

and Human Services, is publishing the following summary of proposed collections for public comment. Interested persons are invited to send comments regarding this burden estimate or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed information collection for the proper performance of the Agency's function; (2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

1. Type of Information Collection Request: Extension without change of a currently approved collection; **Title of Information Collection:** ESRD Beneficiary Selection and Supporting Regulations Contained in 42 CFR 414.330; **Use:** Section 2145 amended section 1881 of the Social Security Act and changes the way the Medicare program pays for home dialysis services. Medicare patients who currently receive dialysis in a facility but later become home dialysis patients must complete the CMS-382 form at the time they go to the home setting. Facilities are required to have all Medicare home dialysis patients choose one of two payment methods. Under Method I, the dialysis facility assumes responsibility for patient care and the facility provides all dialysis equipment and supplies needed to dialyze at home. The facility is required to order, store, deliver, and pay the manufacturers and suppliers for these items. Under Method II, the beneficiary makes his/her own arrangement for securing the necessary supplies and dialysis equipment. Then, the supplier bills the Medicare program (Carrier) for payment.

Form Number: CMS-382 (OMB#: 0938-0372); **Frequency:** Reporting—Yearly; **Affected Public:** Individuals or households; **Number of Respondents:** 7400; **Total Annual Responses:** 7400; **Total Annual Hours:** 617.

To obtain copies of the supporting statement and any related forms for the proposed paperwork collections referenced above, access CMS Web Site

address at <http://www.cms.hhs.gov/PaperworkReductionActof1995>, or E-mail your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov, or call the Reports Clearance Office on (410) 786-1326. To be assured consideration, comments and recommendations for the proposed information collections must be received by the OMB desk officer at the address below, no later than 5 p.m. on December 17, 2007. OMB Human Resources and Housing Branch, Attention: Carolyn Lovett, New Executive Office Building, Room 10235, Washington, DC 20503, Fax Number: (202) 395-6974.

Dated: November 7, 2007.

Michelle Shortt,

Director, Regulations Development Group,
Office of Strategic Operations and Regulatory Affairs.

[FR Doc. E7-22268 Filed 11-15-07; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Comment Request

Title: Computerized Support Enforcement Systems—NPRM.
OMB No.: 0980-0271.

Description: The information being collected is mandated by Section 454(16) of the Social Security Act (the Act), which provides for the establishment and operation by State agencies, in accordance with an initial and annually updated Advance Planning Document (APD) approved under section 452(d) of the Act, of a statewide system meeting the requirements of section 454A of the Act. In addition, section 454A(e)(1) of the Act requires that States create a State Case Registry (SCR) within their statewide automated child support systems to include information on IV-D cases and non-IV-D orders established or modified in the State on or after October 1, 1998. Section 454A(e)(5) of the Act requires States to regularly update their cases in the SCR.

This notice reflects the new transaction set for SCR to Federal Case Registry (FCR) transactions where States are encouraged, but not required, to submit child data from their SCR to the FCR.

The data being collected for the APD are a combination of narratives, budgets and schedules, which are used to provide funding approvals on an annual basis and to monitor and oversee systems development. Child support has separate regulations under 45 CFR 307.15 related to submittal of APDs because the program had supplemental authority for enhanced funding for systems development, a requirement for Independent Validation and Verification (IV&V) reviews for high risk projects, waiver authority, and has substantial penalties for non-compliance with the statutory deadline of October 1, 2000. This information collection reflects the fact that 52 States and Territories are now certified as meeting the automation requirements of the Family Support Act of 1988 (FSA) and the Personal Responsibility and Work Opportunity Reconciliation Act of 1996 (PRWORA), leaving only two States that are not yet PRWORA systems certified and only one State that has not submitted an Implementation APD for compliance with PRWORA automation. The two States not yet certified have a requirement for ongoing IV&V, i.e., that up to five States will require semiannual IV&V services related to their plans to develop entirely new CSE systems to replace their legacy systems and that one State is operating under a waiver for an Alternative Systems Configuration which requires additional information to be provided on an annual basis. States and Territories that opted to keep their APD for child support systems are covered under a separate Information Collection, OMB No. 0992-0005, for 45 CFR Part 95 Subpart F.

The data being collected for the SCR is used to transmit mandatory data elements to the FCR where it is used for matching against other databases for the purposes of location of individuals, assets, employment and other child support related activities.

Respondents: State and Territorial Child Support Agencies.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
307.15(b)(1)(IV&V), Ongoing	2	12	16	384
307.15(b)(1) (IV&V), Semiannual	5	2	16	160
307.5(b), Waiver Option	1	1	80	80

ANNUAL BURDEN ESTIMATES—Continued

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
307.5(a) System, Certification	2	1	240	480
307.11(e)(1)(ii), Collection of Non-IV-D Data for SCR—States	54	25,200	.046	62,597
Collection of Child Data for IV-D Cases for SCR—States	54	12,000	.083	53,784
307.11(e)(1)(ii), Collection of Non-IV-D Data for SCR—Courts	3,045	447	.029	39,472
307.11(e)(3)(v), Collection of Child Data for IV-D cases for SCR—Courts ...	3,045	213	.083	53,833
307.11(f)(1), Case Data Transmitted from SCR to FCR—New cases and case updates	54	52	2.82	7,919

Estimated Total Annual Burden Hours: 218,709.

Additional Information: Copies of the proposed collection may be obtained by writing to the Administration for Children and Families; Office of Administration, Office of Information Services, 370 L'Enfant Promenade, SW., Washington DC 20447, Attn: ACF Reports Clearance Officer. All requests should be identified by the title of the information collection. E-mail address: infocollection@acf.hhs.gov.

OMB Comment: OMB is required to make a decision concerning the collection of information between 30 and 60 days after publication of this document in the **Federal Register**. Therefore, a comment is best assured of having its full effect if OMB receives it within 30 days of publication. Written comments and recommendations for the proposed information collection should be sent directly to the following: Office of Management and Budget, Paperwork Reduction Project, Fax: 202-395-6974, Attn: Desk Officer for the Administration for Children and Families.

November 7, 2007.

Robert Sargis,

Reports Clearance Officer.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Cancer Institute; Notice of Closed Meetings

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. Appendix 2), notice is hereby given of the following meetings.

The meetings will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title U.S.C., as amended. The grant applications and the discussions could disclose

confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Cancer Institute Initial Review Group; Subcommittee A—Cancer Centers.

Date: December 6–7, 2007.

Time: 8 a.m. to 11 a.m.

Agenda: To review and evaluate grant applications.

Place: Doubletree Hotel Bethesda, 8120 Wisconsin Avenue, Bethesda, MD 20814.

Contact Person: Gail J. Bryant, MD, Scientific Review Administrator, Grants Review Branch, Division of Extramural Activities, National Cancer Institute, National Institutes of Health, 6116 Executive Boulevard, Room 8042, Bethesda, MD 20852–7405, (301) 402–0801.

Name of Committee: National Cancer Institute Special Panel; Cancer Center Support Grant.

Dated: December 6, 2007

Time: 2:40 p.m. to 5 p.m.

Agenda: To review and evaluate grant applications.

Place: Double Tree Bethesda, 8120 Wisconsin Avenue, Bethesda, MD 20814.

Contact Person: David E. Maslow, PhD, Chief, Resources and Training Review Branch, Division of Extramural Activities, National Cancer Institute, National Institutes of Health, 6116 Executive Boulevard—Room 8117, Bethesda, MD 20892–7405, (301) 496–2330.

Name of Committee: National Cancer Institute Special Emphasis Panel; Prevention, Control and Population Sciences.

Date: January 30–31, 2008.

Time: 8 a.m. to 5 p.m.

Agenda: To review and evaluate grant applications.

Place: Courtyard Gaithersburg Washingtonian Center, 204 Boardwalk Place, Gaithersburg, MD 20878.

Contact Person: Wlodek Lopaczynski, MD, PhD, Scientific Review Administrator, Research Programs Review Branch, Division of Extramural Activities, National Cancer Institute, 6116 Executive Blvd. Room 8131, Bethesda, MD 20892, 301–594–1402, lopacw@mail.nih.gov.

Name of Committee: National Cancer Institute Special Emphasis Panel; Molecular Oncology.

Date: February 6–8, 2008.

Time: 5:30 p.m. to 3 p.m.

Agenda: To review and evaluate grant applications.

Place: Hyatt Regency Bethesda, One Bethesda Metro Center, 7400 Wisconsin Avenue, Bethesda, MD 20814.

Contact Person: Michael B. Small, PhD, Scientific Review Administrator, Division of Extramural Activities, National Cancer Institute, National Institutes of Health, 6116 Executive Blvd., Room 8127, Bethesda, MD 20892–8328, 301–402–0996, smallm@mail.nih.gov.

Name of Committee: National Cancer Institute Special Emphasis Panel; SPORE in Lymphoma, Myeloma, Pancreas, GI, H&N and Lung Cancers.

Date: February 12–14, 2008.

Time: February 12, 2008, 8 a.m. to 1 p.m.

Agenda: To review and evaluate grant applications.

Place: Marriott Bethesda, 5151 Pooks Hill Rd., Bethesda, MD 20814.

Contact Person: Shamala K. Srinivas, PhD, Scientific Review Administrator, Research Programs Review Branch, Division of Extramural Activities, National Cancer Institute, National Institutes of Health, 6116 Executive Boulevard, Room 8123, Bethesda, MD 20892, 301–594–1224, ss537t@nih.gov.

Name of Committee: National Cancer Institute Special Emphasis Panel; SPORE in Breast, Gynecologic, Genitourinary and Prostate Cancers.

Date: February 12–14, 2008.

Time: 8 a.m. to 5 p.m.

Agenda: To review and evaluate grant applications.

Place: Bethesda Marriott, 5151 Pooks Hill Road, Bethesda, MD 20814.

Contact Person: Caron Lyman, PhD, Scientific Review Administrator, Division of Extramural Activities, National Cancer Institute, National Institutes of Health, 6116 Executive Blvd., Room 8119, Bethesda, MD 20892–8328, 301–451–4761, lymanc@mail.nih.gov.

Name of Committee: National Cancer Institute Special Emphasis Panel; Application of Emerging Technologies for Cancer Research.

Date: March 19–20, 2008.

Time: 8 a.m. to 5 p.m.

Agenda: To review and evaluate grant applications.

Place: Bethesda Marriott Suites, 6711 Democracy Boulevard, Bethesda, MD 20817.

Contact Person: Marvin L. Salin, PhD, Scientific Review Administrator, Special