individual who fails the third testing event must cease examining gynecologic preparations, and until he or she is retested with a 20-slide test set and scores at least 90 percent.

(c) If a laboratory fails to ensure that individuals are tested or those who fail a testing event are retested, or fails to take required remedial actions as described in paragraphs (b)(1), (b)(2) or (b)(3) of this section, HCFA will initiate intermediate sanctions or limit the laboratory’s certificate to exclude gynecologic cytology testing under CLIA, and, if applicable, suspend the laboratory’s Medicare and Medicaid payments for gynecologic cytology testing in accordance with subpart R of this part.

§ 493.855 Condition: Immunohematology.

The specialty of immunohematology includes four subspecialties for the purposes of proficiency testing: ABO group and D (Rho) typing; unexpected antibody detection; compatibility testing; and antibody identification.

§ 493.859 Standard; ABO group and D (Rho) typing.

(a) Failure to attain a score of at least 100 percent of acceptable responses for each analyte or test in each testing event is unsatisfactory analyte performance for the testing event.

(b) Failure to attain an overall testing event score of at least 100 percent is unsatisfactory performance.

(c) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

(3) The laboratory participated in the previous two proficiency testing events.

(d) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(e)(1) For any unsatisfactory testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.
(2) For any unacceptable analyte or unsatisfactory testing event score, remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(f) Failure to achieve satisfactory performance for the same analyte in two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.

(g) Failure to achieve an overall testing event score of satisfactory for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.

§ 493.861 Standard; Unexpected antibody detection.

(a) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.

(b) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

(3) The laboratory participated in the previous two proficiency testing events.

(c) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(d)(1) For any unsatisfactory testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

(2) For any unsatisfactory testing event score, remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(e) Failure to achieve an overall testing event score of satisfactory for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.

§ 493.863 Standard; Compatibility testing.

(a) Failure to attain an overall testing event score of at least 100 percent is unsatisfactory performance.

(b) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

(3) The laboratory participated in the previous two proficiency testing events.

(c) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(d)(1) For any unsatisfactory testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

(2) For any unsatisfactory testing event score, remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(e) Failure to achieve an overall testing event score of satisfactory for two
§ 493.865 Standard; Antibody identification.

(a) Failure to attain an overall testing event score of at least 80 percent is unsatisfactory performance.

(b) Failure to participate in a testing event is unsatisfactory performance and results in a score of 0 for the testing event. Consideration may be given to those laboratories failing to participate in a testing event only if—

(1) Patient testing was suspended during the time frame allotted for testing and reporting proficiency testing results;

(2) The laboratory notifies the inspecting agency and the proficiency testing program within the time frame for submitting proficiency testing results of the suspension of patient testing and the circumstances associated with failure to perform tests on proficiency testing samples; and

(3) The laboratory participated in the previous two proficiency testing events.

(c) Failure to return proficiency testing results to the proficiency testing program within the time frame specified by the program is unsatisfactory performance and results in a score of 0 for the testing event.

(d)(1) For any unsatisfactory testing event for reasons other than a failure to participate, the laboratory must undertake appropriate training and employ the technical assistance necessary to correct problems associated with a proficiency testing failure.

(2) For any unsatisfactory testing event score, remedial action must be taken and documented, and the documentation must be maintained by the laboratory for two years from the date of participation in the proficiency testing event.

(e) Failure to identify the same antibody in two consecutive or two out of three consecutive testing events is unsuccessful performance.

(f) Failure to achieve an overall testing event score of satisfactory for two consecutive testing events or two out of three consecutive testing events is unsuccessful performance.

Subpart I—Proficiency Testing Programs for Tests of Moderate Complexity (Including the Subcategory), High Complexity, or Any Combination of These Tests

SOURCE: 57 FR 7151, Feb. 28, 1992, unless otherwise noted.

§ 493.901 Approval of proficiency testing programs.

In order for a proficiency testing program to receive HHS approval, the program must be offered by a private nonprofit organization or a Federal or State agency, or entity acting as a designated agent for the State. An organization, Federal, or State program seeking approval or reapproval for its program for the next calendar year must submit an application providing the required information by July 1 of the current year. The organization, Federal, or State program must provide technical assistance to laboratories seeking to qualify under the program, and must, for each specialty, subspecialty, and analyte or test for which it provides testing—

(a) Assure the quality of test samples, appropriately evaluate and score the testing results, and identify performance problems in a timely manner;

(b) Demonstrate to HHS that it has—

(i) The technical ability required to—

(1) Prepare or purchase samples from manufacturers who prepare the samples in conformity with the appropriate good manufacturing practices required in 21 CFR parts 606, 640, and 820; and

(ii) Distribute the samples, using rigorous quality control to assure that samples mimic actual patient specimens when possible and that samples are homogeneous, except for specific subspecialties such as cytology, and will be stable within the time frame for analysis by proficiency testing participants;

(2) A scientifically defensible process for determining the correct result for each challenge offered by the program;
§ 493.903 Administrative responsibilities.

The proficiency testing program must—

(a) (1) Provide HHS or its designees and participating laboratories with an electronic or a hard copy, or both, of reports of proficiency testing results and all scores for each laboratory's performance in a format as required by and approved by HCFA for each CLIA-certified specialty, subspecialty, and analyte or test within 60 days after the date by which the laboratory must report proficiency testing results to the proficiency testing program.

(2) Provide HHS with reports of PT results and scores of individual performance in cytology and provide copies of reports to participating individuals, and to all laboratories that employ the individuals, within 15 working days of the testing event;

(b) Furnish to HHS cumulative reports on an individual laboratory's performance and aggregate data on CLIA-certified laboratories for the purpose of establishing a system to make the proficiency testing program's results available, on a reasonable basis, upon request of any person, and include such explanatory information as may be appropriate to assist in the interpretation of the proficiency testing program's results;

(c) Provide HHS with additional information and data upon request and submit such information necessary for HHS to conduct an annual evaluation to determine whether the proficiency testing program continues to meet the requirements of §§ 493.901 through 493.959;

(d) Maintain records of laboratories' performance for a period of five years or such time as may be necessary for any legal proceedings; and

(e) Provide HHS with an annual report and, if needed, an interim report which identifies any previously unrecognized sources of variability in kits, instruments, methods, or PT samples, which adversely affect the programs' ability to evaluate laboratory performance.
§ 493.905 Nonapproved proficiency testing programs.

If a proficiency testing program is determined by HHS to fail to meet any criteria contained in §§ 493.901 through 493.959 for approval of the proficiency testing program, HCFA will notify the program and the program must notify all laboratories enrolled of the non-approval and the reasons for non-approval within 30 days of the notification.

Proficiency Testing Programs by Specialty and Subspecialty

§ 493.909 Microbiology.

(a) Types of services offered by laboratories. In bacteriology, for proficiency testing purposes, there are five types of laboratories:

(1) Those that interpret Gram stains or perform primary inoculation, or both; and refer cultures to another laboratory appropriately certified for the subspecialty of bacteriology for identification;

(2) Those that use direct antigen techniques to detect an organism and may also interpret Gram stains or perform primary inoculation, or perform any combination of these;

(3) Those that, in addition to interpreting Gram stains, performing primary inoculations, and using direct antigen tests, also isolate and identify aerobic bacteria from throat, urine, cervical, or urethral discharge specimens to the genus level and may also perform antimicrobial susceptibility tests on selected isolated microorganisms;

(4) Those that perform the services in paragraph (a)(3) of this section and also isolate and identify anaerobic bacteria from any source.

(b) Program content and frequency of challenge. To be approved for proficiency testing for bacteriology, the annual program must provide a minimum of five samples per testing event. There must be at least three testing events at approximately equal intervals per year. The samples may be provided to the laboratory through mailed shipments or, at HHS’ option, may be provided to HHS or its designee for on-site testing. For the types of laboratories specified in paragraph (a) of this section, an annual program must include samples that contain organisms that are representative of the six major groups of bacteria: anaerobes, Enterobacteriaceae, gram-positive bacilli, gram-positive cocci, gram-negative cocci, and miscellaneous gram-negative bacteria, as appropriate. The specific organisms included in the samples may vary from year to year. The annual program must include samples for bacterial antigen detection, bacterial isolation and identification, Gram stain, and antimicrobial susceptibility testing.

(1) An approved program must furnish HHS with a description of samples that it plans to include in its annual program no later than six months before each calendar year. At least 50 percent of the samples must be mixtures of the principal organism and appropriate normal flora. The program must include other important emerging pathogens (as determined by HHS) and either organisms commonly occurring in patient specimens or opportunistic pathogens. The program must include the following two types of samples; each type of sample must meet the 50 percent mixed culture criterion:

(i) Samples that require laboratories to report only organisms that the testing laboratory considers to be a principal pathogen that is clearly responsible for a described illness (excluding immuno-compromised patients). The program determines the reportable isolates, including antimicrobial susceptibility for any designated isolate; and

(ii) Samples that require laboratories to report all organisms present. Samples must contain multiple organisms frequently found in specimens such as...
urine, blood, abscesses, and aspirates where multiple isolates are clearly significant or where specimens are derived from immuno-compromised patients. The program determines the reportable isolates.

(2) An approved program may vary over time. For example, the types of organisms that might be included in an approved program over time are—

**Anaerobes:**
- Bacteroides fragilis group
- Clostridium perfringens
- Peptostreptococcus anaerobius
- Enterobacteriaceae
- Citrobacter freundii
- Enterobacter aerogenes
- Escherichia coli
- Klebsiella pneumoniae
- Proteus mirabilis
- Salmonella typhimurium
- Serratia marcescens
- Shigella sonnei
- Yersinia enterocolitica

**Gram-positive bacilli:**
- Listeria monocytogenes
- Corynebacterium species CDC Group JK

**Gram-positive cocci:**
- Staphylococcus aureus
- Streptococcus Group A
- Streptococcus Group B
- Streptococcus Group D (S. bovis and enterococcus)
- Streptococcus pneumoniae

**Gram-negative cocci:**
- Branhamella catarrhalis
- Neisseria gonorrhoeae
- Neisseria meningitidis

**Miscellaneous Gram-negative bacteria:**
- Campylobacter jejuni
- Haemophilus influenzae, Type B
- Pseudomonas aeruginosa

(3) For antimicrobial susceptibility testing, the program must provide at least one sample per testing event that includes gram-positive or gram-negative strains that have a predetermined pattern of sensitivity or resistance to the common antimicrobial agents.

(c) Evaluation of a laboratory's performance. HHS approves only those programs that assess the accuracy of a laboratory's responses in accordance with paragraphs (c)(1) through (7) of this section.

(1) The program evaluates characteristics to be interpreted by Gram stain. The program determines the reportable bacteria to be detected by direct antigen techniques or isolation. To determine the accuracy of a laboratory's response for Gram stain interpretation, direct antigen detection, identification, or antimicrobial susceptibility testing, the program must compare the laboratory's response for each sample with the response which reflects agreement of either 90 percent of ten or more referee laboratories or 90 percent or more of all participating laboratories.

(2) To evaluate a laboratory's response for a particular sample, the program must determine a laboratory's type of service in accordance with paragraph (a) of this section. A laboratory must isolate and identify the organisms to the same extent it performs these procedures on patient specimens. A laboratory's performance will be evaluated on the basis of its final answer, for example, a laboratory specified in paragraph (a)(3) of this section will be evaluated on the basis of the average of its scores for paragraphs (c)(3) through (c)(6) as determined in paragraph (c)(7) of this section.

(3) Since laboratories may incorrectly report the presence of organisms in addition to the correctly identified principal organism(s), the grading system must provide a means of deducting credit for additional erroneous organisms that are reported. Therefore, the total number of correct responses for organism isolation and identification submitted by the laboratory divided by the number of organisms present plus the number of incorrect organisms reported by the laboratory must be multiplied by 100 to establish a score for each sample in each testing event. For example, if a sample contained one principal organism and the laboratory reported it correctly but reported the presence of an additional organism, which was not considered reportable, the sample grade would be 1/(1+1)×100=50 percent.

(4) For antimicrobial susceptibility testing, a laboratory must indicate which drugs are routinely included in its test panel when testing patient samples. A laboratory's performance will be evaluated for only those antibiotics for which service is offered. A correct response for each antibiotic will be determined as described in §§493.911(c) (1) using criteria such as
§ 493.913 Mycobacteriology.

(a) Types of services offered by laboratories. In mycobacteriology, there are five types of laboratories for proficiency testing purposes:

(1) Those that interpret acid-fast stains and refer specimen to another laboratory appropriately certified in the subspecialty of mycobacteriology;

(2) Those that interpret acid-fast stains, perform primary inoculation, and refer cultures to another laboratory appropriately certified in the subspecialty of mycobacteriology;

(3) Those that interpret acid-fast stains, isolate and perform identification and/or antimycobacterial susceptibility of Mycobacterium tuberculosis, but refer other mycobacteria species to another laboratory appropriately certified in the subspecialty of mycobacteriology for identification and/or susceptibility tests;

(4) Those that interpret acid-fast stains, isolate, and identify all mycobacteria to the extent required for correct clinical diagnosis, but refer antimycobacterial susceptibility tests to another laboratory appropriately certified in the subspecialty of mycobacteriology;

(5) Those that interpret acid-fast stains, isolate and identify all mycobacteria to the extent required for correct clinical diagnosis, and perform antimycobacterial susceptibility tests on the organisms isolated.

(b) Program content and frequency of challenge. To be approved for proficiency testing for mycobacteriology, the annual program must provide a minimum of five samples per testing event. There must be at least two testing events per year. The samples may be provided through mailed shipments or, at HHS’ option, provided to HHS or its designee for on-site testing events. For types of laboratories specified in paragraphs (a)(1) and (a)(3) through (5) of this section, an annual program must include samples that contain species that are representative of the 5 major (complexes) of mycobacteria encountered in human specimens. The specific mycobacteria included in the samples may vary from year to year.

(1) An approved program must furnish HHS and its agents with a description of samples that it plans to include in its annual program no later than six months before each calendar year. At least 50 percent of the samples must be mixtures of the principal mycobacteria and appropriate normal flora. The program must include mycobacteria commonly occurring in patient specimens and other important emerging mycobacteria (as determined by HHS). The program determines the reportable isolates and correct responses for antimycobacterial susceptibility for any designated isolate.

(2) An approved program may vary over time. For example, the types of
mycobacteria that might be included in an approved program over time are—

TB
  Mycobacterium tuberculosis
  Mycobacterium bovis

Group I
  Mycobacterium kansasii

Group II
  Mycobacterium szulgai

Group III
  Mycobacterium avium-intracellulare
  Mycobacterium terrae

Group IV
  Mycobacterium fortuitum

(3) For antimycobacterial susceptibility testing, the program must provide at least one sample per testing event that includes mycobacterium tuberculosis that has a predetermined pattern of sensitivity or resistance to the common antimycobacterial agents.

(4) For laboratories specified in paragraphs (a)(1) and (a)(2), the program must provide at least five samples per testing event that includes challenges that are acid-fast and challenges which do not contain acid-fast organisms.

(c) Evaluation of a laboratory’s performance. HHS approves only those programs that assess the accuracy of a laboratory’s response in accordance with paragraphs (c)(1) through (6) of this section.

(1) The program determines the reportable mycobacteria to be detected by acid-fast stain, for isolation and identification, and for antimycobacterial susceptibility. To determine the accuracy of a laboratory’s response, the program must compare the laboratory’s response for each sample with the response that reflects agreement of either 90 percent of ten or more referee laboratories or 90 percent or more of all participating laboratories.

(2) To evaluate a laboratory’s response for a particular sample, the program must determine a laboratory’s type of service in accordance with paragraph (a) of this section. A laboratory must interpret acid-fast stains and isolate and identify the organisms to the same extent it performs these procedures on patient specimens. A laboratory’s performance will be evaluated on the basis of the average of its scores as determined in paragraph (c)(6) of this section.

(3) Since laboratories may incorrectly report the presence of organisms in addition to the correctly identified principal organism(s), the grading system must provide a means of deducting credit for additional erroneous organisms reported. Therefore, the total number of correct responses submitted by the laboratory divided by the number of organisms present plus the number of incorrect organisms reported by the laboratory must be multiplied by 100 to establish a score for each sample in each testing event. For example, if a sample contained one principal organism and the laboratory reported it correctly but reported the presence of an additional organism, which was not present, the sample grade would be

\[
\frac{1}{(1+1)} \times 100 = 50\% 
\]

(4) For antimycobacterial susceptibility testing, a laboratory must indicate which drugs are routinely included in its test panel when testing patient samples. A laboratory’s performance will be evaluated for only those antibiotics for which susceptibility testing is routinely performed on patient specimens. A correct response for each antibiotic will be determined as described in §493.913(c)(1). Grading is based on the number of correct susceptibility responses reported by the laboratory divided by the actual number of correct susceptibility responses as determined by the program, multiplied by 100. For example, if a laboratory offers susceptibility testing using three antimycobacterial agents and the laboratory reports correct response for two of the three antimycobacterial agents, the laboratory’s grade would be

\[
\frac{2}{3} \times 100 = 67\% 
\]

(5) The performance criterion for qualitative tests is the presence or absence of acid-fast organisms. The score for acid-fast organism detection is the number of correct responses divided by the number of samples to be tested, multiplied by 100.

(6) The score for a testing event in mycobacteriology is the average of the scores determined under paragraphs (c)(3) through (c)(5) of this section.
§ 493.915 Mycology.

(a) Types of services offered by laboratories. In mycology, there are four types of laboratories for proficiency testing purposes that may perform different levels of service for yeasts, dimorphic fungi, dermatophytes, and aerobic actinomycetes:

(1) Those that isolate and identify only yeasts and/or dermatophytes to the genus level;

(2) Those that isolate and identify yeasts and/or dermatophytes to the species level;

(3) Those that isolate and perform identification of all organisms to the genus level; and

(4) Those that isolate and perform identification of all organisms to the species level.

(b) Program content and frequency of challenge. To be approved for proficiency testing for mycology, the annual program must provide a minimum of five samples per testing event. There must be at least three testing events at approximately equal intervals per year. The samples may be provided through mailed shipments or, at HHS's option, may be provided to HHS or its designee for on-site testing. An annual program must include samples that contain organisms that are representative of five major groups of fungi: Yeast or yeast-like fungi; dimorphic fungi; dematiaceous fungi; dermatophytes; and saprophytes, including opportunistic fungi. The specific fungi included in the samples may vary from year to year.

(1) An approved program must, before each calendar year, furnish HHS with a description of samples that it plans to include in its annual program no later than six months before each calendar year. At least 50 percent of the samples must be mixtures of the principal organism and appropriate normal background flora. Other important emerging pathogens (as determined by HHS) and organisms commonly occurring in patient specimens must be included periodically in the program.

(2) An approved program may vary over time. As an example, the types of organisms that might be included in an approved program over time are—

Candida albicans
Candida (other species)
Cryptococcus neoformans
Sporothrix schenckii
Exophiala jeanselmei
Fonsecae pedrosoi
M. microsporum sp.
Acremonium sp.
Trichophyton sp.
Aspergillus fumigatus
Nocardia sp.
Blastomyces dermatitidis

NOTE:1 Provided as a nonviable sample.

(c) Evaluation of a laboratory's performance. HHS approves only those programs that assess the accuracy of a laboratory's response, in accordance with paragraphs (c)(1) through (5) of this section.

(1) The program determines the reportable organisms. To determine the accuracy of a laboratory's response, the program must compare the laboratory's response for each sample with the response that reflects agreement of either 90 percent of ten or more referee laboratories or 90 percent or more of all participating laboratories.

(2) To evaluate a laboratory's response for a particular sample, the program must determine a laboratory's type of service in accordance with paragraph (a) of this section. A laboratory must isolate and identify the organisms to the same extent it performs these procedures on patient specimens.

(3) Since laboratories may incorrectly report the presence of organisms in addition to the correctly identified principal organism(s), the grading system must deduct credit for additional erroneous organisms reported. Therefore, the total number of correct responses submitted by the laboratory divided by the number of organisms present plus the number of incorrect organisms reported by the laboratory must be multiplied by 100 to establish a score for each sample in each shipment or testing event. For example, if a sample contained one principal organism and the laboratory reported it correctly but reported the presence of an additional organism, which was not
§ 493.917 Parasitology.

(a) Types of services offered by laboratories. In parasitology there are two types of laboratories for proficiency testing purposes—

(1) Those that determine the presence or absence of parasites by direct observation (wet mount) and/or pinworm preparations and, if necessary, refer specimens to another laboratory appropriately certified in the subspecialty of parasitology for identification;

(2) Those that identify parasites using concentration preparations and/or permanent stains.

(b) Program content and frequency of challenge. To be approved for proficiency testing in parasitology, a program must provide a minimum of five samples per testing event. There must be at least three testing events at approximately equal intervals per year. The samples may be provided through mailed shipments or, at HHS's option, may be provided to HHS or its designee for on-site testing. An annual program must include samples that contain parasites that are commonly encountered in the United States as well as those recently introduced into the United States. Other important emerging pathogens (as determined by HHS) and parasites commonly occurring in patient specimens must be included periodically in the program.

(1) An approved program must, before each calendar year furnish HHS with a description of samples that it plans to include in its annual program no later than six months before each calendar year. Samples must include both formalinized specimens and PVA (polyvinyl alcohol) fixed specimens as well as blood smears, as appropriate for a particular parasite and stage of the parasite. The majority of samples must contain protozoa or helminths or a combination of parasites. Some samples must be devoid of parasites.

(2) An approved program may vary over time. As an example, the types of parasites that might be included in an approved program over time are—

Enterobius vermicularis
Entamoeba histolytica
Entamoeba coli
Giardia lamblia
Endolimax nana
Dientamoeba fragilis
Iodamoeba butschlii
Chilomastix mesnili
Hookworm
Ascaris lumbricoides
Strongyloides stercoralis
Trichuris trichiura
Diphyllobothrium latum
Cryptosporidium sp.
Plasmodium falciparum

(3) For laboratories specified in paragraph (a)(1) of this section, the program must provide at least five samples per testing event that include challenges which contain parasites and challenges that are devoid of parasites.

(c) Evaluation of a laboratory's performance. HHS approves only those programs that assess the accuracy of a laboratory's responses in accordance with paragraphs (c)(1) through (6) of this section.

(1) The program must determine the reportable parasites. It may elect to establish a minimum number of parasites to be identified in samples before they are reported. Parasites found in rare numbers by referee laboratories are not considered in scoring a laboratory's performance; such findings are neutral. To determine the accuracy of a laboratory's response, the program must compare the laboratory's response with the response that reflects agreement of either 90 percent of ten or more referee laboratories or 90 percent or more of all participating laboratories.

(2) To evaluate a laboratory's response for a particular sample, the program must determine a laboratory's type of service in accordance with paragraph (a) of this section. A laboratory must determine the presence or absence of a parasite(s) or concentrate and identify the parasites to the same extent it performs these procedures on patient specimens.
§ 493.919 Virology.

(a) Types of services offered by laboratories. In virology, there are two types of laboratories for proficiency testing purposes—

(1) Those that only perform tests that directly detect viral antigens or structures, either in cells derived from infected tissues or free in fluid specimens; and

(2) Those that are able to isolate and identify viruses and use direct antigen techniques.

(b) Program content and frequency of challenge. To be approved for proficiency testing in virology, a program must provide a minimum of five samples per testing event. There must be at least three testing events at approximately equal intervals per year. The samples may be provided to the laboratory through mailed shipments or, at HHS's option, may be provided to HHS or its designee for on-site testing.

An annual program must include viral species that are the more commonly identified viruses. The specific organisms found in the samples may vary from year to year. The annual program must include samples for viral antigen detection and viral isolation and identification.

(1) An approved program must furnish HHS with a description of samples that it plans to include in its annual program no later than six months before each calendar year. The program must include other important emerging viruses (as determined by HHS) and viruses commonly occurring in patient specimens.

(2) An approved program may vary over time. For example, the types of viruses that might be included in an approved program over time are the more commonly identified viruses such as Herpes simplex, respiratory syncytial virus, adenoviruses, enteroviruses, and cytomegaloviruses.

(c) Evaluation of laboratory's performance. HHS approves only those programs that assess the accuracy of a laboratory's response in accordance with paragraphs (c)(1) through (5) of this section.

(1) The program determines the reportable viruses to be detected by direct antigen techniques or isolated by laboratories that perform viral isolation procedures. To determine the accuracy of a laboratory's response, the program must compare the laboratory's response for each sample with the response that reflects agreement of either 90 percent of ten or more referee laboratories or 90 percent or more of all participating laboratories.

(2) To evaluate a laboratory's response for a particular sample, the program must determine a laboratory's type of service in accordance with paragraph (a) of this section. A laboratory must isolate and identify the viruses to the same extent it performs these procedures on patient specimens.

(3) Since laboratories may incorrectly report the presence of viruses in addition to the correctly identified principal virus, the grading system must provide a means of deducting credit for additional erroneous viruses reported. Therefore, the total number of correct responses determined by
virus culture techniques submitted by the laboratory divided by the number of viruses present plus the number of incorrect viruses reported by the laboratory must be multiplied by 100 to establish a score for each sample in each testing event. For example, if a sample contained one principal virus and the laboratory reported it correctly but reported the presence of an additional virus, which was not present, the sample grade would be 1/(1+1)×100=50 percent.

(4) The performance criterion for qualitative antigen tests is presence or absence of the viral antigen. The score for the antigen tests is the number of correct responses divided by the number of samples to be tested for the antigen, multiplied by 100.

(5) The score for a testing event is the average of the sample scores as determined under paragraph (c)(3) and (c)(4) of this section.

§ 493.921 Diagnostic immunology.

The subspecialties under the specialty of immunology for which a program may offer proficiency testing are syphilis serology and general immunology. Specific criteria for these subspecialties are found at §§ 493.923 and 493.927.

§ 493.923 Syphilis serology.

(a) Program content and frequency of challenge. To be approved for proficiency testing in syphilis serology, a program must provide a minimum of five samples per testing event. There must be at least three testing events at approximately equal intervals per year. The samples may be provided through mailed shipments or, at HHS’ option, may be provided to HHS or its designee for on-site testing. An annual program must include samples that cover the full range of reactivity from highly reactive to non-reactive.

(b) Evaluation of test performance. HHS approves only those programs that assess the accuracy of a laboratory’s response in accordance with paragraphs (b)(1) through (4) of this section.

(1) To determine the accuracy of a laboratory’s response for qualitative and quantitative syphilis tests, the program must compare the laboratory’s response with the response that reflects agreement of either 90 percent of ten or more referee laboratories or 90 percent or more of all participating laboratories. The proficiency testing program must indicate the minimum concentration, by method, that will be considered as indicating a positive response. The score for a sample in syphilis serology is the average of scores determined under paragraphs (b)(2) and (b)(3) of this section.

(2) For quantitative syphilis tests, the program must determine the correct response for each method by the distance of the response from the target value. After the target value has been established for each response, the appropriateness of the response must be determined by using fixed criteria. The criterion for acceptable performance for quantitative syphilis serology tests is the target value ±1 dilution.

(3) The criterion for acceptable performance for qualitative syphilis serology tests is reactive or nonreactive.

(4) To determine the overall testing event score, the number of correct responses must be averaged using the following formula:

\[
\frac{\text{Number of acceptable responses for all challenges}}{\text{Total number of all challenges}} \times 100 = \text{Testing event score}
\]


§ 493.927 General immunology.

(a) Program content and frequency of challenge. To be approved for proficiency testing for immunology, the annual program must provide a minimum of five samples per testing event. There must be at least three testing events at approximately equal intervals per year. The annual program must provide samples that cover the full range of reactivity from highly reactive to nonreactive. The samples may be provided through mailed shipments or, at HHS’ option, may be provided to HHS or its designee for on-site testing.

(b) Challenges per testing event. The minimum number of challenges per
testing event the program must provide for each analyte or test procedure is five. Analytes or tests for which laboratory performance is to be evaluated include:

<table>
<thead>
<tr>
<th>Analyte or Test Procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alpha-1 antitrypsin</td>
</tr>
<tr>
<td>Alpha-fetoprotein (tumor marker)</td>
</tr>
<tr>
<td>Antinuclear antibody</td>
</tr>
<tr>
<td>Antistreptolysin O</td>
</tr>
<tr>
<td>Anti-human immunodeficiency virus (HIV)</td>
</tr>
<tr>
<td>Complement C3</td>
</tr>
<tr>
<td>Complement C4</td>
</tr>
<tr>
<td>Hepatitis markers (HBsAg, anti-HBc, HBeAg)</td>
</tr>
<tr>
<td>IgA</td>
</tr>
<tr>
<td>IgG</td>
</tr>
<tr>
<td>IgE</td>
</tr>
<tr>
<td>IgM</td>
</tr>
<tr>
<td>Infectious mononucleosis</td>
</tr>
<tr>
<td>Rheumatoid factor</td>
</tr>
<tr>
<td>Rubella</td>
</tr>
</tbody>
</table>

(c) Evaluation of a laboratory’s analyte or test performance. HHS approves only those programs that assess the accuracy of a laboratory’s responses in accordance with paragraphs (c)(1) through (5) of this section.

(1) To determine the accuracy of a laboratory’s response for quantitative and qualitative immunology tests or analytes, the program must compare the laboratory’s response for each analyte with the response that reflects agreement of either 90 percent of ten or more referee laboratories or 90 percent of all participating laboratories. The proficiency testing program must indicate the minimum concentration that will be considered as indicating a positive response. The score for a sample in general immunology is either the score determined under paragraph (c)(2) or (3) of this section.

(2) For quantitative immunology analytes or tests, the program must determine the correct response for each analyte by the distance of the response from the target value. After the target value has been established for each response, the appropriateness of the response must be determined by using either fixed criteria or the number of standard deviations (SDs) the response differs from the target value.

#### Criteria for Acceptable Performance

The criteria for acceptable performance are—

<table>
<thead>
<tr>
<th>Analyte or test</th>
<th>Criteria for acceptable performance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alpha-1 antitrypsin</td>
<td>Target value ± 3 SD.</td>
</tr>
<tr>
<td>Alpha-fetoprotein (tumor marker)</td>
<td>Target value ± 3 SD.</td>
</tr>
<tr>
<td>Antinuclear antibody</td>
<td>Target value ± 2 dilutions or positive or negative.</td>
</tr>
<tr>
<td>Antistreptolysin O</td>
<td>Target value ± 2 dilution or positive or negative.</td>
</tr>
<tr>
<td>Anti-Human Immunodeficiency virus</td>
<td>Target value ± 25%</td>
</tr>
<tr>
<td>Complement C3</td>
<td>Target value ± 3 SD.</td>
</tr>
<tr>
<td>Complement C4</td>
<td>Target value ± 2 dilutions or positive or negative.</td>
</tr>
<tr>
<td>Hepatitis (HBsAg, anti-HBc, HBeAg)</td>
<td>Target value ± 2 dilutions or positive or negative.</td>
</tr>
<tr>
<td>IgA</td>
<td>Target value ± 3 SD.</td>
</tr>
<tr>
<td>IgG</td>
<td>Target value ± 3 SD.</td>
</tr>
<tr>
<td>IgE</td>
<td>Target value ± 25%</td>
</tr>
<tr>
<td>IgM</td>
<td>Target value ± 3 SD.</td>
</tr>
<tr>
<td>Infectious mononucleosis</td>
<td>Target value ± 2 dilutions or positive or negative.</td>
</tr>
<tr>
<td>Rheumatoid factor</td>
<td>Target value ± 2 dilutions or positive or negative.</td>
</tr>
<tr>
<td>Rubella</td>
<td>Target value ± 2 dilutions or immune or nonimmune or positive or negative.</td>
</tr>
</tbody>
</table>

(3) The criterion for acceptable performance for qualitative general immunology tests is positive or negative.

(4) To determine the analyte testing event score, the number of acceptable analyte responses must be averaged using the following formula:

\[
\text{Number of acceptable responses for the analyte} \times \frac{100}{\text{Total number of challenges for the analyte}} = \text{Analyte score for the testing event}
\]

(5) To determine the overall testing event score, the number of correct responses for all analytes must be averaged using the following formula:

\[
\text{Number of acceptable responses for all challenges} \times \frac{100}{\text{Total number of all challenges}} = \text{Testing event score}
\]
§ 493.929 Chemistry.

The subspecialties under the specialty of chemistry for which a proficiency testing program may offer proficiency testing are routine chemistry, endocrinology, and toxicology. Specific criteria for these subspecialties are listed in §§ 493.931 through 493.939.

§ 493.931 Routine chemistry.

(a) Program content and frequency of challenge. To be approved for proficiency testing for routine chemistry, a program must provide a minimum of five samples per testing event. There must be at least three testing events at approximately equal intervals per year. The annual program must provide samples that cover the clinically relevant range of values that would be expected in patient specimens. The specimens may be provided through mailed shipments or, at HHS' option, may be provided to HHS or its designee for on-site testing.

(b) Challenges per testing event. The minimum number of challenges per testing event a program must provide for each analyte or test procedure listed below is five serum, plasma or blood samples.

### Analyte or Test Procedure

<table>
<thead>
<tr>
<th>Analyte or Test Procedure</th>
<th>Criteria for acceptable performance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alanine aminotransferase (ALT/SGPT)</td>
<td>Target value ±20%</td>
</tr>
<tr>
<td>Albumin</td>
<td>Target value ±10%</td>
</tr>
<tr>
<td>Alkaline phosphatase</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>Amylase</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>Aspartate aminotransferase (AST/SGOT)</td>
<td>Target value ±20%</td>
</tr>
<tr>
<td>Bilirubin, total</td>
<td>Target value ±0.4 mg/dL or ±20% (greater)</td>
</tr>
<tr>
<td>Blood gas (pH, pO2, and pCO2)</td>
<td>Target value ±3 SD or ±8% (greater)</td>
</tr>
<tr>
<td>Calcium, total</td>
<td>Target value ±10.0 mg/dL</td>
</tr>
<tr>
<td>Chloride</td>
<td>Target value ±5%</td>
</tr>
<tr>
<td>Cholesterol, total</td>
<td>Target value ±10%</td>
</tr>
<tr>
<td>Cholesterol, high density lipoprotein</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>Creatine kinase</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>Creatine kinase isoenzymes</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>Creatinine</td>
<td>Target value ±0.3 mg/dL or ±15% (greater)</td>
</tr>
<tr>
<td>Glucose (Excluding measurements on devices cleared by FDA for home use)</td>
<td>Target value ±10%</td>
</tr>
<tr>
<td>Iron, total</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>Lactate dehydrogenase (LDH)</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>LDH isoenzymes</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>Magnesium</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>Potassium</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>Total Protein</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>Triglycerides</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>Urea Nitrogen</td>
<td>Target value ±30%</td>
</tr>
<tr>
<td>Uric Acid</td>
<td>Target value ±30%</td>
</tr>
</tbody>
</table>
| (c) Evaluation of a laboratory's analyte or test performance. HHS approves only those programs that assess the accuracy of a laboratory's responses in accordance with paragraphs (c)(1) through (5) of this section.

(1) To determine the accuracy of a laboratory's response for qualitative and quantitative chemistry tests or analytes, the program must compare the laboratory's response for each analyte with the response that reflects agreement of either 90 percent of ten or more referee laboratories or 90 percent or more of all participating laboratories. The score for a sample in routine chemistry is either the score determined under paragraph (c)(2) or (3) of this section.

(2) For quantitative chemistry tests or analytes, the program must determine the correct response for each analyte by the distance of the response from the target value. After the target value has been established for each response, the appropriateness of the response must be determined by using either fixed criteria based on the percentage difference from the target value or the number of standard deviations (SDs) the response differs from the target value.
§ 493.933 Endocrinology.

(a) Program content and frequency of challenge. To be approved for proficiency testing for endocrinology, a program must provide a minimum of five samples per testing event. There must be at least three testing events at approximately equal intervals per year. The annual program must provide samples that cover the clinically relevant range of values that would be expected in patient specimens. The samples may be provided through mailed shipments or, at HHS’ option, may be provided to HHS or its designee for on-site testing.

(b) Challenges per testing event. The minimum number of challenges per testing event a program must provide for each analyte or test procedure is five serum, plasma, blood, or urine samples.

Analyte or Test

<table>
<thead>
<tr>
<th>Analyte or Test</th>
<th>Criteria for acceptable performance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cortisol</td>
<td>Target value ± 25% (greater)</td>
</tr>
<tr>
<td>Free Thyroxine</td>
<td></td>
</tr>
<tr>
<td>Human Chorionic gonadotropin</td>
<td></td>
</tr>
<tr>
<td>T3 Uptake</td>
<td></td>
</tr>
<tr>
<td>Triiodothyronine</td>
<td></td>
</tr>
<tr>
<td>Thyroid-stimulating hormone</td>
<td></td>
</tr>
<tr>
<td>Thyroxine</td>
<td></td>
</tr>
</tbody>
</table>

(c) Evaluation of a laboratory’s analyte or test performance.

HHS approves only those programs that assess the accuracy of a laboratory’s response in accordance with paragraphs (c)(1) through (5) of this section.

(1) To determine the accuracy of a laboratory’s response for qualitative and quantitative endocrinology tests or analytes, a program must compare the laboratory’s response for each analyte with the response that reflects agreement of either 90 percent of ten or more referee laboratories or 90 percent or more of all participating laboratories. The score for a sample in endocrinology is either the score determined under paragraph (c)(2) or (c)(3) of this section.

(2) For quantitative endocrinology tests or analytes, the program must determine the correct response for each analyte by the distance of the response from the target value. After the target value has been established for each response, the appropriateness of the response must be determined by using either fixed criteria based on the percentage difference from the target value or the number of standard deviations (SDs) the response differs from the target value.

Criteria for Acceptable Performance

<table>
<thead>
<tr>
<th>Analyte or test</th>
<th>Criteria for acceptable performance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cortisol</td>
<td>Target value ± 25% (greater)</td>
</tr>
<tr>
<td>Free Thyroxine</td>
<td>Target value ± 3 SD.</td>
</tr>
</tbody>
</table>
Health Care Financing Administration, HHS

§ 493.937

Toxicology.

(a) Program content and frequency of challenge. To be approved for proficiency testing for toxicology, the annual program must provide a minimum of five samples per testing event. There must be at least three testing events at approximately equal intervals per year. The annual program must provide samples that cover the clinically relevant range of values that would be expected in specimens of patients on drug therapy and that cover the level of clinical significance for the particular drug. The samples may be provided through mailed shipments or, at HHS’ option, may be provided to HHS or its designee for on-site testing.

(b) Challenges per testing event. The minimum number of challenges per testing event a program must provide for each analyte or test procedure is five serum, plasma, or blood samples.

Analyte or Test Procedure

| Alcohol (blood) | Phenytoin |
| Blood lead | Primidone |
| Carbamazepine | Procainamide |
| Digoxin | (and metabolite) |
| Ethosuximide | Quinidine |
| Gentamicin | Theophylline |
| Lithium | Tobramycin |
| Phenobarbital | Valproic Acid |

(c) Evaluation of a laboratory’s analyte or test performance. HHS approves only those programs that assess the accuracy of a laboratory’s responses in accordance with paragraphs (c)(1) through (4) of this section.

(1) To determine the accuracy of a laboratory’s responses for quantitative toxicology tests or analytes, the program must compare the laboratory’s response for each analyte with the response that reflects agreement of either 90 percent of ten or more referee laboratories or 90 percent or more of all participating laboratories. The score for a sample in toxicology is the score determined under paragraph (c)(2) of this section.

(2) For quantitative toxicology tests or analytes, the program must determine the correct response for each analyte by the distance of the response from the target value. After the target value has been established for each response, the appropriateness of the response must be determined by using fixed criteria based on the percentage difference from the target value.

Criteria for Acceptable Performance

The criteria for acceptable performance are:

<table>
<thead>
<tr>
<th>Analyte or test</th>
<th>Criteria for acceptable performance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcohol, blood</td>
<td>Target Value ± 25%</td>
</tr>
<tr>
<td>Blood lead</td>
<td>Target Value ±10% or 4 mcg/dL (greater)</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Target Value ± 25%</td>
</tr>
<tr>
<td>Digoxin</td>
<td>Target Value ± 20% or ± 0.2 ng/mL (greater)</td>
</tr>
<tr>
<td>Ethosuximide</td>
<td>Target Value ± 20%</td>
</tr>
<tr>
<td>Gentamicin</td>
<td>Target Value ± 25%</td>
</tr>
<tr>
<td>Lithium</td>
<td>Target Value ≥ 0.3 mmol/L or ≥ 20% (greater)</td>
</tr>
</tbody>
</table>
§493.941 Hematology (including routine hematology and coagulation).

(a) Program content and frequency of challenge. To be approved for proficiency testing for hematology, a program must provide a minimum of five samples per testing event. There must be at least three testing events at approximately equal intervals per year. The annual program must provide samples that cover the full range of values that would be expected in patient specimens. The samples may be provided through mailed shipments or, at HHS' option, may be provided to HHS and or its designee for on-site testing.

(b) Challenges per testing event. The minimum number of challenges per testing event a program must provide for each analyte or test procedure is five.

(c) Evaluation of a laboratory's analyte or test performance. HHS approves only those programs that assess the accuracy of a laboratory's responses in accord with paragraphs (c) (1) through (5) of this section.

(1) To determine the accuracy of a laboratory's responses for qualitative and quantitative hematology tests or analytes, the program must compare the laboratory's response for each analyte with the response that reflects agreement of either 90 percent of ten or more referee laboratories or 90 percent or more of all participating laboratories. The score for a sample in hematology is either the score determined under paragraph (c) (2) or (3) of this section.

(2) For quantitative hematology tests or analytes, the program must determine the correct response for each analyte by the distance of the response from the target value. After the target value has been established for each response, the appropriateness of the response is determined using either fixed criteria based on the percentage difference from the target value or the number of standard deviations (SDs) the response differs from the target value.
Criteria for Acceptable Performance

The criteria for acceptable performance are:

<table>
<thead>
<tr>
<th>Analyte or Test</th>
<th>Criteria for acceptable performance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell identification</td>
<td>90% or greater consensus on identification.</td>
</tr>
<tr>
<td>White blood cell differential</td>
<td>Target = \pm 3SD based on the percentage of different types of white blood cells in the samples.</td>
</tr>
<tr>
<td>Erythrocyte count</td>
<td>Target = \pm 6%.</td>
</tr>
<tr>
<td>Hematocrit (Excluding spun hematocrits)</td>
<td>Target = \pm 6%.</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>Target = \pm 7%.</td>
</tr>
<tr>
<td>Leukocyte count</td>
<td>Target = \pm 15%.</td>
</tr>
<tr>
<td>Platelet count</td>
<td>Target = \pm 25%.</td>
</tr>
<tr>
<td>Fibrinogen</td>
<td>Target = \pm 15%.</td>
</tr>
<tr>
<td>Partial thromboplastin time</td>
<td></td>
</tr>
<tr>
<td>Prothrombin time</td>
<td></td>
</tr>
</tbody>
</table>

(3) The criterion for acceptable performance for the qualitative hematology test is correct cell identification.

(4) To determine the analyte testing event score, the number of acceptable analyte responses must be averaged using the following formula:

\[
\text{Number of acceptable responses for the analyte} \times 100 = \text{Analyte score for the testing event}
\]

\[
\text{Total number of challenges for the analyte}
\]

(5) To determine the overall testing event score, the number of correct responses for all analytes must be averaged using the following formula:

\[
\text{Number of acceptable responses for all challenges} \times 100 = \text{Testing event score}
\]

\[
\text{Total number of all challenges}
\]


§ 493.945 Cytology; gynecologic examinations.

(a) Program content and frequency of challenge. (1) To be approved for proficiency testing for gynecologic examinations (Pap smears) in cytology, a program must provide test sets composed of 10- and 20-glass slides. Proficiency testing programs may obtain slides for test sets from cytology laboratories, provided the slides have been retained by the laboratory for the required period specified in §493.1257. If slide preparations are still subject to retention by the laboratory, they may be loaned to a proficiency testing program if the program provides the laboratory with documentation of the loan of the slides and ensures that slides loaned to it are retrievable upon request. Each test set must include at least one slide representing each of the response categories described in paragraph (b)(3)(ii)(A) of this section, and test sets should be comparable so that equitable testing is achieved within and between proficiency testing providers.

(2) To be approved for proficiency testing in gynecologic cytology, a program must provide announced and unannounced on-site testing for each individual at least once per year and must provide an initial retesting event for each individual within 45 days after notification of test failure and subsequent retesting events within 45 days after completion of remedial action described in §493.855.

(b) Evaluation of an individual's performance. HHS approves only those programs that assess the accuracy of each individual's responses on both 10- and 20-slide test sets in which the slides have been referenced as specified in paragraph (b)(1) of this section.

(1) To determine the accuracy of an individual's response on a particular challenge (slide), the program must compare the individual's response for each slide preparation with the response that reflects the predetermined consensus agreement or confirmation on the diagnostic category, as described in the table in paragraph (b)(3)(ii)(A) of this section. For all slide preparations, a 100% consensus agreement among a minimum of three physicians certified in anatomic pathology is required. In addition, for premalignant and malignant slide preparations, confirmation by tissue biopsy is required either by comparison of the reported biopsy results or reevaluation of biopsy slide material by a physician certified in anatomic pathology.

(2) An individual qualified as a technical supervisor under §493.1449 (b) or
(k) who routinely interprets gynecologic slide preparations only after they have been examined by a cytotechnologist can either be tested using a test set that has been screened by a cytotechnologist in the same laboratory or using a test set that has not been screened. A technical supervisor who screens and interprets slide preparations that have not been previously examined must be tested using a test set that has not been previously screened.

(3) The criteria for acceptable performance are determined by using the scoring system in paragraphs (b)(3)(i) and (ii) of this section.

(i) Each slide set must contain 10 or 20 slides with point values established for each slide preparation based on the significance of the relationship of the interpretation of the slide to a clinical condition and whether the participant in the testing event is a cytotechnologist qualified under §§493.1469 or 493.1483 or functioning as a technical supervisor in cytology qualified under §493.1449 (b) or (k) of this part.

(ii) The scoring system rewards or penalizes the participants in proportion to the distance of their answers from the correct response or target diagnosis and the penalty or reward is weighted in proportion to the severity of the lesion.

(A) The four response categories for reporting proficiency testing results and their descriptions are as follows:

<table>
<thead>
<tr>
<th>Category</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>A ...............</td>
<td>Unsatisfactory for diagnosis due to:</td>
</tr>
<tr>
<td>...............</td>
<td>(1) Scant cellularity.</td>
</tr>
<tr>
<td>...............</td>
<td>(2) Air drying.</td>
</tr>
<tr>
<td>...............</td>
<td>(3) Obscuring material (blood, inflammatory cells, or lubricant).</td>
</tr>
<tr>
<td>B ...............</td>
<td>Normal or Benign Changes—includes:</td>
</tr>
<tr>
<td>...............</td>
<td>(1) Normal, negative or within normal limits.</td>
</tr>
<tr>
<td>...............</td>
<td>(2) Infection other than Human Papillomavirus (HPV) (e.g., Trichomonas vaginalis, changes or morphology consistent with Candida spp., Actinomyces spp. or Herpes simplex virus).</td>
</tr>
<tr>
<td>...............</td>
<td>(3) Reactive and reparative changes (e.g., inflammation, effects of chemotherapy or radiation).</td>
</tr>
<tr>
<td>C ...............</td>
<td>Low Grade Squamous Intraepithelial Lesion—including:</td>
</tr>
</tbody>
</table>

(B) In accordance with the criteria for the scoring system, the charts in paragraphs (b)(3)(ii)(C) and (D) of this section, for technical supervisors and cytotechnologists, respectively, provide a maximum of 10 points for a correct response and a maximum of minus five (−5) points for an incorrect response on a 10-slide test set. For example, if the correct response on a slide is “High Grade Squamous intraepithelial lesion” (category “D” on the scoring system chart) and an examinee calls it “normal or negative” (category “B” on the scoring system chart), then the examinee’s point value on that slide is calculated as minus five (−5). Each slide is scored individually in the same manner. The individual’s score for the testing event is determined by adding the point value achieved for each slide preparation, dividing by the total points for the testing event and multiplying by 100.

(C) Criteria for scoring system for a 10-slide test set. (See table at (b)(3)(ii)(A) of this section for a description of the response categories.) For technical supervisors qualified under §493.1449 (b) or (k):

<table>
<thead>
<tr>
<th>Examinee’s response:</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
</tr>
</thead>
<tbody>
<tr>
<td>Correct response:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>10</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>B</td>
<td>5</td>
<td>10</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>C</td>
<td>5</td>
<td>0</td>
<td>10</td>
<td>5</td>
</tr>
<tr>
<td>D</td>
<td>0</td>
<td>5</td>
<td>5</td>
<td>10</td>
</tr>
</tbody>
</table>

(D) Criteria for scoring system for a 10-slide test set. (See table at paragraph (b)(3)(ii)(A) of this section for a description of the response categories.) For cytotechnologists qualified under §§493.1469 or 493.1483:
§ 493.959 Immunohematology.

(a) Types of services offered by laboratories. In immunohematology, there are four types of laboratories for proficiency testing purposes—

(1) Those that perform ABO group and/or D (Rho) typing;

(2) Those that perform ABO group and/or D (Rho) typing, and unexpected antibody detection;

(3) Those that in addition to paragraph (a)(2) of this section perform compatibility testing; and

(4) Those that perform in addition to paragraph (a)(3) of this section antibody identification.

(b) Program content and frequency of challenge. To be approved for proficiency testing for immunohematology, a program must provide a minimum of five samples per testing event. There must be at least three testing events at approximately equal intervals per year. The annual program must provide samples that cover the full range of interpretation that would be expected in patient specimens. The samples may be provided through mailed shipments or, at HHS’ option, may be provided to HHS or its designee for on-site testing.

(c) Challenges per testing event. The minimum number of challenges per testing event a program must provide for each analyte or test procedure is five.

(Analyte or Test Procedure)

ABO group (excluding subgroups)
D (Rho) typing
Unexpected antibody detection
Compatibility testing
Antibody identification

(d) Evaluation of a laboratory’s analyte or test performance. HHS approves only those programs that assess the accuracy of a laboratory’s response in accordance with paragraphs (d)(1) through (5) of this section.

(1) To determine the accuracy of a laboratory’s response, a program must compare the laboratory’s response for each analyte with the response that reflects agreement of either 100 percent of ten or more referee laboratories or 95 percent or more of all participating laboratories except for unexpected antibody detection and antibody identification. To determine the accuracy of a laboratory’s response for unexpected antibody detection and antibody identification, a program must compare the laboratory’s response for each analyte with the response that reflects agreement of either 95 percent of ten or more referee laboratories or 95 percent or more of all participating laboratories. The score for a sample in immunohematology is either the score determined under paragraph (d)(2) or (3) of this section.
§493.1101 Criteria for acceptable performance.
The criteria for acceptable performance are—

<table>
<thead>
<tr>
<th>Analyte or test</th>
<th>Criteria for acceptable performance</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABO group</td>
<td>100% accuracy.</td>
</tr>
<tr>
<td>D (Rh) typing</td>
<td>100% accuracy.</td>
</tr>
<tr>
<td>Unexpected antibody detection</td>
<td>80% accuracy.</td>
</tr>
<tr>
<td>Compatibility testing</td>
<td>100% accuracy.</td>
</tr>
<tr>
<td>Antibody identification</td>
<td>80% accuracy.</td>
</tr>
</tbody>
</table>

(3) The criterion for acceptable performance for qualitative immunohematology tests is positive or negative.

(4) To determine the analyte testing event score, the number of acceptable analyte responses must be averaged using the following formula:

\[\text{Number of acceptable responses for the analyte} \times 100 \div \text{Total number of challenges for the analyte event} \]

(5) To determine the overall testing event score, the number of correct responses for all analytes must be averaged using the following formula:

\[\text{Number of acceptable responses for all challenges} \times 100 \div \text{Total number of all challenges} \]

Subpart J—Patient Test Management for Moderate Complexity (Including the Subcategory), High Complexity, or Any Combination of These Tests

SOURCE: 57 FR 7162, Feb. 28, 1992, unless otherwise noted.

§493.1101 Condition: Patient test management; moderate complexity (including the subcategory), or high complexity testing, or any combination of these tests.

Each laboratory performing moderate complexity (including the subcategory) or high complexity testing, or any combination of these tests, must employ and maintain a system that provides for proper patient preparation; proper specimen collection, identification, preservation, transportation, and processing; and accurate result reporting. This system must assure optimum patient specimen integrity and positive identification throughout the preanalytic (pre-testing), analytic (testing), and postanalytic (post-testing) processes and must meet the standards as they apply to the testing performed.

[60 FR 20048, Apr. 24, 1995]

§ 493.1103 Standard; Procedures for specimen submission and handling.

(a) The laboratory must have available and follow written policies and procedures for each of the following, if applicable: Methods used for the preparation of patients; specimen collection; specimen labeling; specimen preservation; conditions for specimen transportation; and specimen processing. Such policies and procedures must assure positive identification and optimum integrity of the patient specimens from the time the specimen(s) are collected until testing has been completed and the results reported.

(b) If the laboratory accepts referral specimens, written instructions must be available to clients and must include, as appropriate, the information specified in paragraph (a) of this section.

(c) Oral explanation of instructions to patients for specimen collection, including patient preparation, may be used as a supplement to written instructions where applicable.


§ 493.1105 Standard; Test requisition.

The laboratory must perform tests only at the written or electronic request of an authorized person. Oral requests for laboratory tests are permitted only if the laboratory subsequently requests written authorization for testing within 30 days. The laboratory must maintain the written authorization or documentation of efforts made to obtain a written authorization. Records of test requisitions or test authorizations must be retained for a minimum of two years. The patient’s chart or medical record, if used as the test requisition, must be retained for a minimum of two years and
§ 493.1109 Standard; Test report.

The laboratory report must be sent promptly to the authorized person, the individual responsible for using the test results or laboratory that initially requested the test. The original report or an exact duplicate of each test report, including final and preliminary report, must be retained by the testing laboratory for a period of at least two years after the date of reporting. Immunohematology reports and transfusion records must be retained by the laboratory for a period of no less than five years in accordance with 21 CFR part 606, subpart I. In addition, records of blood and blood product testing must be maintained for a period not less than five years after processing records have been completed, or six months after the latest expiration date, whichever is the later date, in accordance with 21 CFR 606.160(d). For pathology, test reports must be retained for a period of at least ten years after the date of reporting. This information may be maintained as part of the patient’s chart or medical record which must be readily available to the laboratory and to HHS upon request.

(a) The laboratory must have adequate systems in place to report results in a timely, accurate, reliable and confidential manner, and, ensure patient confidentiality throughout those parts of the total testing process that are under the laboratory’s control.

(a) The patient identification number, accession number, or other unique identification of the specimen;
(b) The date and time of specimen receipt into the laboratory;
(c) The condition and disposition of specimens that do not meet the laboratory’s criteria for specimen acceptability; and
(d) The records and dates of all specimen testing, including the identity of the personnel who performed the test(s), which are necessary to assure proper identification and accurate reporting of patient test results.

§ 493.1111 Standard; Referral of specimens.

A laboratory must refer specimens for testing only to a laboratory possessing a valid certificate authorizing the performance of testing in the specialty or subspecialty of service for the level of complexity in which the referred test is categorized.

(a) The referring laboratory must not revise results or information directly related to the interpretation of results provided by the testing laboratory.

(b) The referring laboratory may permit each testing laboratory to send the test result directly to the authorized person who initially requested the test. The referring laboratory must retain or be able to produce an exact duplicate of each testing laboratory’s report.

(c) The authorized person who orders a test or procedure must be notified by the referring laboratory of the name and address of each laboratory location at which a test was performed.

Subpart K—Quality Control for Tests of Moderate Complexity (Including the Subcategory), High Complexity, or Any Combination of These Tests

SOURCE: 57 FR 7163, Feb. 28, 1992, unless otherwise noted.

§ 493.1201 Condition: General quality control; moderate complexity (including the subcategory) or high complexity testing, or any combination of these tests.

(a) Applicability of subpart K of this part. Subpart K is divided into two sections, general quality control and quality control for specialties and subspecialties. The quality control requirements are specified in §§ 493.1201 through 493.1285 unless—

(1) An alternative procedure specified in the manufacturer’s protocol has been cleared by the Food and Drug Administration (FDA) as meeting certain CLIA requirements for general quality control and specialty/subspecialty quality control, and the manufacturer’s instructions contain the following statement,

Unless this device is modified by a laboratory, the laboratory’s compliance with these quality control instructions will satisfy the applicable requirements of 42 CFR 493.1203(b), or
(2) HHS approves an equivalent procedure that is specified in Appendix C of the State Operations Manual (HCFA Pub. 7).

(b) The laboratory must establish and follow written quality control procedures for monitoring and evaluating the quality of the analytical testing process of each method to assure the accuracy and reliability of patient test results and reports. The laboratory must meet the applicable standards in §§493.1202 through 493.1221 of this subpart, unless an alternative procedure specified in the manufacturer’s protocol has been cleared by the Food and Drug Administration (FDA) as meeting certain CLIA requirements for quality control or HHS approves an equivalent procedure specified in appendix C of the State Operations Manual (HCFA Pub. 7). HCF Pub. 7 is available from the Technical Information Service, U.S. Department of Commerce, 5825 Port Royal Road, Springfield, VA 22161, telephone number (703) 487-4630.

[58 FR 5230, Jan. 19, 1993, as amended at 60 FR 20048, Apr. 24, 1995]

§ 493.1202 Standard; Moderate or high complexity testing, or both: Effective from September 1, 1992 to December 31, 2000.

(a) For each test of high complexity performed, the laboratory must meet all applicable standards of this subpart.

(b) For each test of moderate complexity performed using a standardized method, or method developed in-house, a device not subject to clearance by the FDA (including any commercially distributed instrument, kit or test system subject to the Food, Drug and Cosmetic Act marketed prior to the Medical Device Amendments, Public Law 94-295, enacted on May 28, 1976, and those identified in 21 CFR parts 862, 864, and 866 as exempt from FDA premarket review), or using an instrument, kit or test system cleared by the FDA through the premarket notification (510(k)) or premarket approval (PMA) process for in-vitro diagnostic use, the laboratory must—

(1) Follow the manufacturer’s instructions for instrument or test system operation and test performance;

(2) Have a procedure manual describing the processes for testing and reporting patient test results;

(3) Perform and document calibration procedures or check calibration at least once every six months;

(4) Perform and document control procedures using at least two levels of control materials each day of testing;

(5) Perform and document applicable specialty and subspecialty control procedures as specified under §493.1223;

(6) Perform and document that remedial action has been taken when problems or errors are identified as specified in §493.1219; and

(7) Maintain records of all quality control activities for two years. Quality control records for immunohematology and blood and blood products must be maintained as specified in §493.1221.

[57 FR 7163, Feb. 28, 1992, as amended at 58 FR 5230, Apr. 24, 1993]

§ 493.1203 Standard; Moderate or high complexity testing, or both: Effective beginning December 31, 2000.

For each moderate or high complexity test performed, the laboratory will be in compliance with this section if it:

(a) Meets all applicable quality control requirements specified in this subpart when using a standardized method, a method developed in-house, a device not subject to clearance by the FDA (including any commercially distributed instrument, kit or test system subject to the Food, Drug and Cosmetic Act marketed prior to the Medical Device Amendments, Public Law 94-295, enacted on May 28, 1976, and those identified in 21 CFR parts 862, 864, and 866 as exempt from FDA premarket review), or using an instrument, kit or test system cleared by the FDA through the premarket notification (510(k)) or premarket approval (PMA) process for in-vitro diagnostic use but modified by the laboratory, the laboratory must meet all applicable standards of this subpart.

(b) For all other tests of moderate complexity performed using an instrument, kit or test system cleared by the FDA through the premarket notification (510(k)) or premarket approval (PMA) process for in-vitro diagnostic use, the laboratory must—

(1) Follow the manufacturer’s instructions for instrument or test system operation and test performance;

(2) Have a procedure manual describing the processes for testing and reporting patient test results;

(3) Perform and document calibration procedures or check calibration at least once every six months;

(4) Perform and document control procedures using at least two levels of control materials each day of testing;

(5) Perform and document applicable specialty and subspecialty control procedures as specified under §493.1223;

(6) Perform and document that remedial action has been taken when problems or errors are identified as specified in §493.1219; and

(7) Maintain records of all quality control activities for two years. Quality control records for immunohematology and blood and blood products must be maintained as specified in §493.1221.
§ 493.1204  Standard; Facilities.

The laboratory must provide the space and environmental conditions necessary for conducting the services offered.

(a) The laboratory must be constructed, arranged, and maintained to ensure the space, ventilation, and utilities necessary for conducting all phases of testing, including the preanalytic (pre-testing), analytic (testing), and postanalytic (post-testing), as appropriate.

(b) Safety precautions must be established, posted, and observed to ensure protection from physical, chemical, biochemical and electrical hazards and biohazardous materials.

§ 493.1205  Standard; Test methods, equipment, instrumentation, reagents, materials, and supplies.

The laboratory must utilize test methods, equipment, instrumentation, reagents, materials, and supplies that provide accurate and reliable test results and test reports.

(a) Test methodologies and equipment must be selected and testing performed in a manner that provides test results within the laboratory’s stated performance specifications for each test method as determined under §493.1213.

(b) The laboratory must have appropriate and sufficient equipment, instruments, reagents, materials, and supplies for the type and volume of testing performed and for the maintenance of quality during the preanalytic, analytic, and postanalytic phases of testing.
(2) Components of reagent kits of different lot numbers are not interchanged unless otherwise specified by the manufacturer.

§ 493.1211 Standard; Procedure manual.

(a) A written procedure manual for the performance of all analytical methods used by the laboratory must be readily available and followed by laboratory personnel. Textbooks may be used as supplements to these written descriptions but may not be used in lieu of the laboratory's written procedures for testing or examining specimens.

(b) The procedure manual must include, when applicable to the test procedure:

(1) Requirements for specimen collection and processing, and criteria for specimen rejection;

(2) Procedures for microscopic examinations, including the detection of inadequately prepared slides;

(3) Step-by-step performance of the procedure, including test calculations and interpretation of results;

(4) Preparation of slides, solutions, calibrators, controls, reagents, stains and other materials used in testing;

(5) Calibration and calibration verification procedures;

(6) The reportable range for patient test results as established or verified in § 493.1213;

(7) Control procedures;

(8) Remedial action to be taken when calibration or control results fail to meet the laboratory's criteria for acceptability;

(9) Limitations in methodologies, including interfering substances;

(10) Reference range (normal values);

(11) Imminent life-threatening laboratory results or "panic values";

(12) Pertinent literature references;

(13) Appropriate criteria for specimen storage and preservation to ensure specimen integrity until testing is completed;

(14) The laboratory's system for reporting patient results including, when appropriate, the protocol for reporting panic values;

(15) Description of the course of action to be taken in the event that a test system becomes inoperable; and

(16) Criteria for the referral of specimens including procedures for specimen submission and handling as described in §493.1103.

(c) Manufacturers' package inserts or operator manuals may be used, when applicable, to meet the requirements of paragraphs (b)(1) through (b)(13) of this section. Any of the items under paragraphs (b)(1) through (b)(13) of this section not provided by the manufacturer must be provided by the laboratory.

(d) Procedures must be approved, signed, and dated by the director.

(e) Procedures must be re-approved, signed and dated if the directorship of the laboratory changes.

(f) Each change in a procedure must be approved, signed, and dated by the current director of the laboratory.

(g) The laboratory must maintain a copy of each procedure with the dates of initial use and discontinuance. These records must be retained for two years after a procedure has been discontinued.

§ 493.1213 Standard; Establishment and verification of method performance specifications.

Prior to reporting patient test results, the laboratory must verify or establish, for each method, the performance specifications for the following performance characteristics: accuracy; precision; analytical sensitivity and specificity, if applicable; the reportable range of patient test results; the reference range(s) (normal values); and any other applicable performance characteristic.

(a) The provisions of this section are not retroactive. Laboratories are not required to verify or establish performance specifications for any test method of moderate or high complexity in use prior to September 1, 1992.

(b)(1) Each laboratory that introduces a new procedure for patient testing using a device (instrument, kit, or test system) cleared by the FDA as meeting certain CLIA requirements for quality control, must demonstrate that, prior to reporting patient test results, it can obtain the performance specifications for accuracy, precision,
§ 493.1215 Standard; Equipment maintenance and function checks.

The laboratory must perform equipment maintenance and function checks that include electronic, mechanical and operational checks necessary for the proper test performance and test result reporting of equipment, instruments and test systems, to assure accurate and reliable test results and reports.

(a) Maintenance of equipment, instruments, and test systems. (1) For manufacturers’ equipment, instruments or test systems cleared by the FDA as meeting certain CLIA requirements for quality control, the laboratory must—

(i) Perform maintenance as defined by the manufacturer and with at least the frequency specified by the manufacturer; and

(ii) Document all maintenance performed.

(2) For methods or devices, as specified in either §493.1202(a) or (b) or §493.1203(a), the laboratory must—

(i) Establish a maintenance protocol that ensures equipment, instrument, and test system performance necessary for accurate and reliable test results and test result reporting;

(ii) Perform maintenance with at least the frequency specified in paragraph (a)(2)(i) of this section; and

(iii) Document all maintenance performed.

(b) Function checks of equipment, instruments, and test systems. (1) For manufacturers’ equipment, instruments, or test systems cleared by the FDA as meeting certain CLIA requirements for quality control, the laboratory must—

(i) Perform function checks as defined by the manufacturer and with at least the frequency specified by the manufacturer; and

(ii) Document all function checks performed.

(2) For methods or devices, as specified in either §493.1202(a) or (b) or §493.1203(a), the laboratory must—

(i) Define a function check protocol that ensures equipment, instrument, and test system performance necessary for accurate and reliable test results and test result reporting;

(ii) Perform function checks including background or baseline checks specified in paragraph (b)(2)(i) of this section. Function checks must be within the laboratory’s established limits before patient testing is conducted; and

(iii) Document all function checks performed.

§ 493.1217 Standard; Calibration and calibration verification procedures.

Calibration and calibration verification procedures are required to substantiate the continued accuracy of the test method throughout the laboratory’s reportable range for patient test results. Calibration is the process of testing and adjusting an instrument, kit,
or test system to provide a known relationship between the measurement response and the value of the substance that is being measured by the test procedure. Calibration verification is the assaying of calibration materials in the same manner as patient samples to confirm that the calibration of the instrument, kit, or test system has remained stable throughout the laboratory’s reportable range for patient test results. The reportable range of patient test results is the range of test result values over which the laboratory can establish or verify the accuracy of the instrument, kit or test system measurement response. Calibration and calibration verification must be performed and documented as required in this section unless otherwise specified in §§ 493.1223 through 493.1285.

(a) For laboratory test procedures that are performed using instruments, kits, or test systems that have been cleared by the FDA as meeting certain CLIA requirements for quality control, the laboratory must, at a minimum, follow the manufacturer’s instructions for calibration and calibration verification procedures using calibration materials specified by the manufacturer.

(b) For each method or device, as specified in either §493.1202 (a) or (b) or §493.1203(a), the laboratory must—

(1) Perform calibration procedures—
   (i) At a minimum, in accordance with manufacturer’s instructions, if provided, using calibration materials provided or specified, as appropriate, and with at least the frequency recommended by the manufacturer; and
   (ii) In accordance with criteria established by the laboratory, as required under §493.1213(b)(2)(i)—

   (A) Including the number, type, and concentration of calibration materials, acceptable limits for calibration, and the frequency of calibration; and
   (B) Using calibration materials appropriate for the methodology and, if possible, traceable to a reference method or reference material of known value; and

(2) Perform calibration verification procedures—

   (i) In accordance with the manufacturer’s calibration verification instructions when they meet or exceed the requirements specified in paragraph (b)(2)(ii) of this section; or
   (ii) In accordance with criteria established by the laboratory—

   (A) Including the number, type, and concentration of calibration materials, acceptable limits for calibration verification and frequency of calibration verification;
   (B) Using calibration materials appropriate for—

   (1) The methodology and, if possible, traceable to a reference method or reference material of known value; and
   (2) Verifying the laboratory’s established reportable range of patient test results, which must include at least a minimal (or zero) value, a mid-point value, and a maximum value at the upper limit of that range; and

   (C) At least once every six months and whenever any of the following occur:

   (1) A complete change of reagents for a procedure is introduced, unless the laboratory can demonstrate that changing reagent lot numbers does not affect the range used to report patient test results, and control values are not adversely affected by reagent lot number changes;

   NOTE: If reagents are obtained from a manufacturer and all of the reagents for a test are packaged together, the laboratory is not required to perform calibration verification for each package of reagents, provided the packages of reagents are received in the same shipment and contain the same lot number.

   (2) There is major preventive maintenance or replacement of critical parts that may influence test performance;

   (3) Controls reflect an unusual trend or shift or are outside of the laboratory’s acceptable limits and other means of assessing and correcting unacceptable control values have failed to identify and correct the problem; or

   (4) The laboratory’s established schedule for verifying the reportable range for patient test results requires more frequent calibration verification than specified in paragraphs
§493.1218 Standard; Control procedures.

Control procedures are performed on a routine basis to monitor the stability of the method or test system; control and calibration materials provide a means to indirectly assess the accuracy and precision of patient test results. Control procedures must be performed as defined in this section unless otherwise specified in §§493.1223 through 493.1285 of this subpart.

(a) For each device cleared by the FDA as meeting certain CLIA requirements for quality control, the laboratory must, at a minimum, follow the manufacturer's instructions for control procedures. In addition, the laboratory must meet the requirements under paragraphs (c) through (e) of this section and, as applicable, paragraph (f) of this section.

(b) For each device, as specified in either §493.1202(a) or (b) or §493.1203(a), the laboratory must evaluate instrument and reagent stability and operator variance in determining the number, type, and frequency of testing calibration or control materials and establish criteria for acceptability used to monitor test performance during a run of patient specimen(s). A run is an interval within which the accuracy and precision of a testing system is expected to be stable, but cannot be greater than 24 hours or less than the frequency recommended by the manufacturer. For each procedure, the laboratory must monitor test performance using calibration materials or control materials or a combination thereof.

(1) For qualitative tests, the laboratory must include a positive and negative control with each run of patient specimens.

(2) For quantitative tests, the laboratory must include at least two samples of different concentrations of either calibration materials, control materials, or a combination thereof with the frequency determined in §493.1218(b), but not less frequently than once each run of patient specimens.

(3) For electrophoretic determinations—

(i) At least one control sample must be used in each electrophoretic cell; and

(ii) The control sample must contain fractions representative of those routinely reported in patient specimens.

(4) Each day of use, the laboratory must evaluate the detection phase of direct antigen systems using an appropriate positive and negative control material (organism or antigen extract). When direct antigen systems include an extraction phase, the system must be checked each day of use using a positive organism.

(5) If calibration materials and control materials are not available, the laboratory must have an alternative mechanism to assure the validity of patient test results.

(c) Control samples must be tested in the same manner as patient specimens.

(d) When calibration or control materials are used, statistical parameters (e.g., mean and standard deviation) for each lot number of calibration material and each lot of control material must be determined through repetitive testing.

(1) The stated values of an assayed control material may be used as the target values provided the stated values correspond to the methodology and instrumentation employed by the laboratory and are verified by the laboratory.

(2) Statistical parameters for unassayed materials must be established over time by the laboratory through concurrent testing with calibration materials or control materials having previously determined statistical parameters.

(e) Control results must meet the laboratory's criteria for acceptability prior to reporting patient test results.

(f) Reagent and supply checks. (1) The laboratory must check each batch or shipment of reagents, discs, stains, antisera and identification systems (systems using two or more substrates) when prepared or opened for positive and negative reactivity, as well as graded reactivity if applicable.
§ 493.1221 Standard; Quality control records.

The laboratory must document and maintain records of all quality control activities specified in §§ 493.1202 through 493.1285 of this subpart and retain records for at least two years. Immunohematology quality control records must be maintained for a period of no less than five years. In addition, quality control records for blood and blood products must be maintained for a period not less than five years after processing records have been

§ 493.1219 Standard; Remedial actions.

Remedial action policies and procedures must be established by the laboratory and applied as necessary to maintain the laboratory’s operation for testing patient specimens in a manner that assures accurate and reliable patient test results and reports. The laboratory must document all remedial actions taken when—

(a) Test systems do not meet the laboratory’s established performance specifications, as determined in § 493.1213 of this section, which include but are not limited to—

(1) Equipment or methodologies that perform outside of established operating parameters or performance specifications;

(2) Patient test values that are outside of the laboratory’s reportable range of patient test results; and

(3) The determination that the laboratory’s reference range for a test procedure is inappropriate for the laboratory’s patient population.

(b) Results of control and calibration materials fail to meet the laboratory’s established criteria for acceptability. All patient test results obtained in the unacceptable test run or since the last acceptable test run must be evaluated to determine if patient test results have been adversely affected and the laboratory must take the remedial action necessary to ensure the reporting of accurate and reliable patient test results;

(c) The laboratory cannot report patient test results within its established time frames. The laboratory must determine, based on the urgency of the patient test(s) requested, the need to notify the appropriate individual of the delayed testing; and

(d) Errors in the reported patient test results are detected. The laboratory must—

(1) Promptly notify the authorized person ordering or individual utilizing the test results of reporting errors;

(2) Issue corrected reports promptly to the authorized person ordering the test or the individual utilizing the test results; and

(3) Maintain exact duplicates of the original report as well as the corrected report for two years.
§ 493.1223 Condition: Quality control—specialties and subspecialties for tests of moderate or high complexity, or both.

The laboratory must establish and follow written quality control procedures for monitoring and evaluating the quality of the analytical testing process of each method to assure the accuracy and reliability of patient test results and reports. Except as specified in §493.1202(c), the laboratory must meet the applicable general requirements specified in §§493.1201 through 493.1221. In addition, the laboratory must meet the applicable requirements of §§493.1225 through 493.1285 unless an alternative procedure specified in the manufacturer’s protocol has been cleared by the Food and Drug Administration (FDA) as meeting certain CLIA requirements for quality control or HCFA approves an equivalent procedure specified in appendix C of the State Operations Manual (HCFA Pub. 7). Failure to meet any of the applicable conditions in §§493.1225 through 493.1285 will result in intermediate sanctions, loss of Medicare or Medicaid approval, and/or revocation of CLIA certification for the entire specialty or subspecialty to which the condition applies, in accordance with subpart R of this part.

[58 FR 5232, Jan. 19, 1993]

§ 493.1225 Condition: Microbiology.

The laboratory must meet the applicable quality control requirements in §§493.1201 through 493.1221 and in §§493.1227 through 493.1235 of this subpart for the subspecialties for which it is certified under the specialty of microbiology.

§ 493.1227 Condition: Bacteriology.

To meet the quality control requirements for bacteriology, the laboratory must comply with the applicable requirements in §§493.1201 through 493.1221 and with paragraphs (a) through (c) of this section. All quality control activities must be documented.

(a) The laboratory must check positive and negative reactivity with control organisms—

(1) Each day of use for catalase, coagulase, beta-lactamase, and oxidase reagents and DNA probes;

(2) Each week of use for Gram and acid-fast stains, bacitracin, optochin, ONPG, X, and V discs or strips; and

(3) Each month of use for antisera.

(b) Each day tests are performed, the laboratory must use the appropriate control organism(s) to check the procedure.

§ 493.1229 Condition: Mycobacteriology.

To meet the quality control requirements for mycobacteriology, the laboratory must comply with the applicable requirements in §§493.1201 through 493.1221 of this subpart and with paragraphs (a) through (d) of this section. All quality control activities must be documented.

(a) Each day of use, the laboratory must check the iron uptake test with at least one acid-fast organism that produces a positive reaction and with an organism that produces a negative reaction and check all other reagents or test procedures used for mycobacteria identification with at least one acid-fast organism that produces a positive reaction.

(b) The laboratory must check fluorochrome acid-fast stains for positive and negative reactivity each week of use.

(c) The laboratory must check acid-fast stains each week of use with an acid-fast organism that produces a positive reaction.

(d) For susceptibility tests performed on Mycobacterium tuberculosis isolates,
the laboratory must check the procedure each week of use with a strain of Mycobacterium tuberculosis susceptible to all antimycobacterial agents tested.

§ 493.1231 Condition: Mycology.

To meet the quality control requirements for mycology, the laboratory must comply with the applicable requirements in §§ 493.1201 through 493.1221 of this subpart and with paragraphs (a) through (d) of this section. All quality control activities must be documented.

(a) Each day of use, the laboratory using the auxanographic medium for nitrate assimilation must check the nitrate reagent with a peptone control.

(b) Each week of use, the laboratory must check all reagents used with biochemical tests and other test procedures for mycological identification with an organism that produces a positive reaction.

(c) Each week of use, the laboratory must check acid-fast stains for positive and negative reactivity.

(d) For susceptibility tests, the laboratory must test each drug each day of use with at least one control strain that is susceptible to the drug. The laboratory must establish control limits. Criteria for acceptable control results must be met prior to reporting patient results.

§ 493.1233 Condition: Parasitology.

To meet the quality control requirements for parasitology, the laboratory must comply with the applicable requirements in §§ 493.1201 through 493.1221 of this subpart and with paragraphs (a) through (c) of this section. All quality control activities must be documented.

(a) The laboratory must have available a reference collection of slides or photographs, and, if available, gross specimens for identification of parasites and use these references in the laboratory for appropriate comparison with diagnostic specimens.

(b) The laboratory must calibrate and use the calibrated ocular micrometer for determining the size of ova and parasites, if size is a critical parameter.

(c) Each month of use, the laboratory must check permanent stains using a fecal sample control that will demonstrate staining characteristics.

§ 493.1235 Condition: Virology.

To meet the quality control requirements for virology, the laboratory must comply with the applicable requirements in §§ 493.1201 through 493.1221 of this subpart and with paragraphs (a) through (c) of this section. All quality control activities must be documented.

(a) The laboratory must have available host systems for the isolation of viruses and test methods for the identification of viruses that cover the entire range of viruses that are etiologically related to clinical diseases for which services are offered.

(b) The laboratory must maintain records that reflect the systems used and the reactions observed.

(c) In tests for the identification of viruses, the laboratory must simultaneously culture uninoculated cells or cell substrate controls as a negative control to detect erroneous identification results.

§ 493.1237 Condition: Diagnostic immunology.

The laboratory must meet the applicable quality control requirements in §§ 493.1201 through 493.1221 and §§ 493.1233 through 493.1241 of this subpart for the subspecialties for which it is certified under the specialty of diagnostic immunology.

§ 493.1239 Condition: Syphilis serology.

To meet the quality control requirements for syphilis serology, the laboratory must comply with the applicable requirements in §§ 493.1201 through 493.1221 of this subpart and with paragraphs (a) through (e) of this section. All quality control activities must be documented.

(a) For laboratories performing syphilis testing, the equipment, glassware, reagents, controls, and techniques for tests for syphilis must conform to manufacturers' specifications.

(b) The laboratory must run serologic tests on patient specimens concurrently with a positive serum control of known titer or controls of graded reactivity plus a negative control.
§ 493.1241 Condition: General immunology.

To meet the quality control requirements for general immunology, the laboratory must comply with the applicable requirements in §§ 493.1201 through 493.1221 of this subpart and with paragraphs (a) through (d) of this section. All quality control activities must be documented.

(a) The laboratory must run serologic tests on patient specimens concurrently with a positive serum control of known titer or controls of graded reactivity, if applicable, plus a negative control.

(b) The laboratory must employ controls that evaluate all phases of the test system (antigens, complement, erythrocyte indicator systems, etc.) to ensure reactivity and uniform dosages when positive and negative controls alone are not sufficient.

(c) The laboratory may not report test results unless the predetermined reactivity pattern of the controls is observed.

(d) All facilities manufacturing blood and blood products for transfusion or serving as referral laboratories for these facilities must meet the syphilis serology testing requirements of 21 CFR 640.5(a).

§ 493.1243 Condition: Chemistry.

The laboratory must meet the applicable quality control requirements in §§ 493.1201 through 493.1221 and §§ 493.1245 through 493.1249 of this subpart for the subspecialties for which it is certified under the specialty of chemistry.

§ 493.1245 Condition: Routine chemistry.

To meet the quality control requirements for routine chemistry, the laboratory must comply with the applicable requirements in §§ 493.1201 through 493.1221. All quality control activities must be documented. In addition, for blood gas analyses, the laboratory must—

(a) Calibrate or verify calibration according to the manufacturer's specifications and with at least the frequency recommended by the manufacturer;

(b) Test one sample of control material each eight hours of testing;

(c) Use a combination of calibrators and control materials that include both low and high values on each day of testing; and

(d) Include one sample of calibration material or control material each time patients are tested unless automated instrumentation internally verifies calibration at least every thirty minutes.

§ 493.1247 Condition: Endocrinology.

To meet the quality control requirements for endocrinology, the laboratory must comply with the applicable requirements contained in §§ 493.1201 through 493.1221 of this subpart. All quality control activities must be documented.

§ 493.1249 Condition: Toxicology.

To meet the quality control requirements for toxicology, the laboratory must comply with the applicable requirements in §§ 493.1201 through 493.1221 of this subpart. All quality control activities must be documented. In addition, for drug abuse screening using thin layer chromatography—

(a) Each plate must be spotted with at least one sample of calibration material containing all drug groups identified by thin layer chromatography which the laboratory reports; and

(b) At least one control sample must be included in each chamber, and the control sample must be processed through each step of patient testing, including extraction procedures.
§ 493.1251 Condition: Urinalysis.

Except for those tests categorized as waived, to meet the quality control requirements for urinalysis, the laboratory must comply with the applicable requirements in §§493.1201 through 493.1221.

[58 FR 5232, Jan. 19, 1993]

§ 493.1253 Condition: Hematology.

To meet the quality control requirements for hematology, the laboratory must comply with the applicable requirements in §§493.1201 through 493.1221 of this subpart and with paragraphs (a) through (d) of this section. All quality control activities must be documented.

(a) Cell counts performed manually using a hemocytometer must be tested in duplicate. One control is required for each eight hours of operation.

(b) For non-manual hematology testing systems, excluding coagulation, the laboratory must include two levels of controls each eight hours of operation.

(c) For all non-manual coagulation testing systems, the laboratory must include two levels of control each eight hours of operation and each time a change in reagents occurs.

(d) For manual coagulation tests—

(1) Each individual performing tests must test two levels of controls before testing patient samples and each time a change in reagents occurs; and

(2) Patient and control specimens must be tested in duplicate.


§ 493.1255 Condition: Pathology.

The laboratory must meet the applicable quality control requirements in §§493.1201 through 493.1221 and §§493.1257 through 493.1261 of this subpart for the subspecialties for which it is certified under the specialty of pathology. All quality control activities must be documented.

§ 493.1257 Condition: Cytology.

To meet the quality control requirements for cytology, the laboratory must comply with the applicable requirements in §§493.1201 through 493.1221 of this subpart and paragraphs (a) through (g) of this section.

(a) The laboratory must assure that—

(1) All gynecologic smears are stained using a Papanicolaou or modified Papanicolaou staining method;

(2) Effective measures are taken to prevent cross-contamination between gynecologic and nongynecologic specimens during the staining process;

(3) Nongynecologic specimens that have a high potential for cross-contamination are stained separately from other nongynecologic specimens, and the stains are filtered or changed following staining;

(4) Diagnostic interpretations are not reported on unsatisfactory smears; and

(5) All cytology slide preparations are evaluated on the premises of a laboratory certified to conduct testing in the subspecialty of cytology.

(b) The laboratory is responsible for ensuring that—

(1) Each individual engaged in the evaluation of cytology preparations by nonautomated microscopic technique examines no more than 100 slides (one patient per slide, gynecologic or nongynecologic, or both) in a 24 hour period, irrespective of the site or laboratory. This limit represents an absolute maximum number of slides and is not to be employed as a performance target for each individual. Previously examined negative, reactive, reparative, atypical, premalignant or malignant gynecologic cases as defined in paragraph (c)(1) of this section, previously examined nongynecologic cytology preparations, and tissues pathology slides examined by a technical supervisor qualified under §493.1449(b) or (k) are not included in the 100 slide limit. (For this section, all references to technical supervisor refer to individuals qualified under §§493.1449(b) and (k).)

(2) For purposes of workload calculations, each slide preparation (gynecologic and nongynecologic) made using automated, semi-automated, or other liquid-based slide preparatory techniques which result in cell dispersion over one-half or less of the total available slide area and which is examined by nonautomated microscopic technique counts as one-half slide.

(3) Records are maintained of the total number of slides examined by...
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each individual during each 24 hour period, irrespective of the site or laboratory, and the number of hours each individual spends examining slides in the 24 hour period;

(i) The maximum number of 100 slides described in paragraph (b)(1) of this section is examined in no less than an 8 hour workday;

(ii) For the purposes of establishing workload limits for individuals examining slides by nonautomated microscopic technique on other than an 8 hour workday basis (includes full-time employees with duties other than slide examination and part-time employees), a period of 8 hours must be used to pro-rate the number of slides that may be examined. Use the formula—

\[
\text{No. of hours examining slides} \times \frac{100}{8}
\]

to determine maximum slide volume to be examined.

(c) The individual qualified under §§493.1449 (b) or (k) who provides technical supervision of cytology must ensure that—

(1) All gynecologic smears interpreted to be showing reactive or reparative changes, atypical squamous or glandular cells of undetermined significance, or to be in the premalignant (dysplasia, cervical intraepithelial neoplasia or all squamous intraepithelial lesions including human papillomavirus-associated changes) or malignant category are confirmed by a technical supervisor in cytology. The report must be signed to reflect the review or, if a computer report is generated with signature, it must reflect an electronic signature authorized by the technical supervisor in cytology;

(2) All nongynecologic cytologic preparations are reviewed by the technical supervisor in cytology. The report must be signed to reflect the review or, if a computer report is generated with signature, it must reflect an electronic signature authorized by the technical supervisor;

(3) The slide examination performance of each cytotechnologist is evaluated and documented, including performance evaluation through the re-examination of normal and negative cases and feedback on the reactive, reparative, atypical, malignant or premalignant cases as defined in paragraph (c)(1) of this section; and

(4) A maximum number of slides, not to exceed the maximum workload limit described in paragraph (b) of this section is established by the technical supervisor for each individual examining slide preparations by nonautomated microscopic technique.

(i) The actual workload limit must be documented for each individual and established in accordance with the individual’s capability based on the performance evaluation as described in paragraph (c)(3) of this section.

(ii) Records are available to document that each individual’s workload limit is reassessed at least every 6 months and adjusted when necessary.

(d) The laboratory must establish and follow a program designed to detect errors in the performance of cytologic examinations and the reporting of results.

(1) The laboratory must establish a program that includes a review of slides from at least 10 percent of the gynecologic cases interpreted to be negative for reactive, reparative, atypical, premalignant or malignant conditions as defined in paragraph (c)(1) of this section that are examined by each individual not qualified under §§493.1449 (b) or (k). This review must be done by a technical supervisor in cytology, a cytology general supervisor qualified under §493.1469, or a cytotechnologist qualified under §493.1483 who has the experience specified in §493.1469(b)(2).

(i) The review must include negative cases selected at random from the total caseload and from patients or groups of patients that are identified as having a high probability of developing cervical cancer, based on available patient information;

(ii) Records of initial examinations and rescreening results must be available; and

(iii) The review must be completed before reporting patient results on those cases selected.

(2) The laboratory must compare clinical information, when available, with cytology reports and must compare all malignant and premalignant
§ 493.1259 Condition: Histopathology.

To meet the quality control requirements for histopathology, a laboratory must comply with the applicable requirements in §§ 493.1201 through 493.1221 of this subpart and paragraphs (a) through (e) of this section. All quality control activities must be documented.

(a) A control slide of known reactivity must be included with each slide or group of slides for differential or special stains. Reaction(s) of the control slide with each special stain must be documented.

(b) The laboratory must retain stained slides at least ten years from the date of examination and retain specimen blocks at least two years from the date of examination.

(c) The laboratory must retain remnants of tissue specimens in a manner that assures proper preservation of the tissue specimens until the portions submitted for microscopic examination have been examined and a diagnosis made by an individual qualified under §§ 493.1449(b) or 493.1449(1)(I) of this part. In addition, an individual who meets the requirements of §§ 493.1449(b), 493.1449(1)(I) or 493.1449(1)(II), may examine and provide reports for specimens for skin pathology; an individual meeting the requirements of §§ 493.1449(b) or 493.1449(1)(I) may examine and provide reports for ophthalmic pathology; an individual meeting the requirements of §§ 493.1449(b) or 493.1449(m) may examine and provide reports for oral pathology specimens.

(d) All tissue pathology reports must be signed by an individual qualified as specified in paragraph (c) of this section. If a computer report is generated
§ 493.1261 Condition: Oral pathology.

To meet the quality control requirements for oral pathology, the laboratory must comply with the applicable requirements in §§ 493.1201 through 493.1221 and § 493.1259 of this subpart. All quality control activities must be documented.

§ 493.1263 Condition: Radiobioassay.

To meet quality control requirements for radiobioassay, the laboratory must comply with the applicable requirements of §§ 493.1201 through 493.1221 of this subpart. All quality control activities must be documented.

§ 493.1265 Condition: Histocompatibility.

In addition to meeting the applicable requirements for general quality control in §§ 493.1201 through 493.1221, for quality control for general immunology in § 493.1241 of this subpart and for immunohematology in § 493.1269 of this subpart, the laboratory must comply with the applicable requirements in paragraphs (a) through (d) of this section. All quality control activities must be documented.

(a) For renal allotransplantation, the laboratory must meet the following requirements:

(1) The laboratory must have available and follow criteria for—

(i) Selecting appropriate patient serum samples for crossmatching;
(ii) The technique used in crossmatching;
(iii) Preparation of donor lymphocytes for crossmatching; and
(iv) Reporting crossmatch results;
(2) The laboratory must—

(i) Have available results of final crossmatches before an organ or tissue is transplanted; and
(ii) Make a reasonable attempt and document efforts to have available serum specimens for all potential transplant recipients at initial typing, for periodic screening, for pre-transplantation crossmatch and following sensitizing events, such as transfusion and transplant loss;
(3) The laboratory’s storage and maintenance of both recipient sera and reagents must—

(i) Be at an acceptable temperature range for sera and components;
(ii) Use a temperature alarm system and have an emergency plan for alternate storage; and
(iii) Ensure that all specimens are properly identified and easily retrievable;
(4) The laboratory’s reagent typing sera inventory (applicable only to locally constructed trays) must indicate source, bleeding date and identification number, and volume remaining;
(5) The laboratory must properly label and store cells, complement, buffer, dyes, etc.;
(6) The laboratory must—

(i) HLA type all potential transplant recipients;
(ii) Type cells from organ donors referred to the laboratory; and
(iii) Have available and follow a policy that establishes when antigen redefinition and retyping are required;
(7) The laboratory must have available and follow criteria for—

(i) The preparation of lymphocytes for HLA-A, B and DR typing;
(ii) Selecting typing reagents, whether locally or commercially prepared;
(iii) The assignment of HLA antigens; and
(iv) Assuring that reagents used for typing recipients and donors are adequate to define all major and International Workshop HLA-A, B and DR specificities for which reagents are readily available;
(8) The laboratory must—

(i) Screen potential transplant recipient sera for preformed HLA-A and B antibodies with a suitable lymphocyte panel on sera collected;
(A) At the time of the recipient’s initial HLA typing; and
(B) Thereafter, following sensitizing events and upon request; and
(ii) Use a suitable cell panel for screening patient sera (antibody screen), a screen that contains all the major HLA specificities and common splits—
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(A) If the laboratory does not use commercial panels, it must maintain a list of individuals for fresh panel bleeding; and

(B) If the laboratory uses frozen panels, it must have a suitable storage system;

(9) The laboratory must check—

(i) Each typing tray using—

(A) Positive control sera;

(B) Negative control sera; and

(C) Positive controls for specific cell types when applicable (i.e., T cells, B cells, and monocytes); and

(ii) Each compatibility test (i.e. mixed lymphocyte culture test, homozygous typing cells or DNA analysis) and typing for disease-associated antigens using controls to monitor the test components and each phase of the test system to ensure an acceptable performance level;

(10) Compatibility testing for cellularly-defined antigens must utilize techniques such as the mixed lymphocyte culture test, homozygous typing cells or DNA analysis;

(11) If the laboratory reports the recipient’s or donor’s, or both, ABO blood group and D(Rho) typing, the testing must be performed in accordance with § 493.1269 of this subpart;

(12) If the laboratory utilizes immunologic reagents (such as antibodies or complement) to remove contaminating cells during the isolation of lymphocytes or lymphocyte subsets, the efficacy of the methods must be verified with appropriate quality control procedures;

(13) At least once each month, the laboratory must have each individual performing tests evaluate a previously tested specimen as an unknown to verify his or her ability to reproduce test results. Records of the results for each individual must be maintained; and

(14) The laboratory must participate in at least one national or regional cell exchange program, if available, or develop an exchange system with another laboratory in order to validate inter-laboratory reproducibility.

(b) If the laboratory performs histocompatibility testing for—

(1) Transfusions and other non-renal transplantation, all the requirements specified in this section, as applicable, except for the performance of mixed lymphocyte cultures, must be met;

(2) Bone marrow transplantation, all the requirements specified in this section, including the performance of mixed lymphocyte cultures or other augmented testing to evaluate class II compatibility, must be met; and

(3) Non-renal solid organ transplantation, the results of final crossmatches must be available before transplantation when the recipient has demonstrated presensitization by prior serum screening except for emergency situations. The laboratory must document the circumstances, if known, under which emergency transplants are performed, and records must reflect any information concerning the transplant provided to the laboratory by the patient’s physician.

(c) Laboratories performing HLA typing for disease-associated studies must meet all the requirements specified in this section except for the performance of mixed lymphocyte cultures, antibody screening and crossmatching.

(d) For laboratories performing organ donor HIV testing the requirements of § 493.1241 of this subpart for the transfusion of blood and blood products must be met.

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**Condition:** Immunohematology.

To meet the quality control requirements for immunohematology, the laboratory must comply with the applicable requirements in §§ 493.1201 through 493.1221 of this subpart and with paragraphs (a) through (d) of this section. All quality control activities must be documented.

(a) The laboratory must perform ABO group and D(Rho) typing, unexpected antibody detection, antibody identification and compatibility testing in accordance with manufacturer’s instructions, if provided, and as applicable, with 21 CFR part 606 (with the exception of 21 CFR 606.20a, Personnel) and 21 CFR part 640 et seq.

(b) The laboratory must perform ABO group by concurrently testing unknown red cells with anti-A and anti-B grouping reagents. For confirmation of ABO group, the unknown serum must be tested with known A1 and B red cells.

(c) The laboratory must determine the D(Rho) type by testing unknown red cells with anti-D (anti-Rho) blood grouping reagent.

(d) If required in the manufacturer’s package insert for anti-D reagents, the laboratory must employ a control system capable of detecting false positive D(Rho) test results.

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**Condition:** Transfusion services and bloodbanking.

If a facility provides services for the transfusion of blood and blood products, the facility must be under the adequate control and technical supervision of the pathologist or other doctor of medicine or osteopathy meeting the qualifications in subpart M for technical supervision in immunohematology. The facility must ensure that there are facilities for procurement, safekeeping and transfusion of blood and blood products and that blood and blood products must be available to meet the needs of the physicians responsible for the diagnosis, management, and treatment of patients. The facility meets this condition by complying with the standards in §§ 493.1273 through 493.1285.

[58 FR 5233, Jan. 19, 1993]

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**Standard:** Immunohematological collection, processing, dating periods, labeling and distribution of blood and blood products.

In addition to the requirements in paragraphs (a) through (d) of this section, the facility must also meet the applicable quality control requirements in §§ 493.1201 through 493.1221 of this part.

(a) Blood and blood product collection, processing and distribution must comply with 21 CFR part 640 and 21 CFR part 606, and the testing laboratory must meet the applicable requirements of part 493.

(b) Dating periods for blood and blood products must conform to 21 CFR 630.53.

(c) Labeling of blood and blood products must conform to 21 CFR part 606, subpart G.

(d) Policies to ensure positive identification of a blood or blood product recipient must be established, documented, and followed.

§ 493.1275

**Standard:** Blood and blood products storage facilities.

(a) The blood and blood products must be stored under appropriate conditions, which include an adequate temperature alarm system that is regularly inspected.
§ 493.1355 Condition: Laboratories performing PPM procedures; laboratory director.

The laboratory must have a director who meets the qualification requirements of §493.1357 and provides overall management and direction in accordance with §493.1359.
§ 493.1357 Standard; laboratory director qualifications.

The laboratory director must be qualified to manage and direct the laboratory personnel and the performance of PPM procedures as specified in §493.19(c) and must be eligible to be an operator of a laboratory within the requirements of subpart R of this part.

(a) The laboratory director must possess a current license as a laboratory director issued by the State in which the laboratory is located, if the licensing is required.

(b) The laboratory director must meet one of the following requirements:

1. Be a physician, as defined in §493.2.
2. Be a midlevel practitioner, as defined in §493.2, authorized by a State to practice independently in the State in which the laboratory is located.
3. Be a dentist, as defined in §493.2.

§ 493.1359 Standard; PPM laboratory director responsibilities.

The laboratory director is responsible for the overall operation and administration of the laboratory, including the prompt, accurate, and proficient reporting of test results. The laboratory director must—

(a) Direct no more than five laboratories; and

(b) Ensure that any procedure listed under §493.19(c)—

1. Is personally performed by an individual who meets the qualification requirements in §493.1363; and

2. Is performed in accordance with applicable requirements in subparts H, J, K, M, and P of this part.

§ 493.1361 Condition: Laboratories performing PPM procedures; testing personnel.

The laboratory must have a sufficient number of individuals who meet the qualification requirements of §493.1363 to perform the functions specified in §493.1365 for the volume and complexity of testing performed.

§ 493.1363 Standard; PPM testing personnel qualifications.

Each individual performing PPM procedures must—

(a) Possess a current license issued by the State in which the laboratory is located if the licensing is required; and

(b) Meet one of the following requirements:

1. Be a physician, as defined in §493.2.
2. Be a midlevel practitioner, as defined in §493.2, under the supervision of a physician or in independent practice if authorized by the State in which the laboratory is located.
3. Be a dentist as defined in §493.2 of this part.

§ 493.1365 Standard; PPM testing personnel responsibilities.

The testing personnel are responsible for specimen processing, test performance, and for reporting test results. Any PPM procedure must be—

(a) Personally performed by one of the following practitioners:

1. A physician during the patient's visit on a specimen obtained from his or her own patient or from a patient of a group medical practice of which the physician is a member or employee.
2. A midlevel practitioner, under the supervision of a physician or in independent practice if authorized by the State in which the laboratory is located, during the patient's visit on a specimen obtained from his or her own patient or from the patient of a clinic, group medical practice, or other health care provider, in which the midlevel practitioner is a member or an employee.
3. A dentist during the patient's visit on a specimen obtained from his or her own patient or from a patient of a group dental practice of which the dentist is a member or an employee; and

(b) Performed using a microscope limited to a brightfield or a phase/contrast microscope.
§493.1405 Standard; Laboratory director qualifications.

The laboratory director must be qualified to manage and direct the laboratory personnel and the performance of moderate complexity tests and must be eligible to be an operator of a laboratory within the requirements of subpart R of this part.

(a) The laboratory director must possess a current license as a laboratory director issued by the State in which the laboratory is located, if such licensing is required; and

(b) The laboratory director must:

(1)(i) Be a doctor of medicine or doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(ii) Be certified in anatomic or clinical pathology, or both, by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(2)(i) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and

(ii) Have had laboratory training or experience consisting of:

(A) At least one year directing or supervising non-waived laboratory testing; or

(B) Beginning September 1, 1993, have at least 20 continuing medical education credit hours in laboratory practice commensurate with the director responsibilities defined in §493.1407; or

(C) Laboratory training equivalent to paragraph (b)(2)(iii)(B) of this section obtained during medical residency. (For example, physicians certified either in hematology or hematology and medical oncology by the American Board of Internal Medicine); or

(3) Hold an earned doctoral degree in a chemical, physical, biological, or clinical laboratory science from an accredited institution; and

(i) Be certified by the American Board of Medical Microbiology, the American Board of Clinical Chemistry, the American Board of Bioanalysis, or the American Board of Medical Laboratory Immunology; or

(ii) Have had at least one year experience directing or supervising non-waived laboratory testing;

(4)(i) Have earned a master's degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution;

(ii) Have at least one year of laboratory training or experience, or both in non-waived testing; and

(iii) In addition, have at least one year of supervisory laboratory experience in non-waived testing; or

(5)(i) Have earned a bachelor's degree in a chemical, physical, or biological science or medical technology from an accredited institution;

(ii) Have at least 2 years of laboratory training or experience, or both in non-waived testing; and

(iii) In addition, have at least 2 years of supervisory laboratory experience in non-waived testing;

(6) Be serving as a laboratory director and must have previously qualified or could have qualified as a laboratory director under §493.1406; or

(7) On or before February 28, 1992, qualified under State law to direct a laboratory in the State in which the laboratory is located.


§493.1406 Standard; Laboratory director qualifications on or before February 28, 1992.

The laboratory director must be qualified to manage and direct the laboratory personnel and test performance.

(a) The laboratory director must possess a current license as a laboratory director issued by the State, if such licensing exists; and

(b) The laboratory director must:

(1) Be a physician certified in anatomic or clinical pathology (or both) by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification;

(2) Be a physician who:
§ 493.1407  Standard; Laboratory director responsibilities.

The laboratory director is responsible for the overall operation and administration of the laboratory, including the employment of personnel who are competent to perform test procedures, and record and report test results promptly, accurately, and proficiently and for assuring compliance with the applicable regulations.

(a) The laboratory director, if qualified, may perform the duties of the technical consultant, clinical consultant, and testing personnel, or delegate these responsibilities to personnel meeting the qualifications of §§ 493.1409, 493.1415, and 493.1421, respectively.

(b) If the laboratory director reapporions performance of his or her responsibilities, he or she remains responsible for ensuring that all duties are properly performed.

(c) The laboratory director must be accessible to the laboratory to provide onsite, telephone or electronic consultation as needed.
(d) Each individual may direct no more than five laboratories.

(e) The laboratory director must—

(1) Ensure that testing systems developed and used for each of the tests performed in the laboratory provide quality laboratory services for all aspects of test performance, which includes the preanalytic, analytic, and postanalytic phases of testing;

(2) Ensure that the physical plant and environmental conditions of the laboratory are appropriate for the testing performed and provide a safe environment in which employees are protected from physical, chemical, and biological hazards;

(3) Ensure that—

(i) The test methodologies selected have the capability of providing the quality of results required for patient care;

(ii) Verification procedures used are adequate to determine the accuracy, precision, and other pertinent performance characteristics of the method; and

(iii) Laboratory personnel are performing the test methods as required for accurate and reliable results;

(4) Ensure that the laboratory is enrolled in an HHS approved proficiency testing program for the testing performed and that—

(i) The proficiency testing samples are tested as required under subpart H of this part;

(ii) The results are returned within the timeframes established by the proficiency testing program;

(iii) All proficiency testing reports received are reviewed by the appropriate staff to evaluate the laboratory’s performance and to identify any problems that require corrective action; and

(iv) An approved corrective action plan is followed when any proficiency testing results are found to be unacceptable or unsatisfactory;

(5) Ensure that the quality control and quality assurance programs are established and maintained to assure the quality of laboratory services provided and to identify failures in quality as they occur;

(6) Ensure the establishment and maintenance of acceptable levels of analytical performance for each test system;

(7) Ensure that all necessary remedial actions are taken and documented whenever significant deviations from the laboratory’s established performance specifications are identified, and that patient test results are reported only when the system is functioning properly;

(8) Ensure that reports of test results include pertinent information required for interpretation;

(9) Ensure that consultation is available to the laboratory’s clients on matters relating to the quality of the test results reported and their interpretation concerning specific patient conditions;

(10) Employ a sufficient number of laboratory personnel with the appropriate education and either experience or training to provide appropriate consultation, properly supervise and accurately perform tests and report test results in accordance with the personnel responsibilities described in this subpart;

(11) Ensure that prior to testing patients’ specimens, all personnel have the appropriate education and experience, receive the appropriate training for the type and complexity of the services offered, and have demonstrated that they can perform all testing operations reliably to provide and report accurate results;

(12) Ensure that policies and procedures are established for monitoring individuals who conduct preanalytical, analytical, and postanalytical phases of testing to assure that they are competent and maintain their competency to process specimens, perform test procedures and report test results promptly and proficiently, and whenever necessary, identify needs for remedial training or continuing education to improve skills;

(13) Ensure that an approved procedure manual is available to all personnel responsible for any aspect of the testing process; and

(14) Specify, in writing, the responsibilities and duties of each consultant and each person, engaged in the performance of the preanalytic, analytic, and postanalytic phases of testing, that identifies which examinations and procedures each individual is authorized to perform, whether supervision is
§ 493.1409  Condition: Laboratories performing moderate complexity testing; technical consultant.

The laboratory must have a technical consultant who meets the qualification requirements of §493.1411 of this subpart and provides technical oversight in accordance with §493.1413 of this subpart.

§ 493.1411 Standard; Technical consultant qualifications.

The laboratory must employ one or more individuals who are qualified by education and either training or experience to provide technical consultation for each of the specialties and subspecialties of service in which the laboratory performs moderate complexity tests or procedures. The director of a laboratory performing moderate complexity testing may function as the technical consultant provided he or she meets the qualifications specified in this section.

(a) The technical consultant must possess a current license issued by the State in which the laboratory is located, if such licensing is required.

(b) The technical consultant must—

(1) (i) Be a doctor of medicine or doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(ii) Be certified in anatomic or clinical pathology, or both, by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(2) (i) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and

(ii) Have at least one year of laboratory training or experience, or both in non-waived testing, in the designated specialty or subspecialty areas of service for which the technical consultant is responsible; or

(3)(i) Hold an earned doctoral or master’s degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution; and

(ii) Have at least one year of laboratory training or experience, or both in non-waived testing, in the designated specialty or subspecialty areas of service for which the technical consultant is responsible; or

(4)(i) Have earned a bachelor’s degree in a chemical, physical or biological science or medical technology from an accredited institution; and

(ii) Have at least 2 years of laboratory training or experience, or both in non-waived testing, in the designated specialty or subspecialty areas of service for which the technical consultant is responsible.

NOTE: The technical consultant requirements for “laboratory training or experience, or both” in each specialty or subspecialty may be acquired concurrently in more than one of the specialties or subspecialties of service, excluding waived tests. For example, an individual who has a bachelor’s degree in biology and additionally has documentation of 2 years of work experience performing tests of moderate complexity in all specialties and subspecialties of service, would be qualified as a technical consultant in a laboratory performing moderate complexity testing in all specialties and subspecialties of service.


§ 493.1413 Standard; Technical consultant responsibilities.

The technical consultant is responsible for the technical and scientific oversight of the laboratory. The technical consultant is not required to be onsite at all times testing is performed; however, he or she must be available to the laboratory on an as needed basis to provide consultation, as specified in paragraph (a) of this section.

(a) The technical consultant must be accessible to the laboratory to provide on-site, telephone, or electronic consultation; and
(b) The technical consultant is responsible for—
(1) Selection of test methodology appropriate for the clinical use of the test results;
(2) Verification of the test procedures performed and the establishment of the laboratory’s test performance characteristics, including the precision and accuracy of each test and test system;
(3) Enrollment and participation in an HHS approved proficiency testing program commensurate with the services offered;
(4) Establishing a quality control program appropriate for the testing performed and establishing the parameters for acceptable levels of analytic performance and ensuring that these levels are maintained throughout the entire testing process from the initial receipt of the specimen, through sample analysis and reporting of test results;
(5) Resolving technical problems and ensuring that remedial actions are taken whenever test systems deviate from the laboratory’s established performance specifications;
(6) Ensuring that patient test results are not reported until all corrective actions have been taken and the test system is functioning properly;
(7) Identifying training needs and assuring that each individual performing tests receives regular in-service training and education appropriate for the type and complexity of the laboratory services performed;
(8) Evaluating the competency of all testing personnel and assuring that the staff maintain their competency to perform test procedures and report test results promptly, accurately and proficiently. The procedures for evaluation of the competency of the staff must include, but are not limited to—
(i) Direct observations of routine patient test performance, including patient preparation, if applicable, specimen handling, processing and testing;
(ii) Monitoring the recording and reporting of test results;
(iii) Review of intermediate test results or worksheets, quality control records, proficiency testing results, and preventive maintenance records;
(iv) Direct observation of performance of instrument maintenance and function checks;
(v) Assessment of test performance through testing previously analyzed specimens, internal blind testing samples or external proficiency testing samples; and
(vi) Assessment of problem solving skills; and
(9) Evaluating and documenting the performance of individuals responsible for moderate complexity testing at least semiannually during the first year the individual tests patient specimens. Thereafter, evaluations must be performed at least annually unless test methodology or instrumentation changes, in which case, prior to reporting patient test results, the individual’s performance must be reevaluated to include the use of the new test methodology or instrumentation.

§ 493.1415 Condition: Laboratories performing moderate complexity testing; clinical consultant.

The laboratory must have a clinical consultant who meets the qualification requirements of § 493.1417 of this part and provides clinical consultation in accordance with § 493.1419 of this part.

§ 493.1417 Standard; Clinical consultant qualifications.

The clinical consultant must be qualified to consult with and render opinions to the laboratory’s clients concerning the diagnosis, treatment and management of patient care. The clinical consultant must—
(a) Be qualified as a laboratory director under § 493.1405(b) (1), (2), or (3)(i); or
(b) Be a doctor of medicine, doctor of osteopathy or doctor of podiatric medicine and possess a license to practice medicine, osteopathy or podiatry in the State in which the laboratory is located.

§ 493.1419 Standard; Clinical consultant responsibilities.

The clinical consultant provides consultation regarding the appropriateness of the testing ordered and interpretation of test results. The clinical consultant must—

(a) Be available to provide clinical consultation to the laboratory's clients;

(b) Be available to assist the laboratory's clients in ensuring that appropriate tests are ordered to meet the clinical expectations;

(c) Ensure that reports of test results include pertinent information required for specific patient interpretation; and

(d) Ensure that consultation is available and communicated to the laboratory's clients on matters related to the quality of the test results reported and their interpretation concerning specific patient conditions.

§ 493.1421 Condition: Laboratories performing moderate complexity testing; testing personnel.

The laboratory must have a sufficient number of individuals who meet the qualification requirements of § 493.1423, to perform the functions specified in § 493.1425 for the volume and complexity of tests performed.

§ 493.1423 Standard; Testing personnel qualifications.

Each individual performing moderate complexity testing must—

(a) Possess a current license issued by the State in which the laboratory is located, if such licensing is required; and

(b) Meet one of the following requirements:

(1) Be a doctor of medicine or doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located or have earned a doctoral, master's, or bachelor's degree in a chemical, physical, biological or clinical laboratory science, or medical technology from an accredited institution; or

(2) Have earned an associate degree in a chemical, physical or biological science or medical laboratory technology from an accredited institution; or

(3) Be a high school graduate or equivalent and have successfully completed an official military medical laboratory procedures course of at least 50 weeks duration and have held the military enlisted occupational specialty of Medical Laboratory Specialist (Laboratory Technician); or

(4)(i) Have earned a high school diploma or equivalent; and

(ii) Have documentation of training appropriate for the testing performed prior to analyzing patient specimens. Such training must ensure that the individual has—

(A) The skills required for proper specimen collection, including patient preparation, if applicable, labeling, handling, preservation or fixation, processing or preparation, transportation and storage of specimens;

(B) The skills required for implementing all standard laboratory procedures;

(C) The skills required for performing each test method and for proper instrument use;

(D) The skills required for performing preventive maintenance, troubleshooting and calibration procedures related to each test performed;

(E) A working knowledge of reagent stability and storage;

(F) The skills required to implement the quality control policies and procedures of the laboratory;

(G) An awareness of the factors that influence test results; and

(H) The skills required to assess and verify the validity of patient test results through the evaluation of quality control sample values prior to reporting patient test results.


§ 493.1425 Standard; Testing personnel responsibilities.

The testing personnel are responsible for specimen processing, test performance, and for reporting test results.

(a) Each individual performs only those moderate complexity tests that are authorized by the laboratory director and require a degree of skill commensurate with the individual's education, training or experience, and technical abilities.
§ 493.1443 Standard; Laboratory director qualifications.

The laboratory director must be qualified to manage and direct the laboratory personnel and performance of high complexity tests and must be eligible to be an operator of a laboratory within the requirements of subpart R.

(a) The laboratory director must possess a current license as a laboratory director issued by the State in which the laboratory is located, if such licensing is required; and

(b) The laboratory director must—

(1)(i) Be a doctor of medicine or doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(ii) Be certified in anatomic or clinical pathology, or both, by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(2) Be a doctor of medicine, a doctor of osteopathy or doctor of podiatric medicine licensed to practice medicine, osteopathy or podiatry in the State in which the laboratory is located; and

(i) Have at least one year of laboratory training during medical residency (for example, physicians certified either in hematology or hematology and medical oncology by the American Board of Internal Medicine); or

(ii) Have at least 2 years of experience directing or supervising high complexity testing; or

(3) Hold an earned doctoral degree in a chemical, physical, biological or clinical laboratory science from an accredited institution and—

(i) Be certified by the American Board of Medical Microbiology, the American Board of Clinical Chemistry, the American Board of Bioanalysis, the American Board of Medical Laboratory Immunology or other board deemed comparable by HHS; or

(ii) Until December 31, 2000, must have at least—

(A) Two years of laboratory training or experience, or both;

(B) Two years of experience directing or supervising high complexity testing; and

(C) On December 31, 2000, individuals must meet the qualifications specified in paragraph (b)(3)(i) of this section;

(4) Be serving as a laboratory director and must have previously qualified or could have qualified as a laboratory director under regulations at 42 CFR 493.1415, published March 24, 1990 at 55 FR 9539, on or before February 28, 1992; or

(5) On or before February 28, 1992, be qualified under State law to direct a laboratory in the State in which the laboratory is located; or

(6) For the subspecialty of oral pathology, be certified by the American
§ 493.1445 Standard; Laboratory director responsibilities.

The laboratory director is responsible for the overall operation and administration of the laboratory, including the employment of personnel who are competent to perform test procedures, record and report test results promptly, accurately and proficiently, and for assuring compliance with the applicable regulations.

(a) The laboratory director, if qualified, may perform the duties of the technical supervisor, clinical consultant, general supervisor, and testing personnel, or delegate these responsibilities to personnel meeting the qualifications under §§ 493.1447, 493.1453, 493.1459, and 493.1487, respectively.

(b) If the laboratory director reassesses performance of his or her responsibilities, he or she remains responsible for ensuring that all duties are properly performed.

(c) The laboratory director must be accessible to the laboratory to provide onsite, telephone or electronic consultation as needed.

(d) Each individual may direct no more than five laboratories.

(e) The laboratory director must—

(1) Ensure that testing systems developed and used for each of the tests performed in the laboratory provide quality laboratory services for all aspects of test performance, which includes the preanalytic, analytic, and postanalytic phases of testing;

(2) Ensure that the physical plant and environmental conditions of the laboratory are appropriate for the testing performed and provide a safe environment in which employees are protected from physical, chemical, and biological hazards;

(3) Ensure that—

(i) The test methodologies selected have the capability of providing the quality of results required for patient care;

(ii) Verification procedures used are adequate to determine the accuracy, precision, and other pertinent performance characteristics of the method; and

(iii) Laboratory personnel are performing the test methods as required for accurate and reliable results;

(4) Ensure that the laboratory is enrolled in an HHS-approved proficiency testing program for the testing performed and that—

(i) The proficiency testing samples are tested as required under subpart H of this part;

(ii) The results are returned within the timeframes established by the proficiency testing program;

(iii) All proficiency testing reports received are reviewed by the appropriate staff to evaluate the laboratory’s performance and to identify any problems that require corrective action; and

(iv) An approved corrective action plan is followed when any proficiency testing result is found to be unacceptable or unsatisfactory;

(5) Ensure that the quality control and quality assurance programs are established and maintained to assure the quality of laboratory services provided and to identify failures in quality as they occur;

(6) Ensure the establishment and maintenance of acceptable levels of analytical performance for each test system;

(7) Ensure that all necessary remedial actions are taken and documented whenever significant deviations from the laboratory’s established performance characteristics are identified, and that patient test results are reported only when the system is functioning properly;

(8) Ensure that reports of test results include pertinent information required for interpretation;

(9) Ensure that consultation is available to the laboratory’s clients on matters relating to the quality of the test results reported and their interpretation concerning specific patient conditions;

(10) Ensure that a general supervisor provides on-site supervision of high complexity test performance by testing
personnel qualified under § 493.1489(b)(4);

(11) Employ a sufficient number of laboratory personnel with the appropriate education and either experience or training to provide appropriate consultation, properly supervise and accurately perform tests and report test results in accordance with the personnel responsibilities described in this subpart;

(12) Ensure that prior to testing patients' specimens, all personnel have the appropriate education and experience, receive the appropriate training for the type and complexity of the services offered, and have demonstrated that they can perform all testing operations reliably to provide and report accurate results;

(13) Ensure that policies and procedures are established for monitoring individuals who conduct preanalytical, analytical, and postanalytical phases of testing to assure that they are competent and maintain their competency to process specimens, perform test procedures and report test results promptly and proficiently, and whenever necessary, identify needs for remedial training or continuing education to improve skills;

(14) Ensure that an approved procedure manual is available to all personnel responsible for any aspect of the testing process; and

(15) Specify, in writing, the responsibilities and duties of each consultant and each supervisor, as well as each person engaged in the performance of the preanalytic, analytic, and postanalytic phases of testing, that identifies which examinations and procedures each individual is authorized to perform, whether supervision is required for specimen processing, test performance or result reporting and whether supervisory or director review is required prior to reporting patient test results.

§ 493.1447 Condition: Laboratories performing high complexity testing; technical supervisor.

The laboratory must have a technical supervisor who meets the qualification requirements of § 493.1449 of this subpart and provides technical supervision in accordance with § 493.1451 of this subpart.


The laboratory must employ one or more individuals who are qualified by education and either training or experience to provide technical supervision for each of the specialties and subspecialties of service in which the laboratory performs high complexity tests or procedures. The director of a laboratory performing high complexity testing may function as the technical supervisor provided he or she meets the qualifications specified in this section.

(a) The technical supervisor must possess a current license issued by the State in which the laboratory is located, if such licensing is required; and

(b) The laboratory may perform anatomic and clinical laboratory procedures and tests in all specialties and subspecialties of services except histocompatibility and clinical cytogentic services provided the individual functioning as the technical supervisor—

(1) Is a doctor of medicine or doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(2) Is certified in both anatomic and clinical pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possesses qualifications that are equivalent to those required for such certification.

(c) If the requirements of paragraph (b) of this section are not met and the laboratory performs tests in the subspecialty of bacteriology, the individual functioning as the technical supervisor must—

(1)(i) Be a doctor of medicine or doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(ii) Be certified in clinical pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(2)(i) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine,
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osteopathy, or podiatry in the State in which the laboratory is located; and

(ii) Have at least one year of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of bacteriology; or

(3)(i) Have an earned doctoral degree in a chemical, physical, biological or clinical laboratory science from an accredited institution; and

(ii) Have at least one year of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of bacteriology; or

(4)(i) Have earned a master's degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution; and

(ii) Have at least two years of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of bacteriology.

(d) If the requirements of paragraph (b) of this section are not met and the laboratory performs tests in the subspecialty of mycobacteriology, the individual functioning as the technical supervisor must—

(1)(i) Be a doctor of medicine or doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(ii) Be certified in clinical pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(2)(i) Be a doctor of medicine, doctor of osteopathy, or doctor or podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and

(ii) Have at least one year of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of mycobacteriology; or

(3)(i) Have an earned doctoral degree in a chemical, physical, biological or clinical laboratory science from an accredited institution; and

(ii) Have at least one year of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of mycobacteriology; or

(4)(i) Have earned a master's degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution; and

(ii) Have at least two years of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of mycobacteriology; or

(5)(i) Have earned a bachelor's degree in a chemical, physical, or biological science or medical technology from an accredited institution; and

(ii) Have at least four years of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of mycobacteriology.

(e) If the requirements of paragraph (b) of this section are not met and the laboratory performs tests in the subspecialty of mycology, the individual functioning as the technical supervisor must—

(1)(i) Be a doctor of medicine or doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(ii) Be certified in clinical pathology by the American Board of Pathology or
the American osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(2)(i) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and

(ii) Have at least 1 year of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of mycology;
or

(3)(i) Have an earned doctoral degree in a chemical, physical, biological or clinical laboratory science from an accredited institution; and

(ii) Have at least 1 year of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of mycology;
or

(4)(i) Have earned a master’s degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution; and

(ii) Have at least 2 years of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of mycology;
or

(5)(i) Have earned a bachelor’s degree in a chemical, physical or biological science or medical technology from an accredited institution; and

(ii) Have at least 4 years of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of mycology.

(f) If the requirements of paragraph (b) of this section are not met and the laboratory performs tests in the subspecialty of parasitology, the individual functioning as the technical supervisor must—

(1)(i) Be a doctor of medicine or a doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(ii) Be certified in clinical pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(2)(i) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and

(ii) Have at least 1 year of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of parasitology;
or

(3)(i) Have an earned doctoral degree in a chemical, physical, biological or clinical laboratory science from an accredited institution; and

(ii) Have at least 1 year of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of parasitology;
or

(4)(i) Have earned a master’s degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution; and

(ii) Have at least 2 years of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of parasitology;
or

(5)(i) Have earned a bachelor’s degree in a chemical, physical or biological science or medical technology from an accredited institution; and

(ii) Have at least 4 years of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of parasitology.
functioning as the technical supervisor must—

(1)(i) Be a doctor of medicine or doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(ii) Be certified in clinical pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(2)(i) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and

(ii) Have at least 1 year of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of virology; or

(3)(i) Have an earned doctoral degree in a chemical, physical, biological or clinical laboratory science from an accredited institution; and

(ii) Have at least 1 year of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of virology; or

(4)(i) Have earned a master's degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution; and

(ii) Have at least 2 years of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of virology; or

(5)(i) Have earned a bachelor's degree in a chemical, physical or biological science or medical technology from an accredited institution; and

(ii) Have at least 4 years of laboratory training or experience, or both, in high complexity testing within the specialty of microbiology with a minimum of 6 months experience in high complexity testing within the subspecialty of virology.

(h) If the requirements of paragraph (b) of this section are not met and the laboratory performs tests in the specialty of diagnostic immunology, the individual functioning as the technical supervisor must—

(1)(i) Be a doctor of medicine or a doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(ii) Be certified in clinical pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(2)(i) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and

(ii) Have at least 1 year of laboratory training or experience, or both, in high complexity testing for the specialty of diagnostic immunology; or

(3)(i) Have an earned doctoral degree in a chemical, physical, biological or clinical laboratory science from an accredited institution; and

(ii) Have at least 1 year of laboratory training or experience, or both, in high complexity testing for the specialty of diagnostic immunology; or

(4)(i) Have earned a master's degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution; and

(ii) Have at least 2 years of laboratory training or experience, or both, in high complexity testing for the specialty of diagnostic immunology; or

(5)(i) Have earned a bachelor's degree in a chemical, physical or biological science or medical technology from an accredited institution; and

(ii) Have at least 4 years of laboratory training or experience, or both, in high complexity testing for the specialty of diagnostic immunology.

(i) If the requirements of paragraph (b) of this section are not met and the laboratory performs tests in the specialty of chemistry, the individual functioning as the technical supervisor must—

(1)(i) Be a doctor of medicine or doctor of osteopathy licensed to practice
(i) Be certified in clinical pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(ii) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and

(iii) Have at least 1 year of laboratory training or experience, or both, in high complexity testing for the specialty of chemistry; or

(i) Have an earned doctoral degree in a chemical, physical, biological or clinical laboratory science from an accredited institution; and

(ii) Have at least 1 year of laboratory training or experience, or both, in high complexity testing for the specialty of chemistry; or

(i) Have earned a master’s degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution; and

(ii) Have at least 2 years of laboratory training or experience, or both, in high complexity testing for the specialty of chemistry; or

(i) Have earned a bachelor’s degree in a chemical, physical or biological science or medical technology from an accredited institution; and

(ii) Have at least 4 years of laboratory training or experience, or both, in high complexity testing for the specialty of chemistry.

(j) If the requirements of paragraph (b) of this section are not met and the laboratory performs tests in the specialty of hematology, the individual functioning as the technical supervisor must—

(i) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and

(ii) Be certified in clinical pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(ii) Be a doctor of medicine, or doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and

(i) Have at least one year of laboratory training or experience, or both, in high complexity testing for the specialty of hematology (for example, physicians certified either in hematology or hematology and medical oncology by the American Board of Internal Medicine); or

(i) Have an earned doctoral degree in a chemical, physical, biological or clinical laboratory science from an accredited institution; and

(ii) Have at least 1 year of laboratory training or experience, or both, in high complexity testing within the specialty of hematology; or

(i) Have earned a master’s degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution; and

(ii) Have at least 2 years of laboratory training or experience, or both, in high complexity testing for the specialty of hematology; or

(i) Have earned a bachelor’s degree in a chemical, physical or biological science or medical technology from an accredited institution; and

(ii) Have at least 4 years of laboratory training or experience, or both, in high complexity testing for the specialty of hematology.

(k) If the requirements of paragraph (b) of this section are not met and the laboratory performs tests in the subspecialty of cytology, the individual functioning as the technical supervisor must—

(i) Be a doctor of medicine or a doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(ii) Meet one of the following requirements—

(A) Be certified in anatomic pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(B) Be certified by the American Society of Cytopathology to practice cytopathology or possess qualifications
that are equivalent to those required for such certification;
(2) An individual qualified under §493.1449(b) or paragraph (k)(1) of this section may delegate some of the cytology technical supervisor responsibilities to an individual who is in the final year of full-time training leading to certification specified in paragraphs (b) or (k)(1)(iii)(A) of this section provided the technical supervisor qualified under §493.1449(b) or paragraph (k)(1) of this section remains ultimately responsible for ensuring that all of the responsibilities of the cytology technical supervisor are met.

(l) If the requirements of paragraph (b) of this section are not met and the laboratory performs tests in the subspecialty of histopathology, the individual functioning as the technical supervisor must—

(1) Meet one of the following requirements:

(A) Be a doctor of medicine or a doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(B) Be certified in anatomic pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification;

(ii) An individual qualified under §493.1449(b) or paragraph (l)(1) of this section may delegate to an individual who is a resident in a training program leading to certification specified in paragraphs (b) or (l)(2)(i)(B) of this section, the responsibility for examination and interpretation of histopathology specimens.

(3) For tests in ophthalmic pathology, meet one of the following requirements:

(A) Be a doctor of medicine or doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located and—

(B) Must meet one of the following requirements:

(i) An individual qualified under §493.1449(b) or paragraph (l)(3)(i) of this section may delegate to an individual who is a resident in a training program leading to certification specified in paragraphs (b) or (l)(3)(i)(B) of this section, the responsibility for examination and interpretation of ophthalmic pathology specimens;

(m) If the requirements of paragraph (b) of this section are not met and the laboratory performs tests in the subspecialty of oral pathology, the individual functioning as the technical supervisor must meet one of the following requirements:

(2) Be certified in dermatopathology by the American Board of Dermatology and the American Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(3) Be certified in dermatology by the American Board of Dermatology or possess qualifications that are equivalent to those required for such certification; or

(ii) An individual qualified under §493.1449(b) or paragraph (l)(2)(i) of this section may delegate to an individual who is a resident in a training program leading to certification specified in paragraphs (b) or (l)(2)(i)(B) of this section, the responsibility for examination and interpretation of dermatopathology specimens.

(3) For tests in ophthalmic pathology, meet one of the following requirements:

(A) Be a doctor of medicine or doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located and—

(B) Must meet one of the following requirements:

(i) An individual qualified under §493.1449(b) or paragraph (l)(1)(i) of this section may delegate to an individual who is a resident in a training program leading to certification specified in paragraphs (b) or (l)(1)(i)(B) of this section, the responsibility for examination and interpretation of histopathology specimens.

(2) For tests in dermatopathology, meet one of the following requirements:

(A) Be a doctor of medicine or doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located and—

(B) Meet one of the following requirements:

(i) Be certified in anatomic pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(ii) Be certified in dermatopathology by the American Board of Dermatology and the American Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(iii) Be certified by the American Board of Ophthalmology or possess qualifications that are equivalent to those required for such certification and have successfully completed at least 1 year of formal post-residency fellowship training in ophthalmic pathology; or

(ii) An individual qualified under §493.1449(b) or paragraph (l)(3)(i) of this section may delegate to an individual who is a resident in a training program leading to certification specified in paragraphs (b) or (l)(3)(i)(B) of this section, the responsibility for examination and interpretation of ophthalmic pathology specimens; or
(1)(i) Be a doctor of medicine or a doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located and—
   (ii) Be certified in anatomic pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or
(2) Be certified in oral pathology by the American Board of Oral Pathology or possess qualifications for such certification; or
(3) An individual qualified under §493.1449(b) or paragraph (m) (1) or (2) of this section may delegate to an individual who is a resident in a training program leading to certification specified in paragraphs (b) or (m) (1) or (2) of this section, the responsibility for examination and interpretation of oral pathology specimens.
(n) If the requirements of paragraph (b) of this section are not met and the laboratory performs tests in the specialty of radiobioassay, the individual functioning as the technical supervisor must—
   (1)(i) Be a doctor of medicine or a doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and
   (ii) Be certified in clinical pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or
(2)(i) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and
   (ii) Have at least 1 year of laboratory training or experience, or both, in high complexity testing for the specialty of radiobioassay; or
(4)(i) Have earned a master's degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution; and
   (ii) Have at least 2 years of laboratory training or experience, or both, in high complexity testing for the specialty of radiobioassay; or
(5)(i) Have earned a bachelor's degree in a chemical, physical or biological science or medical technology from an accredited institution; and
   (ii) Have at least 4 years of laboratory training or experience, or both, in high complexity testing for the specialty of radiobioassay.
(o) If the laboratory performs tests in the specialty of histocompatibility, the individual functioning as the technical supervisor must either—
   (1)(i) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and
   (ii) Have training or experience that meets one of the following requirements:
      (A) Have 4 years of laboratory training or experience, or both, within the specialty of histocompatibility; or
      (B)(1) Have 2 years of laboratory training or experience, or both, in the specialty of general immunology; and
       (2) Have 2 years of laboratory training or experience, or both, in the specialty of histocompatibility; or
   (2)(i) Have an earned doctoral degree in a biological or clinical laboratory science from an accredited institution; and
   (ii) Have training or experience that meets one of the following requirements:
      (A) Have 4 years of laboratory training or experience, or both, within the specialty of histocompatibility; or
      (B)(1) Have 2 years of laboratory training or experience, or both, in the specialty of general immunology; and
       (2) Have 2 years of laboratory training or experience, or both, in the specialty of histocompatibility.
(p) If the laboratory performs tests in the specialty of clinical cytogenetics, the individual functioning as the technical supervisor must—
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(1)(i) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and

(ii) Have 4 years of training or experience, or both, in genetics, 2 of which have been in clinical cytogenetics; or

(2)(i) Hold an earned doctoral degree in a biological science, including biochemistry, or clinical laboratory science from an accredited institution; and

(ii) Have 4 years of training or experience, or both, in genetics, 2 of which have been in clinical cytogenetics.

(q) If the requirements of paragraph (b) of this section are not met and the laboratory performs tests in the specialty of immunohematology, the individual functioning as the technical supervisor must—

(1)(i) Be a doctor of medicine or a doctor of osteopathy licensed to practice medicine or osteopathy in the State in which the laboratory is located; and

(ii) Be certified in clinical pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications that are equivalent to those required for such certification; or

(2)(i) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located; and

(ii) Have at least one year of laboratory training or experience, or both, in high complexity testing for the specialty of immunohematology.

NOTE: The technical supervisor requirements for “laboratory training or experience or both” in each specialty or subspecialty may be acquired concurrently in more than one of the specialties or subspecialties of service. For example, an individual, who has a doctoral degree in chemistry and additionally has documentation of 1 year of laboratory experience working concurrently in high complexity testing in the specialties of microbiology and chemistry and 6 months of that work experience included high complexity testing in bacteriology, mycology, and mycobacteriology, would qualify as the technical supervisor for the specialty of chemistry and the subspecialties of bacteriology, mycology, and mycobacteriology.


The technical supervisor is responsible for the technical and scientific oversight of the laboratory. The technical supervisor is not required to be on site at all times testing is performed; however, he or she must be available to the laboratory on an as needed basis to provide supervision as specified in (a) of this section.

(a) The technical supervisor must be accessible to the laboratory to provide on-site, telephone, or electronic consultation; and

(b) The technical supervisor is responsible for—

(1) Selection of the test methodology that is appropriate for the clinical use of the test results;

(2) Verification of the test procedures performed and establishment of the laboratory's test performance characteristics, including the precision and accuracy of each test and test system;

(3) Enrollment and participation in an HHS approved proficiency testing program commensurate with the services offered;

(4) Establishing a quality control program appropriate for the testing performed and establishing the parameters for acceptable levels of analytic performance and ensuring that these levels are maintained throughout the entire testing process from the initial receipt of the specimen, through sample analysis and reporting of test results;

(5) Resolving technical problems and ensuring that remedial actions are taken whenever test systems deviate from the laboratory's established performance specifications;

(6) Ensuring that patient test results are not reported until all corrective actions have been taken and the test system is functioning properly;

(7) Identifying training needs and assuring that each individual performing tests receives regular in-service training and education appropriate for the type and complexity of the laboratory services performed;
(8) Evaluating the competency of all testing personnel and assuring that the staff maintain their competency to perform test procedures and report test results promptly, accurately and proficiently. The procedures for evaluation of the competency of the staff must include, but are not limited to—
(i) Direct observations of routine patient test performance, including patient preparation, if applicable, specimen handling, processing and testing;
(ii) Monitoring the recording and reporting of test results;
(iii) Review of intermediate test results or worksheets, quality control records, proficiency testing results, and preventive maintenance records;
(iv) Direct observation of performance of instrument maintenance and function checks;
(v) Assessment of test performance through testing previously analyzed specimens, internal blind testing samples or external proficiency testing samples; and
(vi) Assessment of problem solving skills; and
(9) Evaluating and documenting the performance of individuals responsible for high complexity testing at least semiannually during the first year the individual tests patient specimens. Thereafter, evaluations must be performed at least annually unless test methodology or instrumentation changes, in which case, prior to reporting patient test results, the individual’s performance must be reevaluated to include the use of the new test methodology or instrumentation.

(c) In cytology, the technical supervisor or the individual qualified under §493.1449(k)(2)—
(1) May perform the duties of the cytology general supervisor and the cytotechnologist, as specified in §§493.1471 and 493.1485, respectively;
(2) Must establish the workload limit for each individual examining slides;
(3) Must reassess the workload limit for each individual examining slides at least every 6 months and adjust as necessary;
(4) Must perform the functions specified in §493.1257(c);
(5) Must ensure that each individual examining gynecologic preparations participates in an HHS approved cytology proficiency testing program, as specified in §493.945 and achieves a passing score, as specified in §493.855; and
(6) If responsible for screening cytology slide preparations, must document the number of cytology slides screened in 24 hours and the number of hours devoted during each 24-hour period to screening cytology slides.

§493.1453 Condition: Laboratories performing high complexity testing; clinical consultant.

The laboratory must have a clinical consultant who meets the requirements of §493.1455 of this subpart and provides clinical consultation in accordance with §493.1457 of this subpart.

§493.1455 Standard; Clinical consultant qualifications.

The clinical consultant must be qualified to consult with and render opinions to the laboratory’s clients concerning the diagnosis, treatment and management of patient care. The clinical consultant must—
(a) Be qualified as a laboratory director under §493.1443(b)(1), (2), or (3)(i) or, for the subspecialty of oral pathology, §493.1443(b)(6); or
(b) Be a doctor of medicine, doctor of osteopathy, doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located.

§493.1457 Standard; Clinical consultant responsibilities.

The clinical consultant provides consultation regarding the appropriateness of the testing ordered and interpretation of test results. The clinical consultant must—
(a) Be available to provide consultation to the laboratory’s clients;
(b) Be available to assist the laboratory’s clients in ensuring that appropriate tests are ordered to meet the clinical expectations;
(c) Ensure that reports of test results include pertinent information required for specific patient interpretation; and
§ 493.1459  Condition: Laboratories performing high complexity testing; general supervisor.

The laboratory must have one or more general supervisors who, under the direction of the laboratory director and supervision of the technical supervisor, provides day-to-day supervision of testing personnel and reporting of test results. In the absence of the director and technical supervisor, the general supervisor must be responsible for the proper performance of all laboratory procedures and reporting of test results.

(a) The general supervisor must possess a current license issued by the State in which the laboratory is located, if such licensing is required; and
(b) The general supervisor must be qualified as—
   (1) Laboratory director under § 493.1443; or
   (2) Technical supervisor under § 493.1449.

(c) If the requirements of paragraph (b)(1) or paragraph (b)(2) of this section are not met, the individual functioning as the general supervisor must—
   (1) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located or have earned a doctoral or master’s degree in a chemical, physical, biological or clinical laboratory science, or medical technology from an accredited institution; and
   (ii) Have at least 2 years of laboratory training or experience, or both, in high complexity testing; or
   (3) Be a high school graduate or equivalent; and
   (ii) Have had at least 10 years of laboratory training or experience, or both, in high complexity testing, including at least 6 years of supervisory experience between September 1, 1982 and September 1, 1992.

(d) For blood gas analysis, the individual providing general supervision must—
   (1) Be qualified under §§ 493.1461(b)(1) or (2), or § 493.1461(c); or
   (2) Meet one of the following requirements:
      (A) Have graduated from a medical laboratory or clinical laboratory training program approved or accredited by the Accrediting Bureau of Health Education Schools (ABHES), the Commission on Allied Health Education Accreditation (CAHEA), or other organization approved by HHS.
      (B) Be a high school graduate or equivalent and have successfully completed an official U.S. military medical laboratory procedures course of at least 50 weeks duration and have held the military enlisted occupational specialty of Medical Laboratory Specialist (Laboratory Technician).
      (i) Meet one of the following requirements:
         (A) Have graduated from a medical laboratory or clinical laboratory training program approved or accredited by the Accrediting Bureau of Health Education Schools (ABHES), the Commission on Allied Health Education Accreditation (CAHEA), or other organization approved by HHS.
         (B) Be a high school graduate or equivalent; and
         (ii) Have had at least 10 years of laboratory training or experience, or both, in high complexity testing, including at least 6 years of supervisory experience between September 1, 1982 and September 1, 1992.

   (3) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located or have earned a doctoral or master’s degree in a chemical, physical, biological or clinical laboratory science, or medical technology from an accredited institution; and
   (ii) Have at least 2 years of laboratory training or experience, or both, in high complexity testing; or
   (3)(i) Except as specified in paragraph (3)(ii) of this section, have previously qualified as a general supervisor under § 493.1462 on or before February 28, 1992.
   (ii) Exception. An individual who achieved a satisfactory grade in a proficiency examination for technologist given by HHS between March 1, 1966 and December 31, 1987, qualifies as a general supervisor if he or she meets the requirements of § 493.1462 on or before January 1, 1994.

(4) On or before September 1, 1992, have served as a general supervisor of high complexity testing and as of April 24, 1995—
   (i) Meet one of the following requirements:
      (A) Have graduated from a medical laboratory or clinical laboratory training program approved or accredited by the Accrediting Bureau of Health Education Schools (ABHES), the Commission on Allied Health Education Accreditation (CAHEA), or other organization approved by HHS.
      (B) Be a high school graduate or equivalent and have successfully completed an official U.S. military medical laboratory procedures course of at least 50 weeks duration and have held the military enlisted occupational specialty of Medical Laboratory Specialist (Laboratory Technician).
      (i) Meet one of the following requirements:
         (A) Have graduated from a medical laboratory or clinical laboratory training program approved or accredited by the Accrediting Bureau of Health Education Schools (ABHES), the Commission on Allied Health Education Accreditation (CAHEA), or other organization approved by HHS.
         (B) Be a high school graduate or equivalent; and
         (ii) Have had at least 10 years of laboratory training or experience, or both, in high complexity testing, including at least 6 years of supervisory experience between September 1, 1982 and September 1, 1992.

   (d) For blood gas analysis, the individual providing general supervision must—
   (1) Be qualified under §§ 493.1461(b)(1) or (2), or § 493.1461(c); or
(2)(i) Have earned a bachelor’s degree in respiratory therapy or cardiovascular technology from an accredited institution; and

(ii) Have at least one year of laboratory training or experience, or both, in blood gas analysis; or

(3)(i) Have earned an associate degree related to pulmonary function from an accredited institution; and

(ii) Have at least two years of training or experience, or both in blood gas analysis.

(e) The general supervisor requirement is met in histopathology, oral pathology, dermatopathology, and ophthalmic pathology because all tests and examinations, must be performed:

(1) In histopathology, by an individual who is qualified as a technical supervisor under §§ 493.1449(b) or 493.1449(1);

(2) In dermatopathology, by an individual who is qualified as a technical supervisor under §§ 493.1449(b) or 493.1449(1) or (2);

(3) In ophthalmic pathology, by an individual who is qualified as a technical supervisor under §§ 493.1449(b) or 493.1449(1)(3); and

(4) In oral pathology, by an individual who is qualified as a technical supervisor under §§ 493.1449(b) or 493.1449(m).


§ 493.1463 Standard: General supervisor responsibilities.

The general supervisor is responsible for day-to-day supervision or oversight of the laboratory operation and personnel performing testing and reporting test results.

(a) The general supervisor—(1) Must be accessible to testing personnel at all times testing is performed to provide on-site, telephone or electronic consultation to resolve technical problems in accordance with policies and procedures established either by the laboratory director or technical supervisor;

(2) Is responsible for providing day-to-day supervision of high complexity test performance by a testing personnel qualified under § 493.1489;

(3) Except as specified in paragraph (c) of this section, must be onsite to provide direct supervision when high
§ 493.1467 Condition: Laboratories performing high complexity testing; cytology general supervisor.

For the subspecialty of cytology, the laboratory must have a general supervisor who meets the qualification requirements of §493.1469 of this subpart, and provides supervision in accordance with §493.1471 of this subpart.

§ 493.1469 Standard: Cytology general supervisor qualifications.

The cytology general supervisor must be qualified to supervise cytology services. The general supervisor in cytology must possess a current license issued by the State in which the laboratory is located, if such licensing is required, and must—

(a) Be qualified as a technical supervisor under §493.1449 (b) or (k); or

(b)(1) Be qualified as a cytotechnologist under §493.1483; and

(2) Have at least 3 years of full-time (2,080 hours per year) experience as a cytotechnologist within the preceding 10 years.

§ 493.1471 Standard: Cytology general supervisor responsibilities.

The technical supervisor of cytology may perform the duties of the cytology general supervisor or delegate the responsibilities to an individual qualified under §493.1469.

(a) The cytology general supervisor is responsible for the day-to-day supervision or oversight of the laboratory operation and personnel performing testing and reporting test results.

(b) The cytology general supervisor must—

(1) Be accessible to provide on-site, telephone, or electronic consultation to resolve technical problems in accordance with policies and procedures established by the technical supervisor of cytology;

(2) Document the slide interpretation results of each gynecologic and nongynecologic cytology case he or she examined or reviewed (as specified under §493.1257(d));

(3) For each 24-hour period, document the total number of slides he or she examined or reviewed in the laboratory as well as the total number of slides examined or reviewed in any other laboratory or for any other employer; and

(4) Document the number of hours spent examining slides in each 24-hour period.

§ 493.1481 Condition: Laboratories performing high complexity testing; cytotechnologist.

For the subspecialty of cytology, the laboratory must have a sufficient number of cytotechnologists who meet the qualifications specified in §493.1483 to perform the functions specified in §493.1485.

§ 493.1483 Standard: Cytotechnologist qualifications.

Each person examining cytology slide preparations must meet the qualifications of §493.1449 (b) or (k), or—

(a) Possess a current license as a cytotechnologist issued by the State in
§ 493.1489 Testing personnel qualifications.

Each individual performing high complexity testing must—
(a) Possess a current license issued by the State in which the laboratory is located, if such licensing is required; and
(b) Meet one of the following requirements:
   (1) Be a doctor of medicine, doctor of osteopathy, or doctor of podiatric medicine licensed to practice medicine, osteopathy, or podiatry in the State in which the laboratory is located or have earned a doctoral, master's or bachelor's degree in a chemical, physical, biological or clinical laboratory science, or medical technology from an accredited institution;
   (2)(i) On or before September 1, 1994, have full-time experience of at least 2 years or equivalent examining cytology slide preparations within the preceding 5 years in the United States under the supervision of a physician qualified under §493.1449(b) or (k)(1); and
   (ii) On or before September 1, 1995, have met the requirements in either paragraph (b)(1) or (2) of this section.

The cytotechnologist is responsible for documenting—
(a) The slide interpretation results of each gynecologic and nongynecologic cytology case he or she examined or reviewed (as specified in §493.1257(d));
(b) For each 24-hour period, the total number of slides examined or reviewed in the laboratory as well as the total number of slides examined or reviewed in any other laboratory or for any other employer; and
(c) The number of hours spent examining slides in each 24-hour period.

§ 493.1487 Condition: Laboratories performing high complexity testing; testing personnel.

The laboratory has a sufficient number of individuals who meet the qualification requirements of §493.1489 of this subpart to perform the functions specified in §493.1495 of this subpart for the volume and complexity of testing performed.

§ 493.1485 Standard; Cytotechnologist responsibilities.

The cytotechnologist is responsible for documenting—
(a) The slide interpretation results of each gynecologic and nongynecologic cytology case he or she examined or reviewed (as specified in §493.1257(d));
(b) Meet one of the following requirements:
   (1) Have graduated from a school of cytotechnology accredited by the Committee on Allied Health Education and Accreditation or other organization approved by HHS; or
   (2) Be certified in cytotechnology by a certifying agency approved by HHS; or
   (3) Before September 1, 1992—
      (i) Have successfully completed 2 years in an accredited institution with at least 12 semester hours in science, 8 hours of which are in biology; and
      (A) Have had 12 months of training in a school of cytotechnology accredited by an accrediting agency approved by HHS; or
      (B) Have received 6 months of formal training in a school of cytotechnology accredited by an accrediting agency approved by HHS and 6 months of full-time experience in cytotechnology in a laboratory acceptable to the pathologist who directed the formal 6 months of training; or
      (ii) Have achieved a satisfactory grade to qualify as a cytotechnologist in a proficiency examination approved by HHS and designed to qualify persons as cytotechnologists; or
   (4) Before September 1, 1994, have full-time experience of at least 2 years or equivalent within the preceding 5 years examining slide preparations under the supervision of a physician qualified under §493.1449(b) or (k)(1), and before January 1, 1969, must have—
      (i) Graduated from high school;
      (ii) Completed 6 months of training in cytotechnology in a laboratory directed by a pathologist or other physician providing cytology services; and
      (iii) Completed 2 years of full-time supervised experience in cytotechnology; or
   (5)(i) On or before September 1, 1994, have full-time experience of at least 2 years or equivalent examining cytology slide preparations within the preceding 5 years in the United States under the supervision of a physician qualified under §493.1449(b) or (k)(1); and
   (ii) On or before September 1, 1995, have met the requirements in either paragraph (b)(1) or (2) of this section.
§ 493.1491 Technologist qualifications on or before February 28, 1992.

In order to qualify as high complexity testing personnel under § 493.1489(b)(3), the individual must have met or could have met the following qualifications for technologist as they were in effect on or before February 28, 1992. Each technologist must—

(ii) Have education and training equivalent to that specified in paragraph (b)(2)(i) of this section that includes—

(A) At least 60 semester hours, or equivalent, from an accredited institution that, at a minimum, include either—

(1) 24 semester hours of medical laboratory technology courses; or

(2) 24 semester hours of science courses that include—

(i) Six semester hours of chemistry;

(ii) Six semester hours of biology; and

(iii) Twelve semester hours of chemistry, biology, or medical laboratory technology in any combination; and

(B) Have laboratory training that includes either of the following:

(1) Completion of a clinical laboratory training program approved or accredited by the ABHES, the CAHEA, or other organization approved by HHS. (This training may be included in the 60 semester hours listed in paragraph (b)(2)(ii)(A) of this section.)

(2) At least 3 months documented laboratory training in each specialty in which the individual performs high complexity testing.

(3) Have previously qualified or could have qualified as a technologist under § 493.1491 on or before February 28, 1992;

(4) On or before April 24, 1995 be a high school graduate or equivalent and have either—

(i) Graduated from a medical laboratory or clinical laboratory training program approved or accredited by ABHES, CAHEA, or other organization approved by HHS; or

(ii) Successfully completed an official U.S. military medical laboratory procedures training course of at least 50 weeks duration and have held the military enlisted occupational specialty of Medical Laboratory Specialist (Laboratory Technician);

(5)(i) Until September 1, 1997—

(A) Have earned a high school diploma or equivalent; and

(B) Have documentation of training appropriate for the testing performed before analyzing patient specimens. Such training must ensure that the individual has—

(1) The skills required for proper specimen collection, including patient preparation, if applicable, labeling, handling, preservation or fixation, processing or preparation, transportation and storage of specimens;

(2) The skills required for implementing all standard laboratory procedures;

(3) The skills required for performing each test method and for proper instrument use;

(4) The skills required for performing preventive maintenance, troubleshooting, and calibration procedures related to each test performed;

(5) A working knowledge of reagent stability and storage;

(6) The skills required to implement the quality control policies and procedures of the laboratory;

(7) An awareness of the factors that influence test results; and

(8) The skills required to assess and verify the validity of patient test results through the evaluation of quality control values before reporting patient test results; and

(ii) As of September 1, 1997, be qualified under § 493.1489(b)(1), (b)(2), or (b)(4), except for those individuals qualified under paragraph (b)(5)(i) of this section who were performing high complexity testing on or before April 24, 1995;

(6) For blood gas analysis—

(i) Be qualified under § 493.1489(b)(1), (b)(2), (b)(3), (b)(4), or (b)(5);

(ii) Have earned a bachelor’s degree in respiratory therapy or cardiovascular technology from an accredited institution; or

(iii) Have earned an associate degree related to pulmonary function from an accredited institution; or

(7) For histopathology, meet the qualifications of § 493.1449 (b) or (l) to perform tissue examinations.

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(a) Possess a current license as a laboratory technologist issued by the State, if such licensing exists; and

(b)(1) Have earned a bachelor’s degree in medical technology from an accredited university; or

(2) Have successfully completed 3 years of academic study (a minimum of 90 semester hours or equivalent) in an accredited college or university, which met the specific requirements for entrance into a school of medical technology accredited by an accrediting agency approved by the Secretary, and has successfully completed a course of training of at least 12 months in such a school;

(3) Have earned a bachelor’s degree in one of the chemical, physical, or biological sciences and, in addition, has at least 1 year of pertinent full-time laboratory experience or training, or both, in the specialty or subspecialty in which the individual performs tests; or

(4)(i) Have successfully completed 3 years (90 semester hours or equivalent) in an accredited college or university with the following distribution of courses—

(A) For those whose training was completed before September 15, 1963. At least 24 semester hours in chemistry and biology courses of which—

(1) At least 6 semester hours were in inorganic chemistry and at least 3 semester hours were in other chemistry courses; and

(2) At least 12 semester hours in biology courses pertinent to the medical sciences; or

(B) For those whose training was completed after September 14, 1963.

(1) 16 semester hours in chemistry courses that included at least 6 semester hours in inorganic chemistry and that are acceptable toward a major in chemistry;

(2) 16 semester hours in biology courses that are pertinent to the medical sciences and are acceptable toward a major in the biological sciences; and

(3) 3 semester hours of mathematics;

(ii) Has experience, training, or both, covering several fields of medical laboratory work of at least 1 year and of such quality as to provide him or her with education and training in medical technology equivalent to that described in paragraphs (b)(1) and (2) of this section; or

(5) With respect to individuals first qualifying before July 1, 1971, the technologist—

(i) Was performing the duties of a laboratory technologist at any time between July 1, 1961, and January 1, 1968, and

(ii) Has had at least 10 years of pertinent laboratory experience prior to January 1, 1968. (This required experience may be met by the substitution of education for experience); or

(6) Achieves a satisfactory grade in a proficiency examination approved by HHS.

[58 FR 39155, July 22, 1993]

§ 493.1495 Standard; Testing personnel responsibilities.

The testing personnel are responsible for specimen processing, test performance and for reporting test results.

(a) Each individual performs only those high complexity tests that are authorized by the laboratory director and require a degree of skill commensurate with the individual’s education, training or experience, and technical abilities.

(b) Each individual performing high complexity testing must—

(1) Follow the laboratory’s procedures for specimen handling and processing, test analyses, reporting and maintaining records of patient test results;

(2) Maintain records that demonstrate that proficiency testing samples are tested in the same manner as patient specimens;

(3) Adhere to the laboratory’s quality control policies, document all quality control activities, instrument and procedural calibrations and maintenance performed;

(4) Follow the laboratory’s established policies and procedures whenever test systems are not within the laboratory’s established acceptable levels of performance;

(5) Be capable of identifying problems that may adversely affect test performance or reporting of test results and either must correct the problems or immediately notify the general supervisor, technical supervisor, clinical consultant, or director;
§ 493.1701 Condition: Quality assurance; moderate complexity (including the subcategory) or high complexity testing, or any combination of these tests.

Each laboratory performing moderate complexity (including the subcategory) or high complexity testing, or any combination of these tests, must establish and follow written policies and procedures for a comprehensive quality assurance program that is designed to monitor and evaluate the ongoing and overall quality of the total testing process (preanalytic, analytic, postanalytic). The laboratory’s quality assurance program must evaluate the effectiveness of its policies and procedures; identify and correct problems; assure the accurate, reliable and prompt reporting of test results; and assure the adequacy and competency of the staff. As necessary, the laboratory must revise policies and procedures based upon the results of those evaluations. The laboratory must meet the standards as they apply to the services offered, complexity of testing performed and test results reported, and the unique practices of each testing entity. All quality assurance activities must be documented.

[60 FR 20050, Apr. 24, 1995]

§ 493.1703 Standard; Patient test management assessment.

The laboratory must have an ongoing mechanism for monitoring and evaluating the systems required under subpart J, Patient Test Management. The laboratory must monitor, evaluate, and revise, if necessary, based on the results of its evaluations, the following:

(a) The criteria established for patient preparation, specimen collection, labeling, preservation and transportation;

(b) The information solicited and obtained on the laboratory’s test requisition for its completeness, relevance, and necessity for the testing of patient specimens;

(c) The use and appropriateness of the criteria established for specimen rejection;

(d) The completeness, usefulness, and accuracy of the test report information necessary for the interpretation or utilization of test results;

(e) The timely reporting of test results based on testing priorities (STAT, routine, etc.); and

(f) The accuracy and reliability of test reporting systems, appropriate storage of records and retrieval of test results.

§ 493.1705 Standard; Quality control assessment.

The laboratory must have an ongoing mechanism to evaluate the corrective actions taken under §493.1219, Remedial actions. Ineffective policies and procedures must be revised based on the outcome of the evaluation. The mechanism must evaluate and review the effectiveness of corrective actions taken for—
(a) Problems identified during the evaluation of calibration and control data for each test method;
(b) Problems identified during the evaluation of patient test values for the purpose of verifying the reference range of a test method; and
(c) Errors detected in reported results.

§ 493.1707 Standard; Proficiency testing assessment.
Under subpart H of this part, Proficiency Testing, the corrective actions taken for any unacceptable, unsatisfactory, or unsuccessful proficiency testing result(s) must be evaluated for effectiveness.

§ 493.1709 Standard; Comparison of test results.
(a) If a laboratory performs the same test using different methodologies or instruments, or performs the same test at multiple testing sites, the laboratory must have a system that twice a year evaluates and defines the relationship between test results using the different methodologies, instruments, or testing sites.
(b) If a laboratory performs tests that are not included under subpart I of this part, Proficiency Testing Programs, the laboratory must have a system for verifying the accuracy of its test results at least twice a year.

[58 FR 5236, Jan. 19, 1993]

§ 493.1711 Standard; Relationship of patient information to patient test results.
For internal quality assurance, the laboratory must have a mechanism to identify and evaluate patient test results that appear inconsistent with relevant criteria such as—
(a) Patient age;
(b) Sex;
(c) Diagnosis or pertinent clinical data, when provided;
(d) Distribution of patient test results when available; and
(e) Relationship with other test parameters, when available within the laboratory.

§ 493.1713 Standard; Personnel assessment.
The laboratory must have an ongoing mechanism to evaluate the effectiveness of its policies and procedures for assuring employee competence and, if applicable, consultant competence.

§ 493.1715 Standard; Communications.
The laboratory must have a system in place to document problems that occur as a result of breakdowns in communication between the laboratory and the authorized individual who orders or receives the results of test procedures or examinations. Corrective actions must be taken, as necessary, to resolve the problems and minimize communication breakdowns.

[58 FR 5236, Jan. 19, 1993]

§ 493.1717 Standard; Complaint investigations.
The laboratory must have a system in place to assure that all complaints and problems reported to the laboratory are documented. Investigations of complaints must be made, when appropriate, and, as necessary, corrective actions are instituted.

§ 493.1719 Standard; Quality assurance review with staff.
The laboratory must have a mechanism for documenting and assessing problems identified during quality assurance reviews and discussing them with the staff. The laboratory must take corrective actions that are necessary to prevent recurrences.

§ 493.1721 Standard; Quality assurance records.
The laboratory must maintain documentation of all quality assurance activities including problems identified and corrective actions taken. All quality assurance records must be available to HHS and maintained for a period of 2 years.

[58 FR 5236, Jan. 19, 1993]

Subpart Q—Inspection

SOURCE: 57 FR 7184, Feb. 28, 1992, unless otherwise noted.
§ 493.1771 Condition: Inspection requirements applicable to all CLIA-certified and CLIA-exempt laboratories.

(a) Each laboratory issued a CLIA certificate must meet the requirements in § 493.1773 and the specific requirements for its certificate type, as specified in §§ 493.1775 through 493.1780.

(b) All CLIA-exempt laboratories must comply with the inspection requirements in §§ 493.1773 and 493.1780, when applicable.

§ 493.1773 Standard: Basic inspection requirements for all laboratories issued a CLIA certificate and CLIA-exempt laboratories.

(a) A laboratory issued a certificate must permit HCFA or a HCFA agent to conduct an inspection to assess the laboratory’s compliance with the requirements of this part. A CLIA-exempt laboratory and a laboratory that requests, or is issued a certificate of accreditation, must permit HCFA or a HCFA agent to conduct validation and complaint inspections.

(b) General requirements. As part of the inspection process, HCFA or a HCFA agent may require the laboratory to do the following:

1. Test samples, including proficiency testing samples, or perform procedures.

2. Permit interviews of all personnel concerning the laboratory’s compliance with the applicable requirements of this part.

3. Permit laboratory personnel to be observed performing all phases of the total testing process (preanalytic, analytic, and postanalytic).

4. Permit HCFA or a HCFA agent access to all areas encompassed under the certificate including, but not limited to, the following:

   i. Specimen procurement and processing areas.

   ii. Storage facilities for specimens, reagents, supplies, records, and reports.

   iii. Testing and reporting areas.

5. Provide HCFA or a HCFA agent with copies or exact duplicates of all records and data it requires.

6. Accessible records and data. A laboratory must have all records and data accessible and retrievable within a reasonable time frame during the course of the inspection.

(d) Requirement to provide information and data. A laboratory must provide, upon request, all information and data needed by HCFA or a HCFA agent to make a determination of the laboratory’s compliance with the applicable requirements of this part.

(e) Reinspection. HCFA or a HCFA agent may reinspect a laboratory at any time to evaluate the ability of the laboratory to provide accurate and reliable test results.

(f) Complaint inspection. HCFA or a HCFA agent may conduct an inspection when there are complaints alleging noncompliance with any of the requirements of this part.

(g) Failure to permit an inspection or reinspection. Failure to permit HCFA or a HCFA agent to conduct an inspection or reinspection results in the suspension or cancellation of the laboratory’s participation in Medicare and Medicaid for payment, and suspension or limitation of, or action to revoke the laboratory’s CLIA certificate, in accordance with subpart R of this part.

§ 493.1775 Standard: Inspection of laboratories issued a certificate of waiver or a certificate for provider-performed microscopy procedures.

(a) A laboratory that has been issued a certificate of waiver or a certificate for provider-performed microscopy procedures is not subject to biennial inspections.

(b) If necessary, HCFA or a HCFA agent may conduct an inspection of a laboratory issued a certificate of waiver or a certificate for provider-performed microscopy procedures at any time during the laboratory’s hours of operation to do the following:

1. Determine if the laboratory is operated and testing is performed in a manner that does not constitute an imminent and serious risk to public health.

2. Evaluate a complaint from the public.

3. Determine whether the laboratory is performing tests beyond the scope of the certificate held by the laboratory.
(4) Collect information regarding the appropriateness of tests specified as waived tests or provider-performed microscopy procedures.

(c) The laboratory must comply with the basic inspection requirements of §493.1773.

[63 FR 26737, May 14, 1998]

§ 493.1777 Standard: Inspection of laboratories that have requested or have been issued a certificate of compliance.

(a) Initial inspection. (1) A laboratory issued a registration certificate must permit an initial inspection to assess the laboratory’s compliance with the requirements of this part before HCFA issues a certificate of compliance.

(2) The inspection may occur at any time during the laboratory’s hours of operation.

(b) Subsequent inspections. (1) HCFA or a HCFA agent may conduct subsequent inspections on a biennial basis or with such other frequency as HCFA determines to be necessary to ensure compliance with the requirements of this part.

(2) HCFA bases the nature of subsequent inspections on the laboratory’s compliance history.

(c) Provider-performed microscopy procedures. The inspection sample for review may include testing in the subcategory of provider-performed microscopy procedures.

(d) Compliance with basic inspection requirements. The laboratory must comply with the basic inspection requirements of §493.1773.

[63 FR 26737, May 14, 1998]

§ 493.1780 Standard: Inspection of CLIA-exempt laboratories or laboratories requesting or issued a certificate of accreditation.

(a) Validation inspection. HCFA or a HCFA agent may conduct a validation inspection of any accredited or CLIA-exempt laboratory at any time during its hours of operation.

(b) Complaint inspection. HCFA or a HCFA agent may conduct a complaint inspection of a CLIA-exempt laboratory or a laboratory requesting or issued a certificate of accreditation at any time during its hours of operation upon receiving a complaint applicable to the requirements of this part.

(c) Noncompliance determination. If a validation or complaint inspection results in a finding that the laboratory is not in compliance with one or more condition-level requirements, the following actions occur:

(1) A laboratory issued a certificate of accreditation is subject to a full review by HCFA, in accordance with subpart E of this part and §488.11 of this chapter.

(2) A CLIA-exempt laboratory is subject to appropriate enforcement actions under the approved State licensure program.

(d) Compliance with basic inspection requirements. CLIA-exempt laboratories and laboratories requesting or issued a certificate of accreditation must comply with the basic inspection requirements in §493.1773.

[63 FR 26737, May 14, 1998]

Subpart R—Enforcement Procedures

SOURCE: 57 FR 2737, Feb. 28, 1992, unless otherwise noted.

§ 493.1800 Basis and scope.

(a) Statutory basis. (1) Section 1846 of the Act—

(i) Provides for intermediate sanctions that may be imposed on laboratories that perform clinical diagnostic tests on human specimens when those laboratories are found to be out of compliance with one or more of the conditions for Medicare coverage of their services; and

(ii) Requires the Secretary to develop and implement a range of such sanctions, including four that are specified in the statute.

(2) The Clinical Laboratories Improvement Act of 1967 (section 353 of the Public Health Service Act) as amended by CLIA ’88—

(i) Establishes requirements for all laboratories that perform clinical diagnostic tests on human specimens;

(ii) Requires a Federal certification scheme to be applied to all such laboratories; and

(iii) Grants the Secretary broad enforcement authority, including—
(A) Use of intermediate sanctions;
(B) Suspension, limitation, or revocation of the certificate of a laboratory that is out of compliance with one or more requirements for a certificate; and
(C) Civil suit to enjoin any laboratory activity that constitutes a significant hazard to the public health.

(3) Section 353 also—
(i) Provides for imprisonment or fine for any person convicted of intentional violation of CLIA requirements;
(ii) Specifies the administrative hearing and judicial review rights of a laboratory that is sanctioned under CLIA; and
(iii) Requires the Secretary to publish annually a list of all laboratories that have been sanctioned during the preceding year.

(b) Scope and applicability. This subpart sets forth—
(1) The policies and procedures that HCFA follows to enforce the requirements applicable to laboratories under CLIA and under section 1846 of the Act; and
(2) The appeal rights of laboratories on which HCFA imposes sanctions.

§ 493.1804 General considerations.
(a) Purpose. The enforcement mechanisms set forth in this subpart have the following purposes:
(1) To protect all individuals served by laboratories against substandard testing of specimens.
(2) To safeguard the general public against health and safety hazards that might result from laboratory activities.
(3) To motivate laboratories to comply with CLIA requirements so that they can provide accurate and reliable test results.

(b) Basis for decision to impose sanctions. (1) HCFA’s decision to impose sanctions is based on one or more of the following:
(i) Deficiencies found by HCFA or its agents in the conduct of inspections to certify or validate compliance with Federal requirements, or through review of materials submitted by the laboratory (e.g., personnel qualifications).
(ii) Unsuccessful participation in proficiency testing.

(2) HCFA imposes one or more of the alternative or principal sanctions specified in §§ 493.1806 and 493.1807 when HCFA or HCFA’s agent finds that a laboratory has condition-level deficiencies.

(c) Imposition of alternative sanctions.
(1) HCFA may impose alternative sanctions in lieu of, or in addition to principal sanctions, (HCFA does not impose alternative sanctions on laboratories that have certificates of waiver because those laboratories are not inspected for compliance with condition-level requirements.)

(2) HCFA may impose alternative sanctions other than a civil money penalty after the laboratory has had an opportunity to respond, but before the hearing specified in § 493.1844.

(d) Choice of sanction: Factors considered. HCFA bases its choice of sanction or sanctions on consideration of one or more factors that include, but are not limited to, the following, as assessed by the State or by HCFA, or its agents:
(1) Whether the deficiencies pose immediate jeopardy.
(2) The nature, incidence, severity, and duration of the deficiencies or noncompliance.
(3) Whether the same condition level deficiencies have been identified repeatedly.
(4) The accuracy and extent of laboratory records (e.g., of remedial action) in regard to the noncompliance, and their availability to the State, to other HCFA agents, and to HCFA.
(5) The relationship of one deficiency or group of deficiencies to other deficiencies.
(6) The overall compliance history of the laboratory including but not limited to any period of noncompliance that occurred between certifications of compliance.
(7) The corrective and long-term compliance outcomes that HCFA hopes to achieve through application of the sanction.
(8) Whether the laboratory has made any progress toward improvement following a reasonable opportunity to correct deficiencies.
(9) Any recommendation by the State agency as to which sanction would be appropriate.
§ 493.1810 Imposition and lifting of alternative sanctions.

(a) Notice of noncompliance and of proposed sanction: Content. If HCFA or its agency identifies condition level noncompliance in a laboratory, HCFA or

§ 493.1807 Additional sanctions: Laboratories that participate in Medicare.

The following additional sanctions are available for laboratories that are out of compliance with one or more CLIA conditions and that have approval to receive Medicare payment for their services.

(1) Principal sanction. Cancellation of the laboratory's approval to receive Medicare payment for its services.

(2) Alternative sanctions. (i) Suspension of payment for tests in one or more specific specialties or subspecialties, performed on or after the effective date of sanction.

(ii) Suspension of payment for all tests in all specialties and subspecialties performed on or after the effective date of sanction.

§ 493.1808 Adverse action on any type of CLIA certificate: Effect on Medicare approval.

(a) Suspension or revocation of any type of CLIA certificate. When HCFA suspends or revokes any type of CLIA certificate, HCFA concurrently cancels the laboratory's approval to receive Medicare payment for its services.

(b) Limitation of any type of CLIA certificate. When HCFA limits any type of CLIA certificate, HCFA concurrently limits Medicare approval to only those specialties or subspecialties that are authorized by the laboratory's limited certificate.

§ 493.1809 Limitation on Medicaid payment.

As provided in section 1902(a)(9)(C) of the Act, payment for laboratory services may be made under the State plan only if those services are furnished by a laboratory that has a CLIA certificate or is licensed by a State whose licensure program has been approved by the Secretary under this part.

[57 FR 7237, Feb. 28, 1992; 57 FR 35761, Aug. 11, 1992]
its agent gives the laboratory written notice of the following:

(1) The condition level noncompliance that it has identified.
(2) The sanction or sanctions that HCFA or its agent proposes to impose against the laboratory.
(3) The rationale for the proposed sanction or sanctions.
(4) The projected effective date and duration of the proposed sanction or sanctions.
(5) The authority for the proposed sanction or sanctions.
(6) The time allowed (at least 10 days) for the laboratory to respond to the notice.

(b) Opportunity to respond. During the period specified in paragraph (a)(6) of this section, the laboratory may submit to HCFA or its agent written evidence or other information against the imposition of the proposed sanction or sanctions.

(c) Notice of imposition of sanction—(1) Content. HCFA gives the laboratory written notice that acknowledges any evidence or information received from the laboratory and specifies the following:

(i) The sanction or sanctions to be imposed against the laboratory.
(ii) The authority and rationale for the imposing sanction or sanctions.
(iii) The effective date and duration of sanction.

(2) Timing. (i) If HCFA or its agent determines that the deficiencies pose immediate jeopardy, HCFA provides notice at least 5 days before the effective date of sanction.

(ii) If HCFA or its agent determines that the deficiencies do not pose immediate jeopardy, HCFA provides notice at least 15 days before the effective date of the sanction.

(d) Duration of alternative sanctions. An alternative sanction continues until the earlier of the following occurs:

(1) The laboratory corrects all condition level deficiencies.
(2) HCFA’s suspension, limitation, or revocation of the laboratory’s CLIA certificate becomes effective.

(e) Lifting of alternative sanctions—(1) General rule. Alternative sanctions are not lifted until a laboratory’s compliance with all condition level requirements is verified.
(2) Credible allegation of compliance. When a sanctioned laboratory submits a credible allegation of compliance, HCFA’s agent determines whether—

(i) It can certify compliance on the basis of the evidence presented by the laboratory in its allegation; or
(ii) It must revisit to verify whether the laboratory has, in fact, achieved compliance.

(3) Compliance achieved before the date of revisit. If during a revisit, the laboratory presents credible evidence (as determined by HCFA or its agent) that it achieved compliance before the date of revisit, sanctions are lifted as of that earlier date.

§ 493.1812 Action when deficiencies pose immediate jeopardy.

If a laboratory’s deficiencies pose immediate jeopardy, the following rules apply:

(a) HCFA requires the laboratory to take immediate action to remove the jeopardy and may impose one or more alternative sanctions to help bring the laboratory into compliance.

(b) If the findings of a revisit indicate that a laboratory has not eliminated the jeopardy, HCFA suspends or limits the laboratory’s CLIA certificate no earlier than 5 days after the date of notice of suspension or limitation. HCFA may later revoke the certificate.

(c) In addition, if HCFA has reason to believe that the continuation of any activity by any laboratory (either the entire laboratory operation or any specialty or subspecialty of testing) would constitute a significant hazard to the public health, HCFA may bring suit and seek a temporary injunction or restraining order against continuation of that activity by the laboratory, regardless of the type of CLIA certificate the laboratory has and of whether it is State-exempt.

§ 493.1814 Action when deficiencies are at the condition level but do not pose immediate jeopardy.

If a laboratory has condition level deficiencies that do not pose immediate jeopardy, the following rules apply:
(a) Initial action. (1) HCFA may cancel the laboratory’s approval to receive Medicare payment for its services.

(2) HCFA may suspend, limit, or revoke the laboratory’s CLIA certificate.

(3) If HCFA does not impose a principal sanction under paragraph (a)(1) or (a)(2) of this section, it imposes one or more alternative sanctions. In the case of unsuccessful participation in proficiency testing, HCFA may impose the training and technical assistance requirement set forth at §493.1838 in lieu of, or in addition to, one or more alternative sanctions.

(b) Failure to correct condition level deficiencies. If HCFA imposes alternative sanctions for condition level deficiencies that do not pose immediate jeopardy, and the laboratory does not correct the condition level deficiencies within 12 months after the last day of inspection, HCFA—

(1) Cancels the laboratory’s approval to receive Medicare payment for its services, and discontinues the Medicare payment sanctions as of the day cancellation is effective.

(2) Following a revisit which indicates that the laboratory has not corrected its condition level deficiencies, notifies the laboratory that it proposes to suspend, limit, or revoke the certificate, as specified in §493.1816(b), and the laboratory’s right to a hearing; and

(3) May impose (or continue, if already imposed) any alternative sanctions that do not pertain to Medicare payments. (Sanctions imposed under the authority of section 353 of the PHS Act may continue for more than 12 months from the last date of inspection, while a hearing on the proposed suspension, limitation, or revocation of the certificate of compliance, registration certificate, certificate of accreditation, or certificate for PPM procedures is pending.)

(c) Action after hearing. If a hearing decision upholds a proposed suspension, limitation, or revocation of a laboratory’s CLIA certificate, HCFA discontinues any alternative sanctions as of the day it makes the suspension, limitation, or revocation effective.

§493.1816 Action when deficiencies are not at the condition level.

If a laboratory has deficiencies, that are not at the condition level, the following rules apply:

(a) Initial action. The laboratory must submit a plan of correction that is acceptable to HCFA in content and time frames.

(b) Failure to correct deficiencies. If, on revisit, it is found that the laboratory has not corrected the deficiencies within 12 months after the last day of inspection, the following rules apply:

(1) HCFA cancels the laboratory’s approval to receive Medicare payment for its services.

(2) HCFA notifies the laboratory of its intent to suspend, limit, or revoke the laboratory’s CLIA certificate and of the laboratory’s right to a hearing.

§493.1820 Ensuring timely correction of deficiencies.

(a) Timing of visits. HCFA, the State survey agency or other HCFA agent may visit the laboratory at any time to evaluate progress, and at the end of the period to determine whether all corrections have been made.

(b) Deficiencies corrected before a visit. If during a visit, a laboratory produces credible evidence that it achieved compliance before the visit, the sanctions are lifted as of that earlier date.

(c) Failure to correct deficiencies. If during a visit it is found that the laboratory has not corrected its deficiencies, HCFA may propose to suspend, limit, or revoke the laboratory’s CLIA certificate.

(d) Additional time for correcting lower level deficiencies not at the condition level. If at the end of the plan of correction period all condition level deficiencies have been corrected, and there are deficiencies, that are not at the condition level, HCFA may request a revised plan of correction. The revised plan may not extend beyond 12 months from the last day of the inspection that originally identified the cited deficiencies.

(e) Persistence of deficiencies. If at the end of the period covered by the plan of correction, the laboratory still has deficiencies, the rules of §§493.1814 and 493.1816 apply.
§ 493.1826 Suspension of part of Medicare payments.

(a) Application. (1) HCFA may impose this sanction if a laboratory—
(i) Is found to have condition level deficiencies with respect to one or more specialties or subspecialties of tests; and
(ii) Agrees (in return for not having its Medicare approval cancelled immediately) not to charge Medicare beneficiaries or their private insurance carriers for the services for which Medicare payment is suspended.

(2) HCFA suspends Medicare payment for those specialties or subspecialties of tests for which the laboratory is out of compliance with Federal requirements.

(b) Procedures. Before imposing this sanction, HCFA provides notice of sanction and opportunity to respond in accordance with §493.1810.

(c) Duration and effect of sanction. This sanction continues until the laboratory corrects the condition level deficiencies or HCFA cancels the laboratory's approval to receive Medicare payment for its services, but in no event longer than 12 months.

(1) If the laboratory corrects all condition level deficiencies, HCFA resumes Medicare payment effective for all services furnished on or after the date the deficiencies are corrected.

(2) [Reserved]

[57 FR 7237, Feb. 28, 1992; 57 FR 35761, Aug. 11, 1992]

§ 493.1828 Suspension of all Medicare payments.

(a) Application. (1) HCFA may suspend payment for all Medicare-approved laboratory services when the laboratory has condition level deficiencies.

(2) HCFA suspends payment for all Medicare covered laboratory services when the following conditions are met:

(i) Either—

(A) The laboratory has not corrected its condition level deficiencies included in the plan of correction within 3 months from the last date of inspection; or

(B) The laboratory has been found to have the same condition level deficiencies during three consecutive inspections; and

(ii) The laboratory has chosen (in return for not having its Medicare approval immediately cancelled), to not charge Medicare beneficiaries or their private insurance carriers for services for which Medicare payment is suspended.

(3) HCFA suspends payment for services furnished on and after the effective date of sanction.

(b) Procedures. Before imposing this sanction, HCFA provides notice of sanction and opportunity to respond in accordance with §493.1810.

(c) Duration and effect of sanction. (1) Suspension of payment continues until all condition level deficiencies are corrected, but never beyond twelve months.

(2) If all the deficiencies are not corrected by the end of the 12 month period, HCFA cancels the laboratory's approval to receive Medicare payment for its services.

§ 493.1832 Directed plan of correction and directed portion of a plan of correction.

(a) Application. HCFA may impose a directed plan of correction as an alternative sanction for any laboratory that has condition level deficiencies. If HCFA does not impose a directed plan of correction as an alternative sanction for a laboratory that has condition level deficiencies, it at least imposes a directed portion of a plan of correction when it imposes any of the following alternative sanctions:

(1) State onsite monitoring.

(2) Civil money penalty.

(3) Suspension of all or part of Medicare payments.

(b) Procedures—(1) Directed plan of correction. When imposing this sanction, HCFA—

(i) Gives the laboratory prior notice of the sanction and opportunity to respond in accordance with §493.1810;

(ii) Directs the laboratory to take specific corrective action within specific time frames in order to achieve compliance; and

(iii) May direct the laboratory to submit the names of laboratory clients for notification purposes, as specified in paragraph (b)(3) of this section.

(2) Directed portion of a plan of correction. HCFA may decide to notify clients
of a sanctioned laboratory, because of the seriousness of the noncompliance (e.g., the existence of immediate jeopardy) or for other reasons. When imposing this sanction, HCFA takes the following steps—

(i) Directs the laboratory to submit to HCFA, the State survey agency, or other HCFA agent, within 10 calendar days after the notice of the alternative sanction, a list of names and addresses of all physicians, providers, suppliers, and other clients who have used some or all of the services of the laboratory since the last certification inspection or within any other timeframe specified by HCFA.

(ii) Within 30 calendar days of receipt of the information, may send to each laboratory client, via the State survey agency, a notice containing the name and address of the laboratory, the nature of the laboratory's noncompliance, and the kind and effective date of the alternative sanction.

(iii) Sends to each laboratory client, via the State survey agency, notice of the rescission of an adverse action within 30 days of the rescission.

§ 493.1834 Civil money penalty.

(a) Statutory basis. Sections 1846 of the Act and 353(h)(2)(B) of the PHS Act authorize the Secretary to impose civil money penalties on laboratories. Section 1846(b)(3) of the Act specifically provides that incrementally more severe fines may be imposed for repeated or uncorrected deficiencies.

(b) Scope. This section sets forth the procedures that HCFA follows to impose a civil money penalty in lieu of, or in addition to, suspending, limiting, or revoking the certificate of compliance, registration certificate, certificate of accreditation, or certificate for PPM procedures of a laboratory that is found to have condition level deficiencies.

(c) Basis for imposing a civil money penalty. HCFA may impose a civil money penalty against any laboratory determined to have condition level deficiencies regardless of whether those deficiencies pose immediate jeopardy.

(d) Amount of penalty—(1) Factors considered. In determining the amount of the penalty, HCFA takes into account the following factors:

(i) The nature, scope, severity, and duration of the noncompliance.

(ii) Whether the same condition level deficiencies have been identified during three consecutive inspections.

(iii) The laboratory's overall compliance history including but not limited to any period of noncompliance that occurred between certifications of compliance.

(iv) The laboratory's intent or reason for noncompliance.

(v) The accuracy and extent of laboratory records and their availability to HCFA, the State survey agency, or other HCFA agent.

(2) Range of penalty amount.

(i) For a condition level deficiency that poses immediate jeopardy, the range is $3,050-$10,000 per day of noncompliance or per violation.

(ii) For a condition level deficiency that does not pose immediate jeopardy, the range is $50-$3,000 per day of noncompliance or per violation.

(3) Decreased penalty amounts. If the immediate jeopardy is removed, but the deficiency continues, HCFA shifts the penalty amount to the lower range.

(4) Increased penalty amounts. HCFA may, before the hearing, propose to increase the penalty amount for a laboratory that has deficiencies which, after imposition of a lower level penalty
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amount, become sufficiently serious to pose immediate jeopardy.

(e) Procedures for imposition of civil money penalty—(1) Notice of intent. (i) HCFA sends the laboratory written notice of HCFA's intent to impose a civil money penalty.

(ii) The notice includes the following information:

(A) The statutory basis for the penalty.

(B) The proposed daily or per violation amount of the penalty.

(C) The factors (as described in paragraph (d)(1) of this section) that HCFA considered.

(D) The opportunity for responding to the notice in accordance with § 493.1810(c).

(E) A specific statement regarding the laboratory's appeal rights.

(2) Appeal rights. (i) The laboratory has 60 days from the date of receipt of the notice of intent to impose a civil money penalty to request a hearing in accordance with § 493.1844(g).

(ii) If the laboratory requests a hearing, all other pertinent provisions of § 493.1844 apply.

(iii) If the laboratory does not request a hearing, HCFA may reduce the proposed penalty amount by 35 percent.

(f) Accrual and duration of penalty—(1) Accrual of penalty. The civil money penalty begins accruing as follows:

(i) 5 days after notice of intent if there is immediate jeopardy.

(ii) 15 days after notice of intent if there is not immediate jeopardy.

(2) Duration of penalty. The civil money penalty continues to accrue until the earliest of the following occurs:

(i) The laboratory's compliance with condition level requirements is verified on the basis of the evidence presented by the laboratory in its credible allegation of compliance or at the time or revisist.

(ii) Based on credible evidence presented by the laboratory at the time of revisist, HCFA determines that compliance was achieved before the revisist. (In this situation, the money penalty stops accruing as of the date of compliance.)

(iii) HCFA suspends, limits, or revokes the laboratory's certificate of accreditation, or certificate for PPM procedures.

(g) Computation and notice of total penalty amount—(1) Computation. HCFA computes the total penalty amount after the laboratory's compliance is verified or HCFA suspends, limits, or revokes the laboratory's CLIA certificate but in no event before—

(i) The 60 day period for requesting a hearing has expired without a request or the laboratory has explicitly waived its right to a hearing; or

(ii) Following a hearing requested by the laboratory, the ALJ issues a decision that upholds imposition of the penalty.

(2) Notice of penalty amount and due date of penalty. The notice includes the following information:

(i) Daily or per violation penalty amount.

(ii) Number of days or violations for which the penalty is imposed.

(iii) Total penalty amount.

(iv) Due date for payment of the penalty.

(h) Due date for payment of penalty. (1) Payment of a civil money penalty is due 15 days from the date of the notice specified in paragraph (g)(2) of this section.

(2) HCFA may approve a plan for a laboratory to pay a civil money penalty, plus interest, over a period of up to one year from the original due date.

(i) Collection and settlement—(1) Collection of penalty amounts. (i) The determined penalty amount may be deducted from any sums then or later owing by the United States to the laboratory subject to the penalty.

(ii) Interest accrues on the unpaid balance of the penalty, beginning on the due date. Interest is computed at the rate specified in § 405.378(d) of this chapter.

(ii) Settlement. HCFA has authority to settle any case at any time before the ALJ issues a hearing decision.


§ 493.1836 State onsite monitoring.

(a) Application. (1) HCFA may require continuous or intermittent monitoring of a plan of correction by the State.
§ 493.1840 Suspension, limitation, or revocation of any type of CLIA certificate.

(a) Adverse action based on actions of the laboratory’s owner, operator or employee. HCFA may initiate adverse action to suspend, limit or revoke any CLIA certificate if HCFA finds that a laboratory’s owner or operator or one of its employees has—

(1) Been guilty of misrepresentation in obtaining a CLIA certificate;

(2) Performed, or represented the laboratory as entitled to perform, a laboratory examination or other procedure that is not within a category of laboratory examinations or other procedures authorized by its CLIA certificate;

(3) Failed to comply with the certificate requirements and performance standards;

(4) Failed to comply with reasonable requests by HCFA for any information or work on materials that HCFA concludes is necessary to determine the laboratory’s continued eligibility for its CLIA certificate or continued compliance with performance standards set by HCFA;

(5) Refused a reasonable request by HCFA or its agent for permission to inspect the laboratory and its operation and pertinent records during the hours that the laboratory is in operation;

(6) Violated or aided and abetted in the violation of any provisions of CLIA and its implementing regulations;

(7) Failed to comply with an alternative sanction imposed under this subpart; or

(8) Within the preceding two-year period, owned or operated a laboratory that had its CLIA certificate revoked.

(This provision applies only to the owner or operator, not to all of the laboratory’s employees.)
§ 493.1842  Cancellation of Medicare approval.

(a) Basis for cancellation. (1) HCFA always cancels a laboratory’s approval to receive Medicare payment for its services if HCFA suspends or revokes the laboratory’s CLIA certificate.

(2) HCFA may cancel the laboratory’s approval under any of the following circumstances:

(i) The laboratory is out of compliance with a condition level requirement.

(ii) The laboratory fails to submit a plan of correction satisfactory to HCFA.

(iii) The laboratory fails to correct all its deficiencies within the time frames specified in the plan of correction.

(b) Notice and opportunity to respond. Before canceling a laboratory’s approval to receive Medicare payment for its services, HCFA gives the laboratory—

(1) Written notice of the rationale for, effective date, and effect of, cancellation;

(2) Opportunity to submit written evidence or other information against cancellation of the laboratory’s approval.

This sanction may be imposed before the hearing that may be requested by a laboratory, in accordance with the appeals procedures set forth in § 493.1844.

(c) Effect of cancellation. Cancellation of Medicare approval terminates any Medicare payment sanctions regardless of the time frames originally specified.

§ 493.1844  Appeals procedures.

(a) General rules. (1) The provisions of this section apply to all laboratories and prospective laboratories that are dissatisfied with any initial determination under paragraph (b) of this section.

(2) Hearings are conducted in accordance with procedures set forth in subpart D of part 498 of this chapter, except that the authority to conduct hearings and issue decisions may be exercised by ALJs assigned to, or detailed to, the Departmental Appeals Board.

(3) Any party dissatisfied with a hearing decision is entitled to request review of the decision as specified in subpart E of part 498 of this chapter, except that the authority to review the decision may be exercised by the Departmental Appeals Board.

(4) When more than one of the actions specified in paragraph (b) of this section are carried out concurrently, the laboratory has a right to only one hearing on all matters at issue.
(b) Actions that are initial determinations. The following actions are initial determinations and therefore are subject to appeal in accordance with this section:

1. The suspension, limitation, or revocation of the laboratory's CLIA certificate by HCFA because of non-compliance with CLIA requirements.
2. The denial of a CLIA certificate.
3. The imposition of alternative sanctions under this subpart (but not the determination as to which alternative sanction or sanctions to impose).
4. The denial or cancellation of the laboratory's approval to receive Medicare payment for its services.

(c) Actions that are not initial determinations. Actions that are not listed in paragraph (b) of this section are not initial determinations and therefore are not subject to appeal under this section. They include, but are not necessarily limited to, the following:

1. The finding that a laboratory accredited by a HCFA-approved accreditation organization is no longer deemed to meet the conditions set forth in subparts H, J, K, M, P, and Q of this part. However, the suspension, limitation or revocation of a certificate of accreditation is an initial determination and is appealable.
2. The finding that a laboratory determined to be in compliance with condition-level requirements but has deficiencies that are not at the condition level.
3. The determination not to reinstate a suspended CLIA certificate because the reason for the suspension has not been removed or there is insufficient assurance that the reason will not recur.
4. The determination as to which alternative sanction or sanctions to impose, including the amount of a civil money penalty to impose per day or per violation.
5. The denial of approval for Medicare payment for the services of a laboratory that does not have in effect a valid CLIA certificate.
6. The determination that a laboratory's deficiencies pose immediate jeopardy.
7. The amount of the civil money penalty assessed per day or for each violation of Federal requirements.

(d) Effect of pending appeals—(1) Alternative sanctions. The effective date of an alternative sanction (other than a civil money penalty) is not delayed because the laboratory has appealed and the hearing or the hearing decision is pending.

(2) Suspension, limitation, or revocation of a laboratory's CLIA certificate—(i) General rule. Except as provided in paragraph (d)(2)(ii) of this section, suspension, limitation, or revocation of a CLIA certificate is not effective until after a hearing decision by an ALJ is issued.

(ii) Exceptions. (A) If HCFA determines that conditions at a laboratory pose immediate jeopardy, the effective date of the suspension or limitation of a CLIA certificate is not delayed because the laboratory has appealed and the hearing or the hearing decision is pending.

(B) HCFA may suspend or limit a laboratory's CLIA certificate before an ALJ hearing or hearing decision if the laboratory has refused a reasonable request for information (including but not limited to billing information), or for work on materials, or has refused permission for HCFA or a HCFA agent to inspect the laboratory or its operation.

(3) Cancellation of Medicare approval. The effective date of the cancellation of a laboratory's approval to receive Medicare payment for its services is not delayed because the laboratory has appealed and the hearing or hearing decision is pending.

(4) Effect of ALJ decision. (i) An ALJ decision is final unless, as provided in paragraph (a)(3) of this section, one of the parties requests review by the Departmental Appeals Board within 60 days, and the Board reviews the case and issues a revised decision.

(ii) If an ALJ decision upholds a suspension imposed because of immediate jeopardy, that suspension becomes a revocation.

(e) Appeal rights for prospective laboratories—(1) Reconsideration. Any prospective laboratory dissatisfied with a denial of a CLIA certificate, or of approval for Medicare payment for its
services, may initiate the appeals process by requesting reconsideration in accordance with §§498.22 through 498.25 of this chapter.

(2) Notice of reopening. If HCFA reopens an initial or reconsidered determination, HCFA gives the prospective laboratory notice of the revised determination in accordance with §498.30 of this chapter.

(3) ALJ hearing. Any prospective laboratory dissatisfied with a reconsidered determination under paragraph (e)(1) of this section, or a revised reconsidered determination under §498.30 of this chapter, is entitled to a hearing before an ALJ, as specified in paragraph (a)(2) of this section.

(4) Review of ALJ hearing decisions. Any prospective laboratory that is dissatisfied with an ALJ’s hearing decision or dismissal of a request for hearing may file a written request for review by the Departmental Appeals Board as provided in paragraph (a)(3) of this section.

(f) Appeal rights of laboratories—(1) ALJ hearing. Any laboratory dissatisfied with the suspension, limitation, or revocation of its CLIA certificate, with the imposition of an alternative sanction under this subpart, or with cancellation of the approval to receive Medicare payment for its services, is entitled to a hearing before an ALJ as specified in paragraph (a)(2) of this section and has 60 days from the notice of sanction to request a hearing.

(2) Review of ALJ hearing decisions. Any laboratory that is dissatisfied with an ALJ’s hearing decision or dismissal of a request for hearing may file a written request for review by the Departmental Appeals Board, as provided in paragraph (a)(3) of this section.

(3) Judicial review. Any laboratory dissatisfied with the decision to impose a civil money penalty or to suspend, limit, or revoke its CLIA certificate may, within 60 days after the decision becomes final, file with the U.S. Court of Appeals of the circuit in which the laboratory has its principal place of business, a petition for judicial review.

(g) Notice of adverse action. (1) If HCFA suspends, limits, or revokes a laboratory’s CLIA certificate or cancels the approval to receive Medicare payment for its services, HCFA gives notice to the laboratory, and may give notice to physicians, providers, suppliers, and other laboratory clients, according to the procedures set forth at §493.1832. In addition, HCFA notifies the general public each time one of these principal sanctions is imposed.

(2) The notice to the laboratory—

(i) Sets forth the reasons for the adverse action, the effective date and effect of that action, and the appeal rights if any; and

(ii) When the certificate is limited, specifies the specialties or subspecialties of tests that the laboratory is no longer authorized to perform, and that are no longer covered under Medicare.

(3) The notice to other entities includes the same information except the information about the laboratory’s appeal rights.

(h) Effective date of adverse action. (1) When the laboratory’s deficiencies pose immediate jeopardy, the effective date of the adverse action is at least 5 days after the date of the notice.

(2) When HCFA determines that the laboratory’s deficiencies do not pose immediate jeopardy, the effective date of the adverse action is at least 15 days after the date of the notice.

§493.1846 Civil action.

If HCFA has reason to believe that continuation of the activities of any laboratory, including a State-exempt laboratory, would constitute a significant hazard to the public health, HCFA may bring suit in a U.S. District Court to enjoin continuation of the specific activity that is causing the hazard or to enjoin the continued operation of the laboratory if HCFA deems it necessary. Upon proper showing, the court shall issue a temporary injunction or restraining order without bond against continuation of the activity.

§493.1850 Laboratory registry.

(a) Once a year HCFA makes available to physicians and to the general public specific information (including information provided to HCFA by the OIG) that is useful in evaluating the performance of laboratories, including the following:

VerDate 11<MAY>2000 12:19 Oct 31, 2000 Jkt 190166 PO 00000 Frm 00998 Fmt 8010 Sfmt 8010 Y:\SGML\190166T.XXX pfrm02 PsN: 190166T
§ 493.2001 Establishment and function of the Clinical Laboratory Improvement Advisory Committee.

(a) HHS will establish a Clinical Laboratory Improvement Advisory Committee to advise and make recommendations on technical and scientific aspects of the provisions of this part 493.

(b) The Clinical Laboratory Improvement Advisory Committee will be comprised of individuals involved in the provision of laboratory services, utilization of laboratory services, development of laboratory testing or methodology, and others as approved by HHS.

(c) HHS will designate specialized subcommittees as necessary.

(d) The Clinical Laboratory Improvement Advisory Committee or any designated subcommittees will meet as needed, but not less than once each year.

(e) The Clinical Laboratory Improvement Advisory Committee or subcommittee, at the request of HHS, will review and make recommendations concerning:

(1) Criteria for categorizing tests and examinations of moderate complexity (including the subcategory) and high complexity;

(2) Determination of waived tests;

(3) Personnel standards;

(4) Patient test management, quality control, quality assurance standards;

(5) Proficiency testing standards;

(6) Applicability to the standards of new technology; and

(7) Other issues relevant to part 493, if requested by HHS.

(f) HHS will be responsible for providing the data and information, as necessary, to the members of the Clinical Laboratory Improvement Advisory Committee.

§ 493.2001 Establishment and function of the Clinical Laboratory Improvement Advisory Committee.

(1) A list of laboratories that have been convicted, under Federal or State laws relating to fraud and abuse, false billing, or kickbacks.

(2) A list of laboratories that have had their CLIA certificates suspended, limited, or revoked, and the reason for the adverse actions.

(3) A list of persons who have been convicted of violating CLIA requirements, as specified in section 353(1) of the PHS Act, together with the circumstances of each case and the penalties imposed.

(4) A list of laboratories on which alternative sanctions have been imposed, showing—

(i) The effective date of the sanctions;

(ii) The reasons for imposing them;

(iii) Any corrective action taken by the laboratory; and

(iv) If the laboratory has achieved compliance, the verified date of compliance.

(5) A list of laboratories whose accreditation has been withdrawn or revoked and the reasons for the withdrawal or revocation.

(6) All appeals and hearing decisions.

(7) A list of laboratories against which HCFA has brought suit under § 493.1846 and the reasons for those actions.

(8) A list of laboratories that have been excluded from participation in Medicare or Medicaid and the reasons for the exclusion.

(b) The laboratory registry is compiled for the calendar year preceding the date the information is made available and includes appropriate explanatory information to aid in the interpretation of the data. It also contains corrections of any erroneous statements or information that appeared in the previous registry.

Subpart S [Reserved]

Subpart T—Consultations

Source: 57 FR 7185, Feb. 28, 1992, unless otherwise noted.
PART 498—APPEALS PROCEDURES FOR DETERMINATIONS THAT AFFECT PARTICIPATION IN THE MEDICARE PROGRAM AND FOR DETERMINATIONS THAT AFFECT THE PARTICIPATION OF ICFs/MR AND CERTAIN NFs IN THE MEDICAID PROGRAM

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S O U R C E : 52 FR 22446, June 12, 1987, unless otherwise noted.

Subpart A—General Provisions

§ 498.1 Statutory basis.
(a) Section 1866(h) of the Act provides for a hearing and for judicial review of the hearing for any institution or agency dissatisfied with a determination that it is not a provider, or with any determination described in section 1866(b)(2) of the Act.
Health Care Financing Administration, HHS § 498.2

(b) Section 1866(b)(2) of the Act lists determinations that serve as a basis for termination of a provider agreement.

c) Sections 1128(a) and (b) of the Act provide for exclusion of certain individuals or entities because of conviction of crimes related to their participation in Medicare and section 1128(f) provides for hearing and judicial review for exclusions.

d) Section 1156 of the Act establishes certain obligations for practitioners and providers of health care services, and provides sanctions and penalties for those that fail to meet those obligations.

e) Section 1128A(c)(2) of the Act provides that the Secretary may not collect a civil money penalty until the affected entity has had notice and opportunity for a hearing.

(f) Although §1866(h) of the Act is silent regarding appeal rights for suppliers and practitioners, the rules in this part include procedures for review of determinations that affect those two groups.

(g) Section 1128A(c)(2) of the Act provides that the Secretary may not collect a civil money penalty until the affected entity has had notice and opportunity for a hearing.

(h) Section 1128A(c)(2) of the Act provides that the Secretary may not collect a civil money penalty until the affected entity has had notice and opportunity for a hearing.

(i) Section 1819(h) of the Act—

(1) Provides that, for SNFs found to be out of compliance with the requirements for participation, specified remedies may be imposed instead of, or in addition to, termination of the facility’s Medicare provider agreement; and

(2) Makes certain provisions of section 1128A of the Act applicable to civil money penalties imposed on SNFs.

(j) Section 1891(e) of the Act provides that, for home health agencies (HHAs) found to be out of compliance with the conditions of participation, specified remedies may be imposed instead of, or in addition to, termination of the HHA’s Medicare provider agreement.

(k) Section 1891(f) of the Act—

(1) Requires the Secretary to develop a range of such remedies; and

(2) Makes certain provisions of section 1128A of the Act applicable to civil money penalties imposed on HHAs.

§ 498.2 Definitions.

As used in this part—

Affected party means a provider, prospective provider, supplier, prospective supplier, or practitioner that is affected by an initial determination or by any subsequent determination or decision issued under this part, and “party” means the affected party or HCFA (or the OIG, as appropriate).

ALJ stands for Administrative Law Judge.

Departmental Appeals Board or Board means a Board established in the Office of the Secretary to provide impartial review of disputed decisions made by the operating components of the Department.

OHA stands for the Social Security Administration’s Office of Hearings and Appeals.

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

Provider means a hospital, critical access hospital (CAH), skilled nursing facility (SNF), comprehensive outpatient rehabilitation facility (CORF), home health agency (HHA), or hospice, that has in effect an agreement to participate in Medicare, that has in effect an agreement to participate in Medicaid, or a clinic, rehabilitation agency, or public health agency that has a similar agreement but only to furnish outpatient physical therapy or outpatient speech pathology services, and prospective provider means any of the listed entities that seeks to participate in Medicare as a provider or to have any facility or organization determined to be a department of the provider or provider-based entity under §413.65 of this chapter.

Supplier means an independent laboratory, supplier of portable X-ray services, rural health clinic (RHC), Federally qualified health center (FQHC), ambulatory surgical center (ASC), organ procurement organization (OPO), or end-stage renal disease (ESRD) treatment facility that is approved by HCFA as meeting the conditions for coverage of its services, and prospective supplier means any of the listed entities that seeks to be approved for coverage of its services under Medicare. (However, for purposes of the sanctions and penalties that may be imposed by the OIG, the term
§ 498.3 Scope and applicability.

(a) Scope. (1) This part sets forth procedures for reviewing initial determinations that HCFA makes with respect to the matters specified in paragraph (b) of this section, and that the OIG makes with respect to the matters specified in paragraph (c) of this section. It also specifies, in paragraph (d) of this section, administrative actions that are not subject to appeal under this part.

(2) The determinations listed in this section affect participation in the Medicare program. Many of the procedures of this part also apply to other determinations that do not affect participation in Medicare. Some examples follow:

(i) HCFA’s determination to terminate an NF’s Medicaid provider agreement.

(ii) HCFA’s determination to cancel the approval of an ICF/MR under section 1910(b) of the Act.

(iii) HCFA’s determination, under the Clinical Laboratory Improvement Act (CLIA), to impose alternative sanctions or to suspend, limit, or revoke the certificate of a laboratory even though it does not participate in Medicare.

(3) The following parts of this chapter specify the applicability of the provisions of this part 498 to sanctions or remedies imposed on the indicated entities:

(i) Part 431, subpart D—for nursing facilities (NFs).

(ii) Part 488, subpart E (§488.330(e))—for SNFs and NFs.

(iii) Part 493, subpart R (§493.1844)—for laboratories.

(b) Initial determinations by HCFA. HCFA makes initial determinations with respect to the following matters:

(1) Whether a prospective provider qualifies as a provider.

(2) Whether a prospective department of a provider, remote location of a hospital, satellite facility, or provider-based status under §413.65 of this chapter, or whether such a facility or entity currently treated as a department of a provider, remote location of a hospital, satellite facility, or a provider-based entity no longer qualifies for that status under §413.65 of this chapter.

(3) Whether an institution is a hospital qualified to elect to claim payment for all emergency hospital services furnished in a calendar year.

(4) Whether an institution continues to remain in compliance with the qualifications for claiming reimbursement for all emergency services furnished in a calendar year.

(5) Whether a prospective supplier meets the conditions for coverage of its services as those conditions are set forth elsewhere in this chapter.

(6) Whether the services of a supplier continue to meet the conditions for coverage.

(7) Whether a physical therapist in independent practice or a chiropractor meets the requirements for coverage of his or her services as set forth in subpart D of part 486 of this chapter and §410.22 of this chapter, respectively.

VERDICT 11<MAY>2000 12:19 Oct 31, 2000 Jkt 190166 PO 00000 Frm 01002 Fmt 8010 Sfmt 8010 Y:\SGML\190166T.XXX pfrm02 PsN: 190166T
(8) The termination of a provider agreement in accordance with §499.53 of this chapter, or the termination of a rural health clinic agreement in accordance with §405.2404 of this chapter, or the termination of a Federally qualified health center agreement in accordance with §405.2436 of this chapter.

(9) HCFA's cancellation, under section 1910(b) of the Act, of an ICF/MR's approval to participate in Medicaid.

(10) Whether, for purposes of rate setting and reimbursement, an ESRD treatment facility is considered to be hospital-based or independent.

(11) Whether to deny payment under §409.19 or §409.64 of this chapter, pertaining to cardiac pacemakers and the pacemaker registry.

(12) Whether a hospital, skilled nursing facility, home health agency, or hospice program meets or continues to meet the advance directives requirements specified in subpart I of part 489 of this chapter.

(13) With respect to an SNF or NF, a finding of noncompliance that results in the imposition of a remedy specified in §488.406 of this chapter, except the State monitoring remedy.

(14) The level of noncompliance found by HCFA in a SNF or NF but only if a successful challenge on this issue would affect—

(i) The range of civil money penalty amounts that HCFA could collect (The scope of review during a hearing on imposition of a civil money penalty is set forth in §488.438(e) of this chapter); or

(ii) A finding of substandard quality of care that results in the loss of approval for a SNF or NF of its nurse aide training program.

(15) The effective date of a Medicare provider agreement or supplier approval.

(16) The finding of substandard quality of care that leads to the loss by a SNF or NF of the approval of its nurse aide training program.

(c) Initial determinations by the OIG. The OIG makes initial determinations with respect to the following matters:

(1) The termination of a provider agreement in accordance with part 1001, subpart C of this title.

(2) The suspension or exclusion from coverage and the denial of reimburse-

ment for services furnished by a provider, practitioner, or supplier, because of fraud or abuse, or conviction of crimes related to participation in the program, in accordance with part 1001, subpart B of this title.

(3) The imposition of sanctions in accordance with part 1004 of this title.

(d) Administrative actions that are not initial determinations. Administrative actions that are not initial determinations (and therefore not subject to appeal under this part) include but are not limited to the following:

(1) The finding that a provider or supplier determined to be in compliance with the conditions or requirements for participation or for coverage has deficiencies.

(2) The finding that a prospective provider does not meet the conditions of participation set forth elsewhere in this chapter, if the prospective provider is, nevertheless, approved for participation in Medicare on the basis of special access certification, as provided in subpart B of part 488 of this chapter.

(3) The refusal to enter into a provider agreement because the prospective provider is unable to give satisfactory assurance of compliance with the requirements of title XVIII of the Act.

(4) The finding that an entity that had its provider agreement terminated may not file another agreement because the previous agreement have not been removed or there is insufficient assurance that the reason for the exclusion will not recur.

(5) The determination not to reinstate a suspended or excluded practitioner, provider, or supplier because the reason for the suspension or exclusion has not been removed, or there is insufficient assurance that the reason will not recur.

(6) The finding that the services of a laboratory are covered as hospital services or as physician's services, rather than as services of an independent laboratory, because the laboratory is not independent of the hospital or of the physician's office.

(7) The refusal to accept for filing an election to claim payment for all emergency hospital services furnished in a calendar year because the institution—
(i) Had previously charged an individual or other person for services furnished during that calendar year;
(ii) Submitted the election after the close of that calendar year; or
(iii) Had previously been notified of its failure to continue to comply.
(8) The finding that the reason for the revocation of a supplier’s right to accept assignment has not been removed or there is insufficient assurance that the reason will not recur.
(9) The finding that a hospital accredited by the Joint Commission on Accreditation of Hospitals or the American Osteopathic Association is not in compliance with a condition of participation, and a finding that that hospital is no longer deemed to meet the conditions of participation.
(10) With respect to an SNF or NF—
(i) The finding that the SNF’s or NF’s deficiencies posed immediate jeopardy to the health or safety of its residents;
(ii) Except as provided in paragraph (b)(13) of this section, a determination by HCFA as to the facility’s level of noncompliance; and
(iii) The imposition of State monitoring.
(11) The choice of alternative sanction or remedy to be imposed on a provider.
(12) The determination that the accreditation requirements of a national accreditation organization do not provide (or do not continue to provide) reasonable assurance that the entities accredited by the accreditation organization meet the applicable long-term care requirements, conditions for coverage, conditions of certification, conditions of participation, or CLIA condition level requirements.
(13) The determination that requirements imposed on a State’s laboratories under the laws of that State do not provide (or do not continue to provide) reasonable assurance that laboratories licensed or approved by the State meet applicable CLIA requirements.
(14) The choice of alternative sanction or remedy to be imposed on a provider or supplier.
(15) A decision by the State survey agency as to when to conduct an initial survey of a prospective provider or supplier.

(e) Exclusion of civil rights issues. The procedures in this subpart do not apply to the adjudication of issues relating to a provider’s compliance with civil rights requirements that are set forth in part 489 of this chapter. Those issues are handled through the Department’s Office of Civil Rights.

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§ 498.4 NFs subject to appeals process in part 498.

A NF is considered a provider for purposes of this part when it has in effect an agreement to participate in Medicaid, including an agreement to participate in both Medicaid and Medicare and it is a—
(a) State-operated NF; or
(b) Non State-operated NF that is subject to compliance action as a result of—
(1) A validation survey by HCFA; or
(2) HCFA’s review of the State’s survey findings.

§ 498.5 Appeal rights.

(a) Appeal rights of prospective providers. (1) Any prospective provider dissatisfied with an initial determination or revised initial determination that it does not qualify as a provider may request reconsideration in accordance with §498.22(a).
(2) Any prospective provider dissatisfied with a reconsidered determination under paragraph (a)(1) of this section, or a revised reconsidered determination under §498.30, is entitled to a hearing before an ALJ.
§ 498.10 Appeal rights of providers and prospective providers.

(a) Any provider dissatisfied with an initial determination to terminate its provider agreement is entitled to a hearing before an ALJ.

(b) Appeal rights of providers. Any provider dissatisfied with an initial determination to terminate its provider agreement is entitled to a hearing before an ALJ.

(c) Appeal rights of providers and prospective providers. Any provider or prospective provider dissatisfied with a hearing decision may request Departmental Appeals Board review, and has a right to seek judicial review of the Board’s decision.

(d) Appeal rights of prospective suppliers. (1) Any prospective supplier dissatisfied with an initial determination or a revised initial determination that its services do not meet the conditions for coverage may request reconsideration in accordance with § 498.22(a).

(2) Any prospective supplier dissatisfied with a reconsidered determination under paragraph (d)(1) of this section, or a revised reconsidered determination under § 498.30, is entitled to a hearing before an ALJ.

(e) Appeal rights of suppliers. Any supplier dissatisfied with an initial determination that the services subject to the determination no longer meet the conditions for coverage, is entitled to a hearing before an ALJ.

(f) Appeal rights of suppliers and prospective suppliers. (1) Any supplier or prospective supplier dissatisfied with the hearing decision may request Departmental Appeals Board review of the ALJ’s decision.

(2) Suppliers and prospective suppliers do not have a right to judicial review except as provided in paragraph (i) of this section.

(g) Appeal rights for certain practitioners. A physical therapist in independent practice or a chiropractor dissatisfied with a determination that he or she does not meet the requirements for coverage of his or her services has the same appeal rights as suppliers have under paragraphs (d), (e) and (f) of this section.

(h) Appeal rights for nonparticipating hospitals that furnish emergency services. A nonparticipating hospital dissatisfied with a determination or decision that it does not qualify to elect to claim payment for all emergency services furnished during a calendar year has the same appeal rights that providers have under paragraph (a), (b), and (c) of this section.

(i) Appeal rights for suspended or excluded practitioners, providers, or suppliers. (1) Any practitioner, provider, or supplier who has been suspended, or whose services have been excluded from coverage in accordance with § 498.3(c)(2), or has been sanctioned in accordance with § 498.3(c)(3), is entitled to a hearing before an ALJ.

(2) Any suspended or excluded practitioner, provider, or supplier dissatisfied with a hearing decision may request Departmental Appeals Board review and has a right to seek judicial review of the Board’s decision by filing an action in Federal district court.

(j) Appeal rights for Medicaid ICF/MR terminated by HCFA. (1) Any Medicaid ICF/MR that has had its approval cancelled by HCFA in accordance with § 498.3(b)(8) has a right to a hearing before an ALJ, to request Departmental Appeals Board review of the hearing decision, and to seek judicial review of the Board’s decision.

(2) The Medicaid agreement remains in effect until the period for requesting a hearing has expired or, if the facility requests a hearing, until a hearing decision is issued, unless HCFA—

(i) Makes a written determination that continuation of provider status for the SNF or ICF constitutes an immediate and serious threat to the health and safety of patients and specifies the reasons for that determination; and

(ii) Certifies that the facility has been notified of its deficiencies and has failed to correct them.

(k) Appeal rights of NFs. Under the circumstances specified in § 431.153(g) and (h) of this chapter, an NF has a right to a hearing before an ALJ, to request Board review of the hearing decision, and to seek judicial review of the Board’s decision.

§ 498.11 Authority of representatives.

(a) A representative appointed and qualified in accordance with § 498.10 may, on behalf of the represented party—

(1) Give and accept any notice or request pertinent to the proceedings set forth in this part;

(2) Present evidence and allegations as to facts and law in any proceedings affecting that party to the same extent as the party; and

(3) Obtain information to the same extent as the party.

(b) A notice or request may be sent to the affected party, to the party’s representative, or to both. A notice or request sent to the representative has the same force and effect as if it had been sent to the party.

§ 498.13 Fees for services of representatives.

Fees for any services performed on behalf of an affected party by an attorney appointed and qualified in accordance with § 498.10 are not subject to the provisions of section 206 of Title II of the Act, which authorizes the Secretary to specify or limit those fees.

§ 498.15 Charge for transcripts.

A party that requests a transcript of prehearing or hearing proceedings or Board review must pay the actual or estimated cost of preparing the transcript unless, for good cause shown by that party, the payment is waived by the ALJ or the Departmental Appeals Board, as appropriate.

§ 498.17 Filing of briefs with the ALJ or Departmental Appeals Board, and opportunity for rebuttal.

(a) Filing of briefs and related documents. If a party files a brief or related document such as a written argument, contention, suggested finding of fact, conclusion of law, or any other written statement, it must submit an original and one copy to the ALJ or the Departmental Appeals Board, as appropriate. The material may be filed by mail or in person and must include a statement certifying that a copy has been furnished to the other party.

(b) Opportunity for rebuttal. (1) The other party will have 20 days from the date of mailing or personal service to submit any rebuttal statement or additional evidence. If a party submits a rebuttal statement or additional evidence, it must file an original and one copy with the ALJ or the Board and furnish a copy to the other party.

(2) The ALJ or the Board will grant an opportunity to reply to the rebuttal statement only if the party shows good cause.

Subpart B—Initial, Reconsidered, and Revised Determinations

§ 498.20 Notice and effect of initial determinations.

(a) Notice of initial determination—(1) General rule. HCFA or the OIG, as appropriate, mails notice of an initial determination to the affected party, setting forth the basis or reasons for the determination, the effect of the determination, and the party’s right to reconsideration, if applicable, or to a hearing.

(2) Special rules: Independent laboratories and suppliers of portable x-ray services. If HCFA determines that an independent laboratory or a supplier of portable x-ray services no longer meets the conditions for coverage of some or all of its services, the notice—

(i) Specifies an effective date of termination of coverage that is at least 15 days after the date of the notice;

(ii) Is also sent to physicians, hospitals, and other parties that might use the services of the laboratory or supplier; and
(iii) In the case of laboratories, specifies the categories of laboratory tests that are no longer covered.

(3) Special rules: Nonparticipating hospitals that elect to claim payment for emergency services. If HCFA determines that a nonparticipating hospital no longer qualifies to elect to claim payment for all emergency services furnished in a calendar year, the notice—
(i) States the calendar year to which the determination applies;
(ii) Specifies an effective date that is at least 5 days after the date of the notice; and
(iii) Specifies that the determination applies to services furnished, in the specified calendar year, to patients accepted (as inpatients or outpatients) on or after the effective date of the determination.

(4) Other special rules. Additional rules pertaining, for example, to content and timing of notice, notice to the public and to other entities, and time allowed for submittal of additional information, are set forth elsewhere in this chapter, as follows:
Part 405 Subpart X—for rural health clinics.
Part 416—for ambulatory surgical centers.
Part 489—for providers, when their provider agreements have been terminated.
Part 1001, Subpart B—for excluded or suspended providers, suppliers, physicians, or practitioners.
Part 1001, Subpart C—for providers, when their provider agreements are terminated by the OIG.
Part 1004—for sanctioned providers and practitioners.

(b) Effect of initial determination. An initial determination is binding unless it is—
(1) Reconsidered in accordance with § 498.24;
(2) Reversed or modified by a hearing decision in accordance with § 498.78; or
(3) Revised in accordance with § 498.32 or § 498.100.

§ 498.24 Reconsidered determination.
When a request for reconsideration has been properly filed in accordance with § 498.22, HCFA—
(a) Receives written evidence and statements that are relevant and material to the matters at issue and are submitted within a reasonable time after the request for reconsideration;
(b) Considers the initial determination, the findings on which the initial
§ 498.25 Notice and effect of reconsidered determination.

(a) Notice. (1) HCFA mails notice of a reconsidered determination to the affected party.
(2) The notice gives the reasons for the determination.
(3) If the determination is adverse, the notice specifies the conditions or requirements of law or regulations that the affected party fails to meet, and informs the party of its right to a hearing.

(b) Effect. A reconsidered determination is binding unless—
(1) HCFA or the OIG, as appropriate, further revises the revised determination; or
(2) The revised determination is reversed or modified by a hearing decision.

Subpart C—Reopening of Initial or Reconsidered Determinations

§ 498.30 Limitation on reopening.

An initial or reconsidered determination that a prospective provider is a provider or that a hospital qualifies to elect to claim payment for all emergency services furnished in a calendar year may not be reopened. HCFA or the OIG, as appropriate, may on its own initiative, reopen any other initial or reconsidered determination, within 12 months after the date of notice of the initial determination.

§ 498.32 Notice and effect of reopening and revision.

(a) Notice. (1) HCFA or the OIG, as appropriate, gives the affected party notice of reopening and of any revision of the reopened determination.
(2) The notice of revised determination states the basis or reason for the revised determination.
(3) If the determination is that a supplier or prospective supplier does not meet the conditions for coverage of its services, the notice specifies the conditions with respect to which the affected party fails to meet the requirements of law and regulations, and informs the party of its right to a hearing.

(b) Effect. A revised determination is binding unless—
(1) The affected party requests a hearing before an ALJ; or
(2) HCFA or the OIG further revises the revised determination.

Subpart D—Hearings

§ 498.40 Request for hearing.

(a) Manner and timing of request. (1) An affected party entitled to a hearing under § 498.5 may file a request for a hearing with HCFA or the OIG, as appropriate, or with OHA.
(2) The affected party or its legal representative or other authorized official must file the request in writing within 60 days from receipt of the notice of initial, reconsidered, or revised determination unless that period is extended in accordance with paragraph (c) of this section. (Presumed date of receipt is determined in accordance with § 498.22(b)(3)).

(b) Content of request for hearing. The request for hearing must—
(1) Identify the specific issues, and the findings of fact and conclusions of law with which the affected party disagrees; and
(2) Specify the basis for contending that the findings and conclusions are incorrect.

(c) Extension of time for filing a request for hearing. If the request was not filed within 60 days—
(1) The affected party or its legal representative or other authorized official may file with the ALJ a written request for extension of time stating the reasons why the request was not filed timely.
(2) For good cause shown, the ALJ may extend the time for filing the request for hearing.
§ 498.42 Parties to the hearing.
   The parties to the hearing are the affected party and HCFA or the OIG, as appropriate.

§ 498.44 Designation of hearing official.
   (a) The Associate Commissioner for Hearings and Appeals, or his or her delegate designates an ALJ or a member or members of the Departmental Appeals Board to conduct the hearing.
   (b) If appropriate, the Associate Commissioner or the delegate may substitute another ALJ or another member or other members of the Departmental Appeals Board to conduct the hearing.
   (c) As used in this part, "ALJ" includes a member or members of the Departmental Appeals Board who are designated to conduct a hearing.

§ 498.45 Disqualification of Administrative Law Judge.
   (a) An ALJ may not conduct a hearing in a case in which he or she is prejudiced or partial to the affected party or has any interest in the matter pending for decision.
   (b) A party that objects to the ALJ designated to conduct the hearing must give notice of its objections at the earliest opportunity.
   (c) The ALJ will consider the objections and decide whether to withdraw or proceed with the hearing.
       (1) If the ALJ withdraws, another will be designated to conduct the hearing.
       (2) If the ALJ does not withdraw, the objecting party may, after the hearing, present its objections to the Departmental Appeals Board as reasons for changing, modifying, or reversing the ALJ's decision or providing a new hearing before another ALJ.

§ 498.47 Prehearing conference.
   (a) At any time before the hearing, the ALJ may call a prehearing conference for the purpose of delineating the issues in controversy, identifying the evidence and witnesses to be presented at the hearing, and obtaining stipulations accordingly.
   (b) On the request of either party or on his or her own motion, the ALJ may adjourn the prehearing conference and reconvene at a later date.

§ 498.48 Notice of prehearing conference.
   (a) Timing of notice. The ALJ will fix a time and place for the prehearing conference and mail written notice to the parties at least 10 days before the scheduled date.
   (b) Content of notice. The notice will inform the parties of the purpose of the conference and specify what issues are sought to be resolved, agreed to, or excluded.
   (c) Additional issues. Issues other than those set forth in the notice of determination or the request for hearing may be considered at the prehearing conference if:
       (1) Either party gives timely notice to that effect to the ALJ and the other party; or
       (2) The ALJ raises the issues in the notice of prehearing conference or at the conference.

§ 498.49 Conduct of prehearing conference.
   (a) The prehearing conference is open to the affected party or its representative, to the HCFA or OIG representatives and their technical advisors, and to any other persons whose presence the ALJ considers necessary or proper.
   (b) The ALJ may accept the agreement of the parties as to the following:
       (1) Facts that are not in controversy.
       (2) Questions that have been resolved favorably to the affected party after the determination in dispute.
       (3) Remaining issues to be resolved.
       (4) The qualifications of those witnesses.
   (c) The ALJ may request the parties to indicate the following:
       (1) The witnesses that will be present to testify at the hearing.
       (2) The nature of other evidence to be submitted.

§ 498.50 Record, order, and effect of prehearing conference.
   (a) Record of prehearing conference. (1) A record is made of all agreements and stipulations entered into at the prehearing conference.
       (2) The record may be transcribed at the request of either party or the ALJ.
§ 498.52 Order and opportunity to object.

(b) Order and opportunity to object. (1) The ALJ issues an order setting forth the results of the prehearing conference, including the agreements made by the parties as to facts not in controversy, the matters to be considered at the hearing, and the issues to be resolved.

(2) Copies of the order are sent to all parties and the parties have 10 days to file objections to the order.

(3) After the 10 days have elapsed, the ALJ settles the order.

(c) Effect of prehearing conference. The agreements and stipulations entered into at the prehearing conference are binding on all parties, unless a party presents facts that, in the opinion of the ALJ, would make an agreement unreasonable or inequitable.

§ 498.53 Time and place of hearing.

(a) The ALJ fixes a time and place for the hearing and gives the parties written notice at least 10 days before the scheduled date.

(b) The notice informs the parties of the general and specific issues to be resolved at the hearing.

§ 498.54 Joint hearings.

When two or more affected parties have requested hearings and the same or substantially similar matters are at issue, the ALJ may, if all parties agree, fix a single time and place for the prehearing conference or hearing and conduct all proceedings jointly. If joint hearings are held, a single record of the proceedings is made and a separate decision issued with respect to each affected party.

§ 498.56 Hearing on new issues.

(a) Basic rules. (1) Within the time limits specified in paragraph (b) of this section, the ALJ may, at the request of either party, or on his or her own motion, provide a hearing on new issues that impinge on the rights of the affected party.

(2) The ALJ may consider new issues even if HCFA or the OIG has not made initial or reconsidered determinations on them, and even if they arose after the request for hearing was filed or after a prehearing conference.

(3) The ALJ may give notice of hearing on new issues at any time after the hearing request is filed and before the hearing record is closed.

(b) Time limits. The ALJ will not consider any issue that arose on or after any of the following dates:

(1) The effective date of the termination of a provider agreement.

(2) The date on which it is determined that a supplier no longer meets the conditions for coverage of its services.

(3) The effective date of the notice to a hospital of its failure to remain in compliance with the qualifications for claiming reimbursement for all emergency services furnished to Medicare beneficiaries during the calendar year.

(4) The effective date of the suspension, or of the exclusion from coverage of services furnished by a suspended or excluded practitioner, provider, or supplier.

(5) With respect to Medicaid SNFs or ICFs surveyed under section 1910(c) of the Act—

(i) The completion date of the survey or resurvey that is the basis for a proposed cancellation of approval; or

(ii) If approval was canceled before the hearings, because of immediate and serious threat to patient health and safety, the effective date of cancellation.

(c) Notice and conduct of hearing on new issues. (1) Unless the affected party waives its right to appear and present evidence, notice of the time and place of hearing on any new issue will be given to the parties in accordance with §498.52.

(2) After giving notice, the ALJ will, except as provided in paragraph (d) of this section, proceed to hearing on new
issues in the same manner as on an issue raised in the request for hearing. 
(d) Remand to HCFA or the OIG. At the request of either party, or on his or 
her own motion, in lieu of a hearing under paragraph (c) of this section, the 
ALJ may remand the case to HCFA or the OIG for consideration of the new 
issue and, if appropriate, a determination. If necessary, the ALJ may direct 
HCFA or the OIG to return the case to the ALJ for further proceedings. 
§ 498.58 Subpoenas.
(a) Basis for issuance. The ALJ, upon 
his or her own motion or at the request 
of a party, may issue subpoenas if they 
are reasonably necessary for the full 
presentation of a case. 
(b) Timing of request by a party. The 
party must file a written request for a 
subpoena with the ALJ at least 5 days 
before the date set for the hearing. 
(c) Content of request. The request 
must:
(1) Identify the witnesses or docu-
ments to be produced; 
(2) Describe their addresses or loca-
tion with sufficient particularity to 
permit them to be found; and 
(3) Specify the pertinent facts the 
party expects to establish by the wit-
nesses or documents, and indicate why 
those facts could not be established 
without use of a subpoena.
(d) Method of issuance. Subpoenas are 
issued in the name of the Secretary, 
who pays the cost of issuance and the 
fees and mileage of any subpoenaed 
witnesses.
§ 498.60 Conduct of hearing.
(a) Participants in the hearing. The hearing is open to the parties and their representatives and technical advisors, and to any other persons whose presence the ALJ considers necessary or proper.
(b) Hearing procedures. (1) The ALJ inquires fully into all of the matters at 
issue, and receives in evidence the testimony of witnesses and any documents that are relevant and material. 
(2) If the ALJ believes that there is relevant and material evidence available which has not been presented at the hearing, he may, at any time be-
fore mailing of notice of the decision, reopen the hearing to receive that evidence. 
(3) The ALJ decides the order in which the evidence and the arguments of the parties are presented and the conduct of the hearing. 
(c) Scope of review: Civil money penalty. In civil money penalty cases— 
(1) The scope of review is as specified in §488.438(e) of this chapter; and 
(2) HCFA’s determination as to the level of noncompliance of an SNF or NF must be upheld unless it is clearly erroneous.
§ 498.66 Waiver of right to appear and present evidence.

(a) Waiver procedures. (1) If an affected party wishes to waive its right to appear and present evidence at the hearing, it must file a written waiver with the ALJ.

(2) If the affected party wishes to withdraw a waiver, it may do so, for good cause, at any time before the ALJ mails notice of the hearing decision.

(b) Effect of waiver. If the affected party waives the right to appear and present evidence, the ALJ need not conduct an oral hearing except in one of the following circumstances:

(1) The ALJ believes that the testimony of the affected party or its representatives or other witnesses is necessary to clarify the facts at issue.

(2) HCFA or the OIG shows good cause for requiring the presentation of oral evidence.

(c) Dismissal for failure to appear. If, despite the waiver, the ALJ sends notice of hearing and the affected party fails to appear, or to show good cause for the failure, the ALJ will dismiss the appeal in accordance with § 498.69.

(d) Hearing without oral testimony. When there is no oral testimony, the ALJ will—

(1) Make a record of the relevant written evidence that was considered in making the determination being appealed, and of any additional evidence submitted by the parties;

(2) Furnish to each party copies of the additional evidence submitted by the other party; and

(3) Give both parties a reasonable opportunity for rebuttal.

(e) Handling of briefs and related statements. If the parties submit briefs or other written statements of evidence or proposed findings of facts or conclusions of law, those documents will be handled in accordance with § 498.17.

§ 498.68 Dismissal of request for hearing.

(a) The ALJ may, at any time before mailing the notice of the decision, dismiss a hearing request if a party withdraws its request for a hearing or the affected party asks that its request be dismissed.

(b) An affected party may request a dismissal by filing a written notice with the ALJ.

§ 498.69 Dismissal for abandonment.

(a) The ALJ may dismiss a request for hearing if it is abandoned by the party that requested it.

(b) The ALJ may consider a request for hearing to be abandoned if the party or its representative—

(1) Fails to appear at the prehearing conference or hearing without having previously shown good cause for not appearing, and

(2) Fails to respond, within 10 days after the ALJ sends a “show cause” notice, with a showing of good cause.

§ 498.70 Dismissal for cause.

On his or her own motion, or on the motion of a party to the hearing, the ALJ may dismiss a hearing request either entirely or as to any stated issue, under any of the following circumstances:

(a) Res judicata. There has been a previous determination or decision with respect to the rights of the same affected party on the same facts and law pertinent to the same issue or issues which has become final either by judicial affirmation or, without judicial consideration, because the affected party did not timely request reconsideration, hearing, or review, or commence a civil action with respect to that determination or decision.

(b) No right to hearing. The party requesting a hearing is not a proper party or does not otherwise have a right to a hearing.

(c) Hearing request not timely filed. The affected party did not file a hearing request timely and the time for filing has not been extended.

§ 498.71 Notice and effect of dismissal and right to request review.

(a) Notice of the ALJ’s dismissal action is mailed to the parties. The notice advises the affected party of its right to request that the dismissal be vacated as provided in § 498.72.

(b) The dismissal of a request for hearing is binding unless it is vacated by the ALJ or the Departmental Appeals Board.

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§ 498.72 Vacating a dismissal of request for hearing.

An ALJ may vacate any dismissal of a request for hearing if a party files a request to that effect within 60 days from receipt of the notice of dismissal and shows good cause for vacating the dismissal. (Date of receipt is determined in accordance with § 498.22(b)(3).)

§ 498.74 Administrative Law Judge's decision.

(a) Timing, basis and content. As soon as practical after the close of the hearing, the ALJ issues a written decision in the case. The decision is based on the evidence of record and contains separate numbered findings of fact and conclusions of law.

(b) Notice and effect. A copy of the decision is mailed to the parties and is binding on them unless—

(1) A party requests review by the Departmental Appeals Board within the time period specified in § 498.82, and the Board reviews the case;

(2) The Departmental Appeals Board denies the request for review and the party seeks judicial review by filing an action in a United States District Court or, in the case of a civil money penalty, in a United States Court of Appeals;

(3) The decision is revised by an ALJ or the Departmental Appeals Board; or

(4) The decision is a recommended decision directed to the Board.

[52 FR 22446, June 12, 1987, as amended at 61 FR 32351, June 24, 1996]

§ 498.76 Removal of hearing to Departmental Appeals Board.

(a) At any time before the ALJ receives oral testimony, the Board may remove to itself any pending request for a hearing.

(b) Notice of removal is mailed to each party.

(c) The Board conducts the hearing in accordance with the rules that apply to ALJ hearings under this subpart.

§ 498.78 Remand by the Administrative Law Judge.

(a) If HCFA or the OIG requests remand, and the affected party concurs in writing or on the record, the ALJ may remand any case properly before him or her to HCFA or the OIG for a determination satisfactory to the affected party.

(b) The ALJ may remand at any time before notice of hearing decision is mailed.

Subpart E—Departmental Appeals Board Review

§ 498.80 Right to request Departmental Appeals Board review of Administrative Law Judge's decision or dismissal.

Either of the parties has a right to request Departmental Appeals Board review of the ALJ's decision or dismissal order, and the parties are so informed in the notice of the ALJ's action.

§ 498.82 Request for Departmental Appeals Board review.

(a) Manner and time of filing. (1) Any party that is dissatisfied with an ALJ's decision or dismissal of a hearing request, may file a written request for review by the Departmental Appeals Board.

(2) The requesting party or its representative or other authorized official must file the request with the OHA within 60 days from receipt of the notice of decision or dismissal, unless the Board, for good cause shown by the requesting party, extends the time for filing. The rules set forth in § 498.40(c) apply to extension of time for requesting Departmental Appeals Board review. (The date of receipt of notice is determined in accordance with § 498.22(c)(3).)

(b) Content of request for review. A request for review of an ALJ decision or dismissal must specify the issues, the findings of fact or conclusions of law with which the party disagrees, and the basis for contending that the findings and conclusions are incorrect.

§ 498.83 Departmental Appeals Board action on request for review.

(a) Request by HCFA or the OIG. The Departmental Appeals Board may dismiss, deny, or grant a request made by HCFA or the OIG for review of an ALJ decision or dismissal.

(b) Request by the affected party. The Board will grant the affected party's request for review unless it dismisses
§ 498.85 Procedures before the Departmental Appeals Board on review.

The parties are given, upon request, a reasonable opportunity to file briefs or other written statements as to fact and law, and to appear before the Departmental Appeals Board to present evidence or oral arguments. Copies of any brief or other written statement must be sent in accordance with § 498.17.

§ 498.86 Evidence admissible on review.

(a) The Departmental Appeals Board may admit evidence into the record in addition to the evidence introduced at the ALJ hearing, (or the documents considered by the ALJ if the hearing was waived), if the Board considers that the additional evidence is relevant and material to an issue before it.

(b) If it appears to the Board that additional relevant evidence is available, the Board will require that it be produced.

(c) Before additional evidence is admitted into the record—

(1) Notice is mailed to the parties (unless they have waived notice) stating that evidence will be received regarding specified issues; and

(2) The parties are given a reasonable time to comment and to present other evidence pertinent to the specified issues.

(d) If additional evidence is presented orally to the Board, a transcript is prepared and made available to any party upon request.

§ 498.88 Decision or remand by the Departmental Appeals Board.

(a) When the Departmental Appeals Board reviews an ALJ’s decision or order of dismissal, or receives a case remanded by a court, the Board may either issue a decision or remand the case to an ALJ for a hearing and decision or a recommended decision for final decision by the Board.

(b) In a remanded case, the ALJ initiates additional proceedings and takes other actions as directed by the Board in its order of remand, and may take other action not inconsistent with that order.

(c) Upon completion of all action called for by the remand order and any other consistent action, the ALJ promptly makes a decision or, as specified by the Board, certifies the case to the Board with a recommended decision.

(d) The parties have 20 days from the date of a notice of a recommended decision to submit to the Board any exception, objection, or comment on the findings of fact, conclusions of law, and recommended decision.

(e) After the 20-day period, the Board issues its decision adopting, modifying or rejecting the ALJ’s recommended decision.

(f) If the Board does not remand the case to an ALJ, the following rules apply:

(i) The Board’s decision—

(ii) Is based upon the evidence in the hearing record and any further evidence that the Board receives during its review;

(iii) Is in writing and contains separate numbered findings of fact and conclusions of law; and

(iv) May modify, affirm, or reverse the ALJ’s decision.

(2) A copy of the Board’s decision is mailed to each party.
§ 498.90 Effect of Departmental Appeals Board decision.

(a) General rule. The Board's decision is binding unless—

(1) The affected party has a right to judicial review and timely files a civil action in a United States District Court or, in the case of a civil money penalty, in a United States Court of Appeals; or

(2) The Board opens and revises its decision in accordance with § 498.102.

(b) Right to judicial review. Section 498.5 specifies the circumstances under which an affected party has a right to seek judicial review.

(c) Special rules: Civil money penalty.

(1) Finality of Board's decision. When HCFA imposes a civil money penalty, notice of the Board's decision (or denial of review) is the final administrative action that initiates the 60-day period for seeking judicial review.

(2) Timing for collection of civil money penalty. For SNFs and NFs, the rules that apply are those set forth in subpart F of part 488 of this chapter.

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§ 498.95 Extension of time for seeking judicial review.

(a) Any affected party that is dissatisfied with a Departmental Appeals Board decision and is entitled to judicial review must commence civil action within 60 days from receipt of the notice of the Board's decision (or denial of review) is the final administrative action that initiates the 60-day period for seeking judicial review.

(b) The request for extension must be filed in writing with the Board before the 60-day period ends.

(c) For good cause shown, the Board may extend the time for commencing civil action.

Subpart F—Reopening of Decisions Made by Administrative Law Judges or the Departmental Appeals Board

§ 498.100 Basis, timing, and authority for reopening an ALJ or Board decision.

(a) Basis and timing for reopening. An ALJ or Departmental Appeals Board decision may be reopened, within 60 days from the date of the notice of decision, upon the motion of the ALJ or the Board or upon the petition of either party to the hearing.

(b) Authority to reopen. (1) A decision of the Departmental Appeals Board may be reopened only by the Departmental Appeals Board.

(2) A decision of an ALJ may be reopened by that ALJ, by another ALJ if that one is not available, or by the Departmental Appeals Board. For purposes of this paragraph, an ALJ is considered to be unavailable if the ALJ has died, terminated employment, or been transferred to another duty station, is on leave of absence, or is unable to conduct a hearing because of illness.

§ 498.102 Revision of reopened decision.

(a) Revision based on new evidence. If a reopened decision is to be revised on the basis of new evidence that was not included in the record of that decision, the ALJ or the Departmental Appeals Board—

(1) Notifies the parties of the proposed revision; and

(2) Unless the parties waive their right to hearing or appearance—

(i) Grants a hearing in the case of an ALJ revision; and

(ii) Grants opportunity to appear in the case of a Board revision.

(b) Basis for revised decision and right to review. (1) If a revised decision is necessary, the ALJ or the Departmental Appeals Board, as appropriate, renders it on the basis of the entire record.

(2) If the decision is revised by an ALJ, the Departmental Appeals Board may review that revised decision at the request of either party or on its own motion.

§ 498.103 Notice and effect of revised decision.

(a) Notice. The notice mailed to the parties states the basis or reason for the revised decision and informs them of their right to Departmental Appeals Board review of an ALJ revised decision, or to judicial review of a Board reviewed decision.

(b) Effect—(1) ALJ revised decision. An ALJ revised decision is binding unless
it is reviewed by the Departmental Appeals Board.

(2) Departmental Appeals Board revised decision. A Board revised decision is binding unless a party files a civil action in a district court of the United States within the time frames specified in §498.95.