DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Board of Scientific Counselors, National Center for Environmental Health/Agency for Toxic Substances and Disease Registry (BSC, NCEH/ ATSDR), Lead Poisoning Prevention (LPP) Subcommittee; Notice of Meeting

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), the CDC, National Center for Environmental Health/ Agency for Toxic Substances and Disease Registry (NCEH/ATSDR) announces the following meeting of the aforementioned subcommittee:

Time and Date: 12:00 p.m.–1:45 p.m., EST, February 9, 2016.

Place: This meeting will be held by teleconference. To participate in the teleconference, please dial 1–877–315–6535 Passcode: 383520.

Status: The meeting is open to the public, limited only by the conference lines available. The public is welcome to participate during the public comment period, which is tentatively scheduled from 1:30 p.m. to 1:45 p.m.

This Federal Register Notice is being published less than 15 days before the meeting because of the urgent nature of recent events involving the Flint, Michigan water contamination with lead. CDC is convening a meeting of the Lead Subcommittee of the Board of Scientific Counselors to initiate discussion of public health measures and assessments needed in response to this event.

Purpose: The subcommittee will propose strategies and options to the Board of Scientific Counselors (BSC) on ways to prioritize NCEH/ATSDR's activities, improve health outcomes, and address health disparities as it relates to lead exposures. The subcommittee will deliberate on ways to evaluate lead exposure and how to best conduct health evaluations through exposure and epidemiologic studies. Subcommittee proposals on lead prevention practices and national lead poisoning prevention efforts will be provided to the Board of Scientific Counselors for deliberation and possible adoption as formal recommendations to NCEH/ATSDR.

Matters for Discussion: Agenda items will include the following: Blood lead testing and health surveillance strategies for the residents of Flint, Michigan, including methodological approaches for conducting retrospective and

prospective assessments of blood lead levels and associated health outcomes.

Agenda items are subject to change as priorities dictate.

Contact Person for More Information: Sandra Malcom, Committee Management Specialist, NCEH/ATSDR, 4770 Buford Highway, Mail Stop F–45, Chamblee, Georgia 30345; telephone 770/488–0575, Fax: 770/488–3377; Email: smalcom@cdc.gov.

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 Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.

Gary Johnson,

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

[FR Doc. 2016–02574 Filed 2–4–16; 4:15 pm]
BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day-16-0600]

Agency Forms Undergoing Paperwork Reduction Act Review

The Centers for Disease Control and Prevention (CDC) has submitted the following information collection request to the Office of Management and Budget (OMB) for review and approval in accordance with the Paperwork Reduction Act of 1995. The notice for the proposed information collection is published to obtain comments from the public and affected agencies.

Written comments and suggestions from the public and affected agencies concerning the proposed collection of information are encouraged. Your comments should address any of the following: (a) Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility; (b) Evaluate the accuracy of the agencies estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (c) Enhance the quality, utility, and clarity of the information to be collected; (d) Minimize the burden of the collection of information on those who are to respond, including through

the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology, e.g., permitting electronic submission of responses; and (e) Assess information collection costs.

To request additional information on the proposed project or to obtain a copy of the information collection plan and instruments, call (404) 639–7570 or send an email to <code>omb@cdc.gov</code>. Written comments and/or suggestions regarding the items contained in this notice should be directed to the Attention: CDC Desk Officer, Office of Management and Budget, Washington, DC 20503 or by fax to (202) 395–5806. Written comments should be received within 30 days of this notice.

Proposed Project

CDC Model Performance Evaluation Program (MPEP) for Mycobacterium tuberculosis Drug Susceptibility Testing (OMB Control No. 0920–0600, Expires 5/31/2016)—Extension—National Center for HIV/AIDS, Viral Hepatitis, STD, and TB Prevention (NCHHSTP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

As part of the continuing effort to support domestic public health objectives for treatment of tuberculosis (TB), prevention of multi-drug resistance, and surveillance programs, CDC is requesting approval for an extension of three years from the Office of Management and Budget to continue information collection from participants in the Model Performance Evaluation Program for Mycobacterium Tuberculosis Susceptibility Testing. Extension of this information collection will not require changes in the scope of the study, methodology, information collection instruments, or burden on the respondents.

While the overall number of cases of TB in the U.S. has decreased, rates still remain high among foreign-born persons, prisoners, homeless populations, and individuals infected with HIV in major metropolitan areas. To reach the goal of eliminating TB, the Model Performance Evaluation Program for Mycobacterium Tuberculosis Drug Susceptibility Testing is used to monitor and evaluate performance and practices among national laboratories performing M. tuberculosis susceptibility testing. Participation in this program is one way laboratories can ensure high-quality laboratory testing, resulting in accurate and reliable testing results.

Extension of this information collection provides CDC with an

evaluation program to assess the ability of the laboratories to test for drug resistant M. tuberculosis strains. Laboratories also have a self-assessment tool to aid in optimizing their skills in susceptibility testing. The information obtained from the laboratories on susceptibility practices and procedures is used to establish variables related to good performance, assessing training

needs, and aid with the development of practice standards.

Participants in this program include domestic clinical and public health laboratories. Data collection from laboratory participants occurs twice per year. The data collected in this program will include the susceptibility test results of primary and secondary drugs, drug concentrations, and test methods performed by laboratories on a set of performance evaluation (PE) samples. The PE samples are sent to participants twice a year. Participants also report demographic data such as laboratory type and the number of tests performed annually.

There is no cost to respondents to participate other than their time.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondent	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
Domestic Laboratory	Participant Biosafety Compliance Letter of Agreement.	93	2	5/60
	MPEP <i>Mycobacterium tuberculosis</i> Results Worksheet.	93	2	30/60
	Online Survey Instrument	93	2	15/60

Leroy A. Richardson,

Chief, Information Collection Review Office, Office of Scientific Integrity, Office of the Associate Director for Science, Office of the Director, Centers for Disease Control and Prevention.

[FR Doc. 2016–02519 Filed 2–8–16; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Committee to the Director (ACD), Centers for Disease Control and Prevention (CDC): Notice of Charter Renewal

This gives notice under the Federal Advisory Committee Act (Pub. L. 92–463) of October 6, 1972, that the Advisory Committee to the Director, Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (HHS), has been renewed for a 2-year period extending through February 1, 2018.

Contact Person For More Information: Carmen Villar, M.S.W., Designated Federal Officer, Advisory Committee to the Director, CDC, 1600 Clifton Road NE., Mailstop D14, Atlanta, Georgia 30333, Telephone 404–639–7000.

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 Centers for Disease Control and

Prevention and the Agency for Toxic Substances and Disease Registry.

Elaine L. Baker.

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

[FR Doc. 2016–02480 Filed 2–8–16; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[CMS Computer Match No. 2016–12; HHS Computer Match No. 1604; SSA Computer Match No. 1097–1899]

Privacy Act of 1974

AGENCY: Department of Health and Human Services (HHS), Centers for Medicare & Medicaid Services (CMS). **ACTION:** Notice of Computer Matching Program (CMP).

SUMMARY: In accordance with the requirements of the Privacy Act of 1974, as amended, this notice announces the re-establishment of a CMP that CMS plans to conduct with the Social Security Administration (SSA).

DATES: Effective Dates: Comments are invited on all portions of this notice. Public comments are due 30 days after publication. The matching program will become effective no sooner than 40 days after the report of the matching program is sent to the Office of Management and Budget (OMB) and Congress, or 30 days after publication in the Federal Register, whichever is later.

ADDRESSES: The public should send comments to: CMS Privacy Officer,

Division of Security, Privacy Policy & Governance, Information Security & Privacy Group, Office of Enterprise Information, CMS, Room N1–24–08, 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 9:00 a.m.–3:00 p.m., Eastern Time zone.

FOR FURTHER INFORMATION CONTACT:

Elizabeth Kane, Acting Director, Verifications Policy & Operations Division, Eligibility and Enrollment Policy and Operations Group, Center for Consumer Information and Insurance Oversight, CMS, 7501 Wisconsin Avenue, Bethesda, MD 20814, Office Phone: (301) 492–4418, Facsimile: (443) 380–5531, E-Mail: Elizabeth.Kane@cms.hhs.gov.

SUPPLEMENTARY INFORMATION: The Computer Matching and Privacy Protection Act of 1988 (Public Law (Pub. L.) 100-503), amended the Privacy Act (5 U.S.C. 552a) by describing the manner in which computer matching involving Federal agencies could be performed and adding certain protections for individuals applying for and receiving Federal benefits. Section 7201 of the Omnibus Budget Reconciliation Act of 1990 (Pub. L. 101-508) further amended the Privacy Act regarding protections for such individuals. The Privacy Act, as amended, regulates the use of computer matching by Federal agencies when records in a system of records (SOR) are matched with other Federal, state, or local government records. It requires Federal agencies involved in computer matching programs to: