

Dated: April 6, 2009.

Maryam I. Daneshvar,

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

[FR Doc. E9-8337 Filed 4-10-09; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration on Aging

Agency Information Collection Activities; Proposed Collection; Comment Request; SMP (Formerly Senior Medicare Patrol) Program Outcome Measurement

AGENCY: Administration on Aging, HHS.

ACTION: Notice.

SUMMARY: The Administration on Aging (AoA) is announcing an opportunity for public comment on the proposed collection of certain information by the agency. Under the Paperwork Reduction Act of 1995 (the PRA), Federal agencies are required to publish notice in the **Federal Register** concerning each proposed collection of information, including each proposed extension of an existing collection of information, and to allow 60 days for public comment in response to the notice. This notice solicits comments on the information collection requirements relating to SMP (formerly Senior Medicare Patrol) Program outcome measurement.

DATES: Submit written or electronic comments on the collection of information by June 12, 2009.

ADDRESSES: Submit electronic comments on the collection of information to:

doris.summey@aoa.hhs.gov.

Submit written comments on the collection of information to Administration on Aging, Washington, DC 20201. Attention: Doris Summey.

FOR FURTHER INFORMATION CONTACT:

Doris Summey, by telephone: (202) 357-3533 or by e-mail:

doris.summey@aoa.hhs.gov.

SUPPLEMENTARY INFORMATION: Under the PRA (44 U.S.C. 3501-3520), Federal agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. "Collection of information" is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes agency request or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA (44 U.S.C. 3506(c)(2)(A)) requires Federal agencies to provide a 60-day notice in

the **Federal Register** concerning each proposed collection of information, including each proposed extension of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, AoA is publishing notice of the proposed collection of information set forth in this document. With respect to the following collection of information, AoA invites comments on: (1) Whether the proposed collection of information is necessary for the proper performance of AoA's functions, including whether the information will have practical utility; (2) the accuracy of AoA's estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques when appropriate, and other forms of information technology.

Grantees are required by Congress to provide information for use in program monitoring and for Government Performance and Results Act (GPRA) purposes. This information collection reports the number of active volunteers, issues and inquiries received, other SMP program outreach activities, and the number of Medicare dollars recovered, among other SMP Performance outcomes.

AoA estimates the burden of this collection of information as follows: *Respondents:* 54 SMP grantees at 23 hours per month (276 hours per year, per grantee). *Total Estimated Burden Hours:* 14,904 hours per year.

Dated: April 8, 2009.

Edwin L. Walker,

Acting Assistant Secretary for Aging.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-09-09BK]

Proposed Data Collections Submitted for Public Comment and Recommendations

In compliance with the requirement of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 for opportunity for public comment on proposed data collection projects, the

Centers for Disease Control and Prevention (CDC) will publish periodic summaries of proposed projects. To request more information on the proposed projects or to obtain a copy of the data collection plans and instruments, call 404-639-5960 and send comments to Maryam I. Daneshvar, CDC Acting Reports Clearance Officer, 1600 Clifton Road, MS-D74, Atlanta, GA 30333 or send an e-mail to *omb@cdc.gov*.

Comments are invited on: (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency's estimate of the burden of the proposed collection of information; (c) ways to enhance the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology. Written comments should be received within 60 days of this notice.

Proposed Project

Registration of Individuals Displaced by the Hurricanes Katrina and Rita (Pilot Project)—New—Agency for Toxic Substances and Disease Registry (ATSDR), Coordinating Center for Environmental Health and Injury Prevention (CCEHIP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

On August 29, 2005, Hurricane Katrina made landfall on the coast of the Gulf of Mexico near New Orleans, Louisiana, and became one of the most deadly and destructive storms in U.S. history. Also occurring in 2005, Hurricane Rita was the fourth-most intense Atlantic hurricane ever recorded and the most intense tropical cyclone ever observed in the Gulf of Mexico. Following the initial phase of the response, the Federal Emergency Management Agency (FEMA) assumed the primary role for housing displaced persons over the intermediate term. To support those needing temporary housing, FEMA provided over 143,000 travel trailers, park homes, and mobile homes for persons displaced by the above mentioned storms. However, some persons living in trailers complained of an odor or of eye or respiratory tract irritation.

FEMA entered into an Interagency Agreement with the Centers for Disease Control and Prevention (CDC)/ATSDR on August 16, 2007 to conduct a

comprehensive public health assessment, based on objective and credible research, of air quality conditions present in FEMA housing units to guide FEMA policy makers and inform the public as to the actual conditions in the field and any actions required to better promote a safe and healthful environment for the disaster victims FEMA housed in the units. FEMA's agreement with the CDC includes an initial formaldehyde exposure assessment as well as a subsequent long-term study of the health effects among resident children. Formaldehyde testing conducted and evaluated by the CDC pursuant to the initial exposure assessment has identified the need to evaluate the feasibility of establishing a national registry to identify and monitor the

health of disaster victims who occupied FEMA-provided temporary housing units. The establishment of such a registry would complement the long-term health effects study set forth in the FEMA-CDC Interagency Agreement.

The purpose of this study is to assess the feasibility of contacting and enrolling members of the targeted group in a registry; to provide a basis for budgeting and further planning for a comprehensive registry; and to test the acceptance of and response to a questionnaire composed of standardized health questions related to systemic and respiratory symptoms.

A pre-registration dataset will be created before enrollment. This dataset will be populated with contact information of the study population, gathered from two main sources: FEMA datasets (in the case of occupants of

temporary housing units) and data provided by self-identified individuals who were displaced by the hurricanes but did not live in the FEMA temporary trailers, members of a pre-defined population (in the case of occupants of non-temporary housing units).

A computer-assisted telephone interview (CATI) system based on a paper questionnaire will be used during all interviews to collect data for this project. The first part will consist of screening questions to determine eligibility for enrollment. The second part will contain contact information of the registrant and other household members, demographics, and health status questions, focusing on respiratory outcomes and cancer.

There are no costs to the respondents other than their time.

ESTIMATED ANNUALIZED BURDEN HOURS

Respondents	Form	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total burden (in hours)
Temporary housing unit occupant	Screening questionnaire	8,000	1	3/60	400
	Main questionnaire	4,000	1	45/60	3,000
Non-Temporary housing unit occupant.	Screening questionnaire	2,000	1	3/60	100
	Main questionnaire	1,000	1	45/60	750
Total	4,250

Dated: April 7, 2009.

Maryam I. Daneshvar,

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

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BILLING CODE

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Proposed Information Collection Activity; Comment Request

Title: National Survey of Child and Adolescent Well-Being Second Cohort (NSCAW II).

OMB No.: 0970-0202.

Description: The Department of Health and Human Services (HHS) intends to collect follow-up data on a sample of children and families for the National Survey of Child and Adolescent WellBeing (NSCAW). The NSCAW was authorized under Section 427 of the Personal Responsibility and Work Opportunities Reconciliation Act of 1996. The NSCAW is the only source of nationally representative, firsthand information about the functioning and well-being, service needs, and service utilization of children and families who come to the attention of the child welfare system. Information is collected about children's cognitive, social, emotional, behavioral, and adaptive functioning, as well as family and community factors that are likely to influence their functioning. Family

service needs and service utilization also are addressed in the data collection.

Selection of the current NSCAW sample and baseline data collection began in 2007 with a final anticipated sample size of 5,700 children. The proposed data collection will allow for follow-up of this sample 18 months post-baseline, and will follow the same format as that used in the baseline round and will employ, with only modest revisions, the same instruments that were used in the previous round. Data from NSCAW are made available to the research community through licensing arrangements from the National Data Archive on Child Abuse and Neglect at Cornell University.

Respondents: Children and their associated permanent or foster caregivers, caseworkers, and teachers.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
Child Interview	1,520	1	1.33	2,022
Caregiver Interview	1,520	1	1.6	2,432
Caseworker Interview	355	1	1	355