In our review of DND indications, we found sufficient evidence to determine that PET scans are no longer experimental. However, the evidence was insufficient to reach a conclusion that FDG PET is reasonable and necessary in all instances. A sufficient inference of benefit, however, can be drawn to support limited coverage if certain safeguards for patients are provided. This inference is based on both the physiological basis for FDG PET usefulness in a differential diagnosis of fronto-temporal dementia (FTD) and Alzheimer’s disease (AD), as well as, evidence of a positive benefit of PET for patients with several other dementing neurodegenerative diseases for which there is evidence of sufficient quality to warrant coverage.

The purpose of this system is to collect and maintain information on Medicare beneficiaries receiving FDG PET scans for indications for DND when there is not sufficient evidence to reach a firm conclusion that the scan is reasonable and necessary unless they are enrolled in an approved study. Information retrieved from this system will be disclosed to: (1) Support regulatory, reimbursement, and policy functions performed within the agency or by a contractor or consultant; (2) assist another Federal or state agency with information to enable such agency to administer a Federal health benefits program, or to enable such agency to fulfill a requirement of Federal statute or regulation that implements a health benefits program funded in whole or in part with Federal funds; (3) support an agency or program related to the prevention of disease or disability, the restoration or maintenance of health, or payment for medical care; (4) support constituent requests made to a Congressional representative; (5) support litigation involving the agency; and (6) combat fraud and abuse in certain health benefits programs. We have provided background information about the new system in the Supplementary Information section below. Although the Privacy Act requires only that CMS provide an opportunity for interested persons to comment on the proposed routine uses, CMS invites comments on the Secretary (HHS) with respect to whether or not a particular item or service is covered nationally under Title XVIII of the Social Security Act (the Act) section 1869(f)(1)(B). In order to be covered by Medicare, an item or service must fall within one or more benefit categories contained in Part A or Part B, and must not be otherwise excluded from coverage.
all portions of this notice. See EFFECTIVE DATE section for comment period.

EFFECTIVE DATE: CMS has filed a new SOR report with the Chair of the House Committee on Government Reform and Oversight, the Chair of the Senate Committee on Governmental Affairs, and the Administrator, Office of Information and Regulatory Affairs, Office of Management and Budget (OMB) on October 5, 2005. We will not disclose any information under a routine use until 30 days after publication. We may defer implementation of this system or one or more of the routine use statements listed below if we receive comments that persuade us to defer implementation.

ADDRESS: The public should address comments to the CMS Privacy Officer, Mail Stop N2–04–27, 7500 Security Boulevard, Baltimore, Maryland 21244–1850. Comments received will be available for review at this location, by appointment, during regular business hours, Monday through Friday from 9 a.m.–3 p.m., eastern daylight time.

FOR FURTHER INFORMATION CONTACT: Rosemarie Hakim, Epidemiologist, Division of Operations and Committee Management, Coverage and Analysis Division of Operations and Committee Management, Coverage and Analysis Group, Office of Clinical Standards and Quality, CMS, Mail Stop C1–09–06, 7500 Security Boulevard, Baltimore, Maryland 21244–1849. Her telephone number is (410) 786–3934, or she can be reached via e-mail at Rosemarie.Hakim@cms.hhs.gov.

SUPPLEMENTARY INFORMATION: Medicare covers FDG–PET scans for either the differential diagnosis of FTD and AD under specific requirements; or, its use in a CMS approved practical clinical trial focused on the utility of FDG–PET in the diagnosis or treatment of dementia or other neurodegenerative diseases. Specific requirements for each indication are clarified as follows: an FDG–PET scan is considered reasonable and necessary in patients with a recent diagnosis of dementia and documented cognitive decline of at least 6 months, who meet diagnostic criteria for both AD and FTD. These patients have been evaluated for specific alternate neurodegenerative diseases or other causative factors, but the cause of the clinical symptoms remains uncertain.

The following additional conditions must be met before an FDG–PET scan will be covered: (1) The patient’s onset, clinical presentation, or course of cognitive impairment is such that FTD is suspected as an alternative neurodegenerative cause of the cognitive deficit. Specifically, symptoms such as social disinhibition, awkwardness, difficulties with language, or loss of executive function are more prominent early in the course of FTD than the memory loss typical of AD;

(2) The patient has had a comprehensive clinical evaluation (as defined by the American Academy of Neurology) encompassing a medical history from the patient and a well-acquainted informant (including assessment of activities of daily living), physical and mental status examination (including formal documentation of cognitive decline occurring over at least 6 months) aided by cognitive scales or neuropsychological testing, laboratory tests, and structural imaging such as magnetic resonance imaging (MRI) or computed tomography (CT);

(3) The evaluation of the patient has been conducted by a physician experienced in the diagnosis and assessment of dementia;

(4) The evaluation of the patient did not clearly determine a specific neurodegenerative disease or other cause for the clinical symptoms, and information available through FDG–PET is reasonably expected to help clarify the diagnosis between FTD and AD and help guide future treatment;

(5) The FDG–PET scan is performed in a facility that has all the accreditation necessary to operate nuclear medicine equipment. The reading of the scan should be done by an expert in nuclear medicine, radiology, neurology, or psychiatry, with experience interpreting such scans in the presence of dementia and;

(6) A brain single photon emission computed tomography (SPECT) or FDG–PET scan has not been obtained for the same indication. (The indication can be considered to be different in patients who exhibit important changes in scope or severity of cognitive decline, and meet all other qualifying criteria listed above and below (including the judgment that the likely diagnosis remains uncertain.) The results of a prior SPECT or FDG–PET scan must have been inconclusive or, in the case of SPECT, difficult to interpret due to immature or inadequate technology. In these instances, an FDG–PET scan may be covered after one year has passed from the time the first SPECT or FDG–PET scan was performed.)

The referring and billing provider(s) have documented the appropriate evaluation of the Medicare beneficiary. Providers should establish the medical necessity of an FDG–PET scan by ensuring that the following information has been collected and is maintained in the beneficiary medical record: Date of onset of symptoms; diagnosis of clinical syndrome (normal aging; mild cognitive impairment; mild, moderate or severe dementia); mini mental status exam or similar test score; presumptive cause (possible, probable, uncertain AD); any neuropsychological testing performed; results of any structural imaging (MRI or CT) performed; relevant laboratory tests (B12, thyroid hormone); and, number and name of prescribed medications.

The billing provider must furnish a copy of the FDG–PET scan result for use by CMS and its contractors upon request. These verification requirements are consistent with Federal requirements set forth in 42 Code of Federal Regulations (CFR) Section 410.32 generally for diagnostic x-ray tests, diagnostic laboratory tests, and other tests. In summary, section 410.32 requires the billing physician and the referring physician to maintain information in the medical record of each patient to demonstrate medical necessity [410.32(d)(2)] and submit the information demonstrating medical necessity to CMS and/or its agents upon request [410.32(d)(3)(I)] (OMB number 0938–0685).

A FDG–PET scan is considered reasonable and necessary in patients with mild cognitive impairment or only in the context of an approved clinical trial that contains patient safeguards and protections to ensure proper administration, use and evaluation of the FDG–PET scan.

The clinical trial must compare patients who do and do not receive an FDG–PET scan and have as its goal to monitor, evaluate, and improve clinical outcomes. In addition, it must meet the following basic criteria: written protocol on file; Institutional Review Board review and approval; scientific review and approval by two or more qualified individuals who are not part of the research team; and, certification that investigators have not been disqualified. All other uses of FDG–PET for patients with a presumptive diagnosis of dementia-causing neurodegenerative disease (e.g., possible or probable AD, clinically typical FTD, dementia of Lewy bodies, or Creutzfeld-Jacob disease) for which CMS has not specifically indicated coverage continue to be noncovered.

CMS will consider prospective data collection systems to be qualified if they provide assurance that the specific hypotheses are addressed and they collect appropriate data elements. The data collection shall include baseline patient characteristics: Indications for the PET scan; PET scan type and characteristics; FDG PET results; results of all other imaging and provider characteristics; differential diagnosis; and stage; long term patient
outcomes; disease management changes; and treatment received. The clinical data collection must ensure that specific hypotheses are identified prospectively; hospitals and providers are qualified to provide FDG PET and interpret the results; and participating hospitals and providers collect prospective data at the time of payment on all enrolled patients undergoing FDG PETs for DND indications. Data elements will be transmitted to CMS for evaluation of the short and long term benefits of the FDG PET for its beneficiaries and inform future clinical decision making. CMS shall be assured that all applicable patient confidentiality, privacy, and other Federal laws are complied with, including the Standards for Privacy of Individually Identifiable Health Information.

I. Description of the Proposed System of Records

A. Statutory and Regulatory Basis for SOR

The statutory authority for linking coverage decisions to the collection of additional data is derived from Sec. 1862(a)(1)(A) of the Act, which states that Medicare may not provide payment for items and services unless they are “reasonable and necessary” for the treatment of illness or injury. In some cases, CMS will determine that an item or service is only reasonable and necessary when specific data collections accompany the provision of the service. In these cases, the collection of data is required to ensure that the care provided to individual patients will improve health outcomes.

B. Collection and Maintenance of Data in the System

The data collection shall include baseline patient characteristics: Indications for the PET scan; PET scan type and characteristics; FDG PET results; results of all other imaging studies; facility and provider characteristics; differential diagnosis; long term patient outcomes; disease management changes; and DND treatment received. The collected information will contain name, address, telephone number, Health Insurance Claim Number (HICN), geographic location, race/ethnicity, gender, and date of birth, as well as, background information relating to Medicare or Medicaid issues.

II. Agency Policies, Procedures, and Restrictions on the Routine Use

A. Agency Policies, Procedures, and Restrictions on the Routine Use

The Privacy Act permits us to disclose information without an individual’s consent if the information is to be used for a purpose that is compatible with the purpose(s) for which the information was collected. Any such disclosure of data is known as a “routine use.” The government will only release PET DND information that can be associated with an individual as provided for under “Section III. Proposed Routine Use Disclosures of Data in the System.” Both identifiable and non-identifiable data may be disclosed under a routine use.

We will only collect the minimum personal data necessary to achieve the purpose of PET DND. CMS has the following policies and procedures concerning disclosures of information that will be maintained in the system. Disclosure of information from the system will be approved only to the extent necessary to accomplish the purpose of the disclosure and only after CMS:

1. Determines that the use or disclosure is consistent with the reason that the data is being collected, e.g., to collect and maintain information on Medicare beneficiaries receiving PET scans for indications for which there is not sufficient evidence to reach a firm conclusion that the scan is reasonable and necessary unless they are enrolled in an approved study.
2. Determines that:
   a. The purpose for which the disclosure is to be made can only be accomplished if the record is provided in individually identifiable form;
   b. The purpose for which the disclosure is to be made is of sufficient importance to warrant the effect and/or risk on the privacy of the individual that additional exposure of the record might bring; and
   c. There is a strong probability that the proposed use of the data would in fact accomplish the stated purpose(s).
3. Requires the information recipient to:
   a. Establish administrative, technical, and physical safeguards to prevent unauthorized use of disclosure of the record;
   b. Remove or destroy at the earliest time all patient-identifiable information;
   c. Agree to not use or disclose the information for any purpose other than the stated purpose under which the information was disclosed.
4. Determines that the data are valid and reliable.

III. Proposed Routine Use Disclosures of Data in the System

A. The Privacy Act allows us to disclose information without an individual’s consent if the information is to be used for a purpose that is compatible with the purpose(s) for which the information was collected. Any such compatible use of data is known as a “routine use.” The proposed routine use in this system meet the compatibility requirement of the Privacy Act. We are proposing to establish the following routine use disclosures of information maintained in the system:

1. To agency contractors or consultants who have been engaged by the agency to assist in the performance of a service related to this system and who need to have access to the records in order to perform the activity.

2. To another Federal or State agency to:
   a. Contribute to the accuracy of CMS’s proper payment of Medicare benefits,
   b. Enable such agency to administer a Federal health benefits program, or as necessary to enable such agency to fulfill a requirement of a Federal statute or regulation that implements a health benefits program funded in whole or in part with Federal funds, and/or
   c. Assist Federal/state Medicaid programs within the state.

Other Federal or state agencies in their administration of a Federal health program may require PET DND information in order to collect information on Medicare beneficiaries receiving PET scans for sufficient evidence to reach a firm conclusion that the scan is reasonable and necessary.

3. To an individual or organization for a research project or in support of an evaluation project related to the...
prevention of disease or disability, the restoration or maintenance of health, or payment related projects.

The PET DND data will provide for research or in support of evaluation projects, a broader, longitudinal, national perspective of the status of Medicare beneficiaries. CMS anticipates that many researchers will have legitimate requests to use this data in projects that could ultimately improve the care provided to Medicare beneficiaries and the policy that governs the care.

4. To a member of Congress or to a Congressional staff member in response to an inquiry of the Congressional office made at the written request of the constituent about whom the record is maintained.

Beneficiaries sometimes request the help of a member of Congress in resolving an issue relating to a matter before CMS. The member of Congress then writes CMS, and CMS must be able to give sufficient information to be responsive to the inquiry.

5. To the Department of Justice (DOJ), court or adjudicatory body when:
   a. The agency or any component thereof, or
   b. Any employee of the agency in his or her official capacity, or
   c. Any employee of the agency in his or her individual capacity where the DOJ has agreed to represent the employee, or
   d. The United States Government, is a party to litigation or has an interest in such litigation, and by careful review, CMS determines that the records are both relevant and necessary to the litigation and that the use of such records by the DOJ, court or adjudicatory body is compatible with the purpose for which the agency collected the records.

Whenever CMS is involved in litigation, and occasionally when another party is involved in litigation and CMS’ policies or operations could be affected by the outcome of the litigation, CMS would be able to disclose information to the DOJ, court or adjudicatory body involved.

6. To a CMS contractor (including, but not necessarily limited to Medicare administrative contractors, fiscal intermediaries and carriers) that assists in the administration of a CMS-administered health benefits program, or to a grantee of a CMS-administered grant program, when disclosure is deemed reasonably necessary by CMS to prevent, deter, discover, detect, investigate, examine, prosecute, sue with respect to, defend against, correct, remedy, or otherwise combat fraud or abuse in such program.

We contemplate disclosing information under this routine use only in situations in which CMS may enter into a contractual relationship or grant with a third party to assist in accomplishing CMS functions relating to the purpose of combating fraud and abuse.

CMS occasionally contracts out certain of its functions and makes grants when doing so would contribute to effective and efficient operations. CMS must be able to give a contractor or grantee whatever information is necessary for the contractor or grantee to fulfill its duties. In these situations, safeguards are provided in the contract prohibiting the contractor or grantee from using or disclosing the information for any purpose other than that described in the contract and requiring the contractor or grantee to return or destroy all information.

7. To another Federal agency or to an instrumentality of any governmental jurisdiction within or under the control of the United States (including any State or local governmental agency), that administers, or that has the authority to investigate potential fraud or abuse in, a health benefits program funded in whole or in part by Federal funds, when disclosure is deemed reasonably necessary by CMS to prevent, deter, discover, detect, investigate, examine, prosecute, sue with respect to, defend against, correct, remedy, or otherwise combat fraud or abuse in such programs.

Other agencies may require PET DND information for the purpose of combating fraud and abuse in such Federally-funded programs.

B. Additional Provisions Affecting Routine Use Disclosures. This system contains Protected Health Information (PHI) as defined by HHS regulation “Standards for Privacy of Individually Identifiable Health Information” (45 CFR Parts 160 and 164, 65 FR 82462 (12–28–00), Subparts A and E. Disclosures of PHI are authorized by these routine uses may only be made if, and as, permitted or required by the “Standards for Privacy of Individually Identifiable Health Information.”

In addition, our policy will be to prohibit release even if not directly identifiable information, except pursuant to one of the routine uses or if required by law, if we determine there is a possibility that an individual can be identified through implicit deduction based on small cell sizes (instances where the patient population is so small that individuals who are familiar with the enrollees could, because of the small size, use this information to deduce the identity of the beneficiary).

IV. Safeguards

CMS has safeguards in place for authorized users and monitors such users to ensure against excessive or unauthorized use. Personnel having access to the system have been trained in the Privacy Act and information security requirements. Employees who maintain records in this system are instructed not to release data until the intended recipient agrees to implement appropriate management, operational and technical safeguards sufficient to protect the confidentiality, integrity and availability of the information and information systems and to prevent unauthorized access.

This system will conform to all applicable Federal laws and regulations and Federal, HHS, and CMS policies and standards as they relate to information security and data privacy. These laws and regulations include but are not limited to: The Privacy Act of 1974; the Federal Information Security Management Act of 2002; the Computer Fraud and Abuse Act of 1986; the Health Insurance Portability and Accountability Act of 1996; the E-Government Act of 2002, the Clinger-Cohen Act of 1996; the Medicare Modernization Act of 2003, and the corresponding implementing regulations. OMB Circular A–130, Management of Federal Resources, Appendix III, Security of Federal Automated Information Resources also applies. Federal, HHS, and CMS policies and standards include but are not limited to: All pertinent National Institute of Standards and Technology publications; HHS Information Systems Program Handbook and the CMS Information Security Handbook.

V. Effects of the Proposed System of Records on Individual Rights

CMS proposes to establish this system in accordance with the principles and requirements of the Privacy Act and will collect, use, and disseminate information only as prescribed therein. Data in this system will be subject to the authorized releases in accordance with the routine uses identified in this system of records.

CMS will take precautionary measures (see item IV above) to minimize the risks of unauthorized access to the records and the potential harm to individual privacy or other personal or property rights of patients whose data are maintained in the system. CMS will collect only that information necessary to perform the system’s functions. In addition, CMS will make disclosure from the proposed system only with consent of the subject
individual, or his/her legal representative, or in accordance with an applicable exception provision of the Privacy Act. CMS, therefore, does not anticipate an unfavorable effect on individual privacy as a result of information relating to individuals.


Lori Davis,
Acting Chief Operating Officer, Centers for Medicare & Medicaid Services.

SYSTEM NO. 09–70–0561.

SYSTEM NAME
Fluoro-Deoxy Glucose (FDG) Positron Emission Tomography (PET) for Dementia and Neurodegenerative Diseases (DND) (PET DND) HHS/CMS/OCSCQ.

SECURITY CLASSIFICATION:
Level Three Privacy Act Sensitive Data.

SYSTEM LOCATION:
Centers for Medicare & Medicaid Services (CMS) Data Center, 7500 Security Boulevard, North Building, First Floor, Baltimore, Maryland 21244–1850; and at various co-locations of CMS contractors.

CATEGORIES OF INDIVIDUALS COVERED BY THE SYSTEM:
Providers participating in and patients enrolled in one of the following types of prospective clinical studies: a clinical trial of FDG PET that meets the Food and Drug Administration category B investigational device exemption or an FDG PET clinical study that is designed to prospectively collect information at the time of the scan to assist in patient management.

CATEGORIES OF RECORDS IN THE SYSTEM:
The data collection should include baseline patient characteristics: Indications for the PET scan; PET scan type and characteristics; FDG PET results; results of all other imaging studies; facility and provider characteristics; differential diagnosis; long term patient outcomes; disease management changes; and DND treatment received. The collected information will contain name, address, telephone number, Health Insurance Claim Number (HCN) number, geographic location, race/ethnicity, gender, and date of birth, as well as, background information relating to Medicare or Medicaid issues.

AUTHORITY FOR MAINTENANCE OF THE SYSTEM:
The statutory authority for linking coverage decisions to the collection of additional data is derived from Sec. 1862(a)(1)(A) of the Social Security Act, which states that Medicare may not provide payment for items and services unless they are “reasonable and necessary” for the treatment of illness or injury. In some cases, CMS will determine that an item or service is only reasonable and necessary when specific data collections accompany the provisions of the service. In these cases, the collection of data is required to ensure that the care provided to individual patients will improve health outcomes.

PURPOSE(S) OF THE SYSTEM:
The purpose of this system is to collect and maintain information on Medicare beneficiaries receiving FDG PET scans for indications for DND when there is not sufficient evidence to reach a firm conclusion that the scan is reasonable and necessary unless they are enrolled in an approved study. Information retrieved from this system will be disclosed to: (1) Support regulatory, reimbursement, and policy functions performed within the agency or by a contractor or consultant; (2) assist another Federal or state agency with information to enable such agency to administer a Federal health benefits program, or to enable such agency to fulfill a requirement of Federal statute or regulation that implements a health benefits program funded in whole or in part with Federal funds; (3) support an individual or organization for a research project or in support of an evaluation project related to the prevention of disease or disability, the restoration or maintenance of health, or payment related projects; (4) support constituent requests made to a Congressional representative; (5) support litigation involving the agency; and (6) combat fraud and abuse in certain health benefits programs.

OUTLINE OF CONTENTS:
A. The Privacy Act allows us to disclose information without an individual’s consent if the information is to be used for a purpose that is compatible with the purpose(s) for which the information was collected. Any such compatible use of data is known as a “routine use.” The proposed routine uses in this system meet the compatibility requirement of the Privacy Act. We are proposing to establish the following routine use disclosures of information maintained in the system:

1. To agency contractors or consultants who have been engaged by the agency to assist in the performance of a service related to this system and who need to have access to the records in order to perform the activity.

2. To another Federal or state agency to:
   a. Contribute to the accuracy of CMS’s proper payment of Medicare benefits.
   b. Enable such agency to administer a Federal health benefits program, or as necessary to enable such agency to fulfill a requirement of a Federal statute or regulation that implements a health benefits program funded in whole or in part with Federal funds, and/or
   c. Assist Federal/State Medicaid programs within the State.

3. To an individual or organization for a research project or in support of an evaluation project related to the prevention of disease or disability, the restoration or maintenance of health, or payment related projects.

4. To a member of Congress or to a Congressional staff member in response to an inquiry of the Congressional office made at the written request of the constituent about whom the record is maintained.

5. To the Department of Justice (DOJ), court or adjudicatory body when:
   a. The agency or any component thereof, or
   b. Any employee of the agency in his or her official capacity, or
   c. Any employee of the agency in his or her individual capacity where the DOJ has agreed to represent the employee, or
   d. The United States Government, is a party to litigation or has an interest in such litigation, and by careful review, CMS determines that the records are both relevant and necessary to the litigation and that the use of such records by the DOJ, court or adjudicatory body is compatible with the purpose for which the agency collected the records.

6. To a CMS contractor (including, but not necessarily limited to Medicare administrative contractors, fiscal intermediaries and carriers) that assists in the administration of a CMS-administered health benefits program, or to a grantee of a CMS-administered grant program, when disclosure is deemed reasonably necessary by CMS to prevent, deter, discover, detect, investigate, examine, prosecute, sue with respect to, defend against, correct, remedy, or otherwise combat fraud or abuse in such program.

7. To another Federal agency or to an instrumentality of any governmental jurisdiction within or under the control of the United States (including any State or local governmental agency) that administers, or that has the authority to investigate potential fraud or abuse in,
a health benefits program funded in whole or in part by Federal funds, when disclosure is deemed reasonably necessary by CMS to prevent, deter, discover, detect, investigate, examine, prosecute, sue with respect to, defend against, correct, remedy, or otherwise combat fraud or abuse in such programs.

B. Additional Provisions Affecting Routine Use Disclosures. This system contains Protected Health Information (PHI) as defined by Department of Health and Human Services (HHS) regulation “Standards for Privacy of Individually Identifiable Health Information” (45 Code of Federal Regulations (CFR) Parts 160 and 164. 65 Fed. Reg. 82462 (12–28–00), Subparts A and E. Disclosures of PHI authorized by these routine uses may only be made if, and as, permitted or required by the “Standards for Privacy of Individually Identifiable Health Information.”

In addition, our policy will be to prohibit release even if not directly identifiable information, except pursuant to one of the routine uses or if required by law, if we determine there is a possibility that an individual can be identified through implicit deduction based on small cell sizes (instances whether the patient population is so small that individuals who are familiar with the enrollee could, because of the small size, use this information to deduce the identity of the beneficiary).

POLICIES AND PRACTICES FOR STORING, RETRIEVING, ACCESSING, RETAINING, AND DISPOSING OF RECORDS IN THE SYSTEM:

STORAGE:
All records are stored electronically.

RETRIEVABILITY:
The data are retrieved by an individual identifier i.e., name of beneficiary or provider.

SAFEGUARDS:
CMS has safeguards in place for authorized users and monitors such users to ensure against excessive or unauthorized use. Personnel having access to the system have been trained in the Privacy Act and information security requirements. Employees who maintain records in this system are instructed not to release data until the intended recipient agrees to implement appropriate management, operational and technical safeguards sufficient to protect the confidentiality, integrity and availability of the information and information systems and to prevent unauthorized access.

This system will conform to all applicable Federal laws and regulations and Federal, HHS, and CMS policies and standards as they relate to information security and data privacy. These laws and regulations include but are not limited to: The Privacy Act of 1974; the Federal Information Security Management Act of 2002; the Computer Fraud and Abuse Act of 1986; the Health Insurance Portability and Accountability Act of 1996; the E-Government Act of 2002; the Clinger-Cohen Act of 1996; the Medicare Modernization Act of 2003, and the corresponding implementing regulations. OMB Circular A–130, Management of Federal Resources, Appendix III, Security of Federal Automated Information Resources also applies. Federal, HHS, and CMS policies and standards include but are not limited to: All pertinent National Institute of Standards and Technology publications; HHS Information Systems Program Handbook and the CMS Information Security Handbook.

RETENTION AND DISPOSAL:
CMS will retain information for a total period of 10 years. All claims-related records are encompassed by the document preservation order and will be retained until notification from DOJ.

SYSTEM MANAGER AND ADDRESS:
Director, Coverage and Analysis Group, Office of Clinical Standards and Quality, CMS, Mail Stop C1–09–06, 7500 Security Boulevard, Baltimore, Maryland 21244–1850.

NOTIFICATION PROCEDURE:
For the purpose of access, the subject individual should write to the system manager who will require the system name, address, age, gender, and for verification purposes, the subject individual’s name (woman’s maiden name, if applicable).

RECORD ACCESS PROCEDURE:
For the purpose of access, use the same procedures outlines in Notification Procedures above. Requestors should also reasonably specify the record contents being sought. (These procedures are in accordance with Department regulation 45 CFR 5b.5.)

CONTESTING RECORDS PROCEDURES:
The subject individual should contact the system manager named above and reasonably identify the records and specify the information to be contested. State the corrective action sought and the reasons for the correction with supporting justification. (These procedures are in accordance with Department regulation 45 CFR 5b.7.)

RECORD SOURCE CATEGORIES:
Records maintained in this system are derived from Carrier and Fiscal Intermediary Systems of Records, Common Working File System of Records, clinics, institutions, hospitals and group practices performing the procedures, and outside registries and professional interest groups.

SYSTEMS EXEMPTED FROM CERTAIN PROVISIONS OF THE ACT:
None.

BILLING CODE 4120–03–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Administration on Children, Youth and Families; Award Announcement

AGENCY: Administration on Children, Youth and Families, ACF, HHS.

ACTION: Award announcement.

SUMMARY: The Administration on Children, Youth and Families, Family and Youth Services Bureau (FYSB), herein announces the awarding of twenty-eight urgent grant awards in order to enable seventeen Mentoring Children of Prisoner Programs and eleven Training and Technical Assistance providers to respond immediately to hurricane disaster evacuee needs in their States and local communities. The effects of Hurricane Katrina have disrupted the ability of the children whose parents are incarcerated to receive mentoring services due to their forced relocation throughout the nation. As a result, FYSB’s network of mentoring grantees and training and technical assistance providers are uniquely positioned to respond to the increase in the numbers of children of incarcerated parents arriving in their new communities. The following agencies are receiving grant funds for a twelve month project period: Big Brothers Big Sisters of Heart, Macon, Georgia, in the amount of $95,000; State of Alabama Child Abuse and Neglect Prevention Board, Montgomery, Alabama, in the amount of $50,000; YMCA of Greater Louisville, Louisville, Kentucky, in the amount of $50,000; Big Brothers Big Sisters of Mississippi, Jackson, Mississippi, in the amount of $95,000; Family and Children’s Agency, Inc., Norwalk, Connecticut, in the amount of $21,350; America on Track of Santa Ana, California in the amount of $95,000; Volunteers in Prevention, Probation and Prisons, Detroit,