

authorization and appropriation of availability of funds, GSA control of the site to complete archaeological investigations and continuity of the tenant agencies' Program of Requirements as they were understood at the time this study was completed.

Following this thirty (30) day notice in the Federal Register, the GSA will issue a ROD at which time its availability will be announced in the **Federal Register** and local media.

II. Distribution

Copies of the FEIS are being distributed to select stakeholders as well as being made available for public review at the International Falls Public Library, Chamber of Commerce Offices, and Koochiching County Office Building.

Dated: October 6, 2011.

Ann P. Kalayil,

Regional Commissioner, Public Buildings Service, Great Lakes Region.

[FR Doc. 2011-26647 Filed 10-13-11; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration on Aging

Agency Information Collection Activities; Proposed Collection; Comment Request; the Evaluation of the Aging and Disability Resource Center Program

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 the Evaluation of the Aging and Disability Resource Center Program.

DATES: Submit written or electronic comments on the collection of information by December 13, 2011.

ADDRESSES: Submit electronic comments on the collection of information to: Susan Jenkins at Susan.Jenkins@aoa.hhs.gov.

Submit written comments on the collection of information to

Administration on Aging, Washington, DC 20201, Attn. Susan Jenkins.

FOR FURTHER INFORMATION CONTACT: Susan Jenkins at 202.357.3591.

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. The Aging and Disability Resource Center (ADRC) Program is a collaborative effort of the Administration on Aging (AoA) and the Centers for Medicare & Medicaid Services (CMS). ADRCs target services to the elderly and individuals with physical disabilities, serious mental illness, and/or developmental/intellectual disabilities. The ultimate goal of the ADRCs is to serve all individuals with long-term care needs regardless of their age or disability. The statutory authority for the ADRC grant program is contained in Titles II and IV of the Older Americans Act (OAA) (42 U.S.C. 3032), as amended by the Older Americans Act Amendments of 2006, Public Law 109-365. (Catalog of Federal Domestic Assistance 93.048, Title IV Discretionary Projects). 42 U.S.C. 3017 specifies that the Assistant Secretary for

Aging "shall measure and evaluate the impact of all programs authorized by this chapter * * * Evaluations shall be conducted by persons not immediately involved in the administration of the program or project evaluated." This new collection of information is necessary to determine the overall effect of ADRCs on both long term support and service systems and individuals. AoA will gather information about how ADRCs provide services and whether consumers, who access ADRCs, as compared to consumers who access other systems, report that the experience is more personalized, consumer-friendly, streamlined, and efficient. Staff of the Administration on Aging's Office of Program Innovation and Demonstration will use the information to both determine the value of the ADRC model and to improve program operations. The evaluation will include both process and outcome components. The *Agency Data Collection Tool* requests respondents' names and contact information to allow the research team to contact potential respondents. The *Personal Experience Survey* will collect information about consumers' level and type of disability, and demographic characteristics including race and living status. Respondents will be asked to provide their Medicare and/or Medicaid identification numbers to allow for analysis of the effect of the ADRC program on health care utilization and nursing home diversion. The proposed data collection tools may be found on the AoA Web site: [INSERT WEB ADDRESS WHEN DETERMINED]. AoA estimates the burden of this collection at 1,732 hours for individuals and 1,294 hours for organizations—Total Burden for Study 3,026.

Dated: October 7, 2011.

Kathy Greenlee,

Assistant Secretary for Aging.

[FR Doc. 2011-26552 Filed 10-13-11; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30 Day-12-0773]

Agency Forms Undergoing Paperwork Reduction Act Review

The Centers for Disease Control and Prevention (CDC) publishes a list of information collection requests under review by the Office of Management and Budget (OMB) in compliance with the

Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these requests, call the CDC Reports Clearance Officer at (404) 639-5960 or send an e-mail to omb@cdc.gov. Send written comments to CDC Desk Officer, Office of Management and Budget, Washington, DC or by fax to (202) 395-5806. Written comments should be received within 30 days of this notice.

Proposed Project

National Surveillance for Severe Adverse Events Associated with Treatment of Latent Tuberculosis Infection (NSSAE)—Reinstatement with change—Division of Tuberculosis Elimination (DTBE), National Center for HIV/AIDS, Viral Hepatitis, STD, and TB Prevention (NCHHSTP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Between October 2000 and September 2004, the CDC received reports of 50 patients with severe adverse events (SAEs) associated with the use of the two or three-month regimen of rifampin and pyrazinamide (RZ) for the treatment of LTBI; 12 (24%) patients died (Morbidity and Mortality Weekly Report 2003;52[31]:735-9). In 2004, CDC began collecting reports of SAEs associated with any treatment regimen for LTBI. For surveillance purposes, an SAE was

defined as any drug-associated reaction resulting in a patient's hospitalization or death after at least one treatment dose for LTBI. During 2004-2008, CDC received 17 reports of SAEs in 15 adults and two children; all patients had received isoniazid (INH) and had experienced severe liver injury (Morbidity and Mortality Weekly Report 2010; 59:224-9).

The CDC requests approval for a 3-year reinstatement with change of the previously approved National Surveillance for Severe Adverse Events Associated with Treatment of Latent Tuberculosis Infection (OMB No. 0920-0773, expired April 31, 2011). The changes include a shortened data collection form and an increase in the number of respondents. This project will continue the passive reporting system for SAEs associated with therapy for LTBI. The system will rely on medical chart review and/or onsite investigations by TB control staff.

The purpose of this information collection request is to determine the annual number and trends of SAEs associated with treatment of LTBI and identify common characteristics of patients with SAEs during treatment of LTBI. Potential correspondents are any of the 60 reporting areas for the national TB surveillance system (the 50 states, the District of Columbia, New York City,

Puerto Rico, and 7 jurisdictions in the Pacific and Caribbean). Data will be collected using the data collection form for adverse event associated with LTBI treatment (NSSAE). The NSSAE form is completed for each reported hospitalization or death related to treatment of LTBI and contains demographic, clinical, and laboratory information. CDC will analyze and periodically publish reports summarizing national LTBI treatment adverse events statistics and also will conduct special analyses for publication in peer-reviewed scientific journals to further describe and interpret these data.

The Food and Drug Administration (FDA) collects data on adverse events related to drugs through the FDA MedWatch Program. CDC is collaborating with FDA in the reporting of SAEs. Reporting will be conducted through telephone, e-mail, or during CDC site visits. In this request, CDC is requesting approval for approximately 60 burden hours annually, an estimated increase of 36 hours from the previously approved 24 hours. This is due to an estimated increase of reports of SAEs after the publication of the MMWR report on SAEs in 2010. There are no costs to respondents other than their time to gather medical records to complete the reporting form.

ESTIMATE OF ANNUALIZED BURDEN HOURS

Type of respondents	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
Physicians	10	1	1
Nurses	10	1	4
Medical Clerk	10	1	1

Dated: October 7, 2011.
Daniel Holcomb,
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

[30Day-12-11JJ]

Agency Forms Undergoing Paperwork Reduction Act Review

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Proposed Project

Evaluating Locally-Developed HIV Prevention Interventions for African-American MSM in Los Angeles—New—National Center for HIV/AIDS, Viral Hepatitis, STD, TB Prevention (NCHHSTP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Data on HIV cases reported in 33 U.S. states with HIV reporting indicate the burden of HIV/AIDS is most concentrated in the African American population compared to other racial/ethnic groups. Of the 49,704 African American males diagnosed with HIV between 2001 and 2004, 54% of these cases were among men who have sex with men (MSM). In Los Angeles County (LAC), the proportion of HIV/AIDS cases among African American males attributable to male-to-male sexual transmission is even greater (75%). In the absence of an effective vaccine, behavioral interventions represent one of the few methods for reducing high HIV incidence among African American MSM (AAMSM). Unfortunately, in the third decade of the