

ESTIMATE OF ANNUALIZED BURDEN HOURS

Type of respondent	Form name	No. of respondents	No. of responses per respondent	Avg. burden per response (in hours)
All Respondents	Telephone script for permission to seek consent (C1a or C1b).	294	1	3/60
Screened Eligible Respondents—	Tracking Form (C1c)	250	1	5/60
Pregnancy Exposure (group 1)	Consent (C2a or C2b)	250	1	20/60
Lactation Exposure (group 2)
Pregnancy and Lactation Exposure (group 3).
Groups 1, 2 and 3	Enrollment (D1)	250	1	10/60
Group 1 and 3	Initial pregnancy Questionnaire (D2)	200	1	30/60
	Follow-up pregnancy questionnaire (D3)	200	1	20/60
	Initial infant questionnaire (D4)	200	1	20/60
	Follow-up infant questionnaire (D5)	200	1	15/60
Groups 2 and 3	Initial breastfeeding questionnaire (D6) ...	100	1	20/60
	Follow-up breastfeeding questionnaire (D7).	100	1.5	15/60

Dated: November 6, 2008.

Maryam I. Daneshvar,

Acting Reports Clearance Officer, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Board of Scientific Counselors, National Institute for Occupational Safety and Health (BSC, NIOSH)

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92-463), the Centers for Disease Control and Prevention (CDC) announces the following meeting for the aforementioned committee:

Time and Date: 8:30 a.m.–3:30 p.m., December 4, 2008.

Place: Marriott Crystal City at Reagan National, 1999 Jefferson Davis Highway, Arlington, VA 22202.

Status: Open to the public, limited only by the space available. The meeting room accommodates approximately 50 people. Teleconference available toll-free; please dial (866) 700-6634, Participant Pass Code 3756066.

Purpose: The Secretary, the Assistant Secretary for Health, and by delegation the Director, Centers for Disease Control and Prevention, are authorized under Sections 301 and 308 of the Public Health Service Act to conduct directly or by grants or contracts, research, experiments, and demonstrations relating to occupational safety and health and to mine health. The Board of Scientific Counselors shall provide guidance to the Director, National Institute for Occupational Safety and Health on research and prevention programs. Specifically, the Board shall provide guidance on the Institute's research activities related to developing and

evaluating hypotheses, systematically documenting findings and disseminating results. The Board shall evaluate the degree to which the activities of the National Institute for Occupational Safety and Health: (1) Conform to appropriate scientific standards, (2) address current, relevant needs, and (3) produce intended results.

Matters To Be Discussed: Agenda items include a Report by the Acting Director of NIOSH; National Academies (NA) Recommendations for NIOSH Programs; Implementation of NA Recommendations in Agriculture, Forestry and Fishing; Occupational Safety and Health Surveillance Needs; NIOSH Nanotechnology Research Strategic Plan; National Occupational Research Agenda; Future Meetings and Closing Remarks.

Agenda items are subject to change as priorities dictate.

Contact Person for More Information: Roger Rosa, Executive Secretary, BSC, NIOSH, CDC, 395 E Street, SW., Suite 9200, Patriots Plaza Building, Washington, DC 20201, telephone (202) 245-0655, fax (202) 245-0664.

The Director, Management Analysis and Services Office, has been delegated the authority to sign **Federal Register** notices pertaining to announcements of meetings and other committee management activities for both the CDC and the Agency for Toxic Substances and Disease Registry.

Dated: November 6, 2008.

Elaine L. Baker,

Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS-R-74, CMS-R-107, CMS-2786U, CMS-R-285 and CMS-R-245]

Agency Information Collection Activities: Proposed Collection; Comment Request

AGENCY: Centers for Medicare & Medicaid Services, HHS.

In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Centers for Medicare & Medicaid Services (CMS) 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 functions; (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 of a currently approved collection; *Title of Information Collection:* Income and Eligibility Verification System; *Use:* This collection is necessary to verify income and eligibility requirements for Medicaid beneficiaries, as required by Section 1137 of the Social Security Act. *Form Number:* CMS-R-74 (OMB#

0938-0467); *Frequency*: Monthly; *Affected Public*: State, Local or Tribal Governments; *Number of Respondents*: 54; *Total Annual Responses*: 54; *Total Annual Hours*: 124,054.

2. Type of Information Collection

Request: Extension of a currently approved collection; *Title of Information Collection*: Medicaid-Determining Third Party Liability (TPL) State Plan Preprint and Supporting Regulations in 42 CFR 433.138; *Use*: The information collected from Medicaid applicants and beneficiaries as well as from State and local agencies is necessary to determine the legal liability of third parties to pay for medical services in lieu of Medicaid payment. *Form Number*: CMS-R-107 (OMB# 0938-0502); *Frequency*: On occasion; *Affected Public*: Individuals or households and State, Local or Tribal Government; *Number of Respondents*: 2,900,000; *Total Annual Responses*: 2,900,000; *Total Annual Hours*: 510,968.

3. Type of Information Collection

Request: Revision of a currently approved collection; *Title of Information Collection*: Fire Safety Survey Reports; *Use*: The Life Safety Code (LSC) is a compilation of fire safety requirements for new and existing buildings and is updated and published every 3 years by the National Fire Protection Association (NFPA), a private, non-profit organization dedicated to reducing loss of life due to fire. The Medicare regulations have historically incorporated by reference these requirements along with Secretarial waiver authority. The statutory basis for incorporating NFPA's LSC for our providers is under the Secretary's general rulemaking authority at Sections 1102 and 1871 of the Social Security Act. These forms are used by the State Agencies to record data collected to determine compliance with standards specified in 416.44(b) for ambulatory surgical centers (ASCs), and 494.60(e) for End-Stage Renal Disease (ESRD) facilities. The Medicare Health Insurance Program is authorized by Title XVIII of the Social Security Act. The CMS-2786U form is being revised to include ESRD information. *Form Number*: 2786U (OMB# 0938-0242); *Frequency*: Weekly; *Affected Public*: Individuals or households and State, Local or Tribal Government; *Number of Respondents*: 54; *Total Annual Responses*: 2442; *Total Annual Hours*: 4884.

4. Type of Information Collection

Request: Extension of a currently approved collection; *Title of Information Collection*: Request For Retirement Benefit Information; *Use*: Section 1818 of the Social Security Act

provides that former State and local government employees who are age 65 or older, that have been entitled to Premium Part A for at least 7 years, and did not have the premium paid for by a State or a political subdivision of a State, may have the Part A premium reduced to zero. This collection will assist in determining whether individuals currently paying a monthly premium for Medicare Part A coverage are eligible to have their premium reduced to zero. *Form Number*: CMS-R-285 (OMB# 0938-0769); *Frequency*: Monthly; *Affected Public*: State, Local or Tribal Governments; *Number of Respondents*: 1,500; *Total Annual Responses*: 1,500; *Total Annual Hours*: 375.

5. Type of Information Collection

Request: Revision of a currently approved collection; *Title of Information Collection*: Medicare and Medicaid Programs OASIS Collection Requirements as Part of the CoPs for HHAs and Supporting Regulations in 42 CFR, Sections 484.55, 484.205, 484.245, 484.250; *Use*: The Centers for Medicare and Medicaid Services is requesting OMB approval to modify the Outcome and Assessment Information Set (OASIS) data set that home health agencies (HHAs) are required to collect in order to participate in the Medicare program. Proposed revisions to the OASIS data set include: (1) Issues raised by stakeholders, including removing items that are not currently used by CMS for payment or quality, adding items to address clinical domains not currently covered, and modifying item wording or response categories for selected items; and (2) the addition of process items that support measurement of evidence-based practices. Proposed revisions to OASIS items address issues raised by stakeholders, including removing items that are not currently used by CMS for payment or quality, adding items to address clinical domains not currently covered, and modifying item wording or response categories for selected items. These changes and item deletions are and considered to be high priority by CMS and have implications for outcome measurement, risk adjustment of outcome reports, case mix adjustment for prospective payment, data submission procedures and specifications, reporting systems, and provider paperwork burden.

In addition, adopting measures of efficient and high-quality care is central to the direction that CMS would like to take in its Quality Initiative. In concordance with long-standing federal objectives, CMS ultimately plans to create a standard patient assessment

instrument that can be used across all post-acute care settings. The revision of the OASIS instrument is an opportunity to consider various components of quality care and how patients might be better served as they (and information about them and their care) move among health care settings. For this reason, the OASIS C includes process items that support measurement of evidence-based practices across the post-acute care spectrum that have been shown to prevent exacerbation of serious conditions, can improve care received by individual patients, and can provide guidance to agencies on how to improve care and avoid adverse events. *Form Number*: CMS-R-245 (OMB# 0938-0760); *Frequency*: Occasionally; *Affected Public*: Business or other for-profit and not-for-profit institutions; *Number of Respondents*: 10,170; *Total Annual Responses*: 14,960,070; *Total Annual Hours*: 15,590,610.

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

In commenting on the proposed information collections please reference the document identifier or OMB control number. To be assured consideration, comments and recommendations must be submitted in one of the following ways by *January 13, 2009*:

1. *Electronically*. You may submit your comments electronically to <http://www.regulations.gov>. Follow the instructions for "Comment or Submission" or "More Search Options" to find the information collection document(s) accepting comments.

2. *By regular mail*. You may mail written comments to the following address: CMS, Office of Strategic Operations and Regulatory Affairs, Division of Regulations Development, Attention: Document Identifier/OMB Control Number _____, Room C4-26-05, 7500 Security Boulevard, Baltimore, Maryland 21244-1850.

Dated: November 6, 2008.

Michelle Shortt,

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

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