### DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Centers for Disease Control and Prevention

Board of Scientific Counselors, National Center for Injury Prevention and Control, (BSC, NCIPC) Meeting

**AGENCY:** Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (HHS).

**ACTION:** Notice of closed meeting.

**SUMMARY:** In accordance with the Federal Advisory Committee Act, the CDC announces the following meeting for the Board of Scientific Counselors, National Center for Injury Prevention and Control, (BSC, NCIPC).

**DATES:** The meeting will be held on July 12, 2018, 1:00 p.m. to 3:00 p.m., EDT (CLOSED).

ADDRESSES: Teleconference.

#### FOR FURTHER INFORMATION CONTACT:

Gwendolyn H. Cattledge, Ph.D., M.S.E.H., Deputy Associate Director for Science, NCIPC, CDC, 4770 Buford Highway, NE, Mailstop F–63, Atlanta, GA 30341, Telephone (770) 488–1430, Email address: NCIPCBSC@cdc.gov.

SUPPLEMENTARY INFORMATION: The meeting will be closed to the public in accordance with provisions set forth in Section 552b(c)(4) and (6), Title 5 U.S.C., and the Determination of the Director, Management Analysis and Services Office, CDC, pursuant to Public Law 92–463.

Purpose: The Board of Scientific Counselors makes recommendations regarding policies, strategies, objectives, and priorities; and reviews progress toward injury and violence prevention. The Board also provides advice on the appropriate balance of intramural and extramural research, and guidance on the needs, structure, progress and performance of intramural programs. The Board also provides guidance on extramural scientific program matters, including the: (1) Review of extramural research concepts for funding opportunity announcements; (2) conduct of secondary peer review of extramural research grants, cooperative agreements, and contracts applications received in response to the funding opportunity announcements as they relate to the Center's programmatic balance and mission; (3) submission of secondary review recommendations to the Center Director relating to applications to be considered for funding support; (4) review of research portfolios, and (5) review of program proposals.

Matters To Be Considered: The agenda will include discussions on secondary peer review of extramural research grant and cooperative agreement applications received in response to three (3) Notice of Funding Opportunities (NOFO): RFA-CE-18-003, Research on Improving Pediatric mTBI (Mild Traumatic Brain Injury) Outcomes Through Clinician Training, Decision Support, and Discharge Instructions; RFA-CE-18-002, Evaluation of Policies for the Primary Prevention of Multiple Forms of Violence; and RFA-CE-18-004, Research to Evaluate Medication Management of Opioids and Benzodiazepines to Reduce Older Adult Falls. Agenda items are subject to change as priorities dictate.

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 Baker,

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

[FR Doc. 2018–12417 Filed 6–8–18; 8:45 am] BILLING CODE 4163–19–P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifiers: CMS-10394]

Agency Information Collection Activities: Submission for OMB Review; Comment Request

**AGENCY:** Centers for Medicare & Medicaid Services, HHS.

**ACTION:** Notice.

**SUMMARY:** The Centers for Medicare & Medicaid Services (CMS) is announcing an opportunity for the public to comment on ČMS' intention to collect information from the public. Under the Paperwork Reduction Act of 1995 (PRA), federal agencies are required to publish notice in the Federal Register concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, and to allow a second opportunity for public comment on the notice. Interested persons are invited to send comments regarding the burden estimate or any other aspect of this collection of information, including the necessity and

utility of the proposed information collection for the proper performance of the agency's functions, the accuracy of the estimated burden, ways to enhance the quality, utility, and clarity of the information to be collected and the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

DATES: Comments on the collection(s) of information must be received by the OMB desk officer by July 11, 2018.

ADDRESSES: When commenting on the proposed information collections, please reference the document identifier or OMB control number. To be assured consideration, comments and recommendations must be received by the OMB desk officer via one of the following transmissions: OMB, Office of Information and Regulatory Affairs, Attention: CMS Desk Officer, Fax Number: (202) 395–5806 OR, Email: OIRA submission@omb.eop.gov.

To obtain copies of a supporting statement and any related forms for the proposed collection(s) summarized in this notice, you may make your request using one of following:

1. Access CMS' website address at http://www.cms.hhs.gov/ PaperworkReductionActof1995.

2. Email your request, including your address, phone number, OMB number, and CMS document identifier, to *Paperwork@cms.hhs.gov*.

3. Call the Reports Člearance Office at (410) 786–1326.

**FOR FURTHER INFORMATION CONTACT:** Reports Clearance Office at (410) 786–

SUPPLEMENTARY INFORMATION: Under the Paperwork Reduction Act of 1995 (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. The term "collection of information" is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes agency requests 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 publish a 30-day notice in the Federal Register concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, CMS is publishing this notice that summarizes the following proposed collection(s) of information for public comment:

1. Type of Information Collection Request: Extension of a currently

approved collection;

Title of Information Collection: Application to be Qualified Entity to Receive Medicare Data for Performance Measurement: Use: The Patient Protection and Affordable Care Act (ACA) was enacted on March 23, 2010 (Pub. L. 111–148). ACA amends section 1874 of the Social Security Act by adding a new subsection (e) to make standardized extracts of Medicare claims data under Parts A, B, and D available to qualified entities to evaluate the performance of providers of services and suppliers. This is the application needed to determine an organization's eligibility as a qualified entity. To implement the requirements outlined in the legislation, CMS established the Qualified Entity Certification Program (QECP) to evaluate an organization's eligibility across three areas: Organizational and governance capabilities, addition of claims data from other sources (as required in the statute), and data privacy and security. This collection covers the application through which organizations provide information to CMS to determine whether they will be approved as a qualified entity. Form Number: CMS-10394 (OMB control number: 0938-1144); Frequency: Reporting-Yearly; Affected Public: Private Sector (State, Local, or Tribal Governments, Business or other for-profits, Not-for-Profit Institutions); Number of Respondents: 30; Total Annual Responses: 10; Total Annual Hours: 5,000, (For policy questions regarding this collection contact Kari Gaare at 410-786-8612.)

Dated: June 5, 2018.

#### William N. Parham, III,

Director, Paperwork Reduction Staff, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. 2018–12437 Filed 6–8–18; 8:45 am] BILLING CODE 4120–01–P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Administration for Children and Families

#### Submission for OMB Review; Comment Request

*Title:* The Evaluation of Child Welfare Information Gateway.

OMB No.: New Collection.

(Note: Some of the data collection activities proposed for the Evaluation of Child Welfare Information Gateway were previously approved via Fast Track OMB Clearance. We are seeking regular OMB approval so that future evaluation findings may be publicly disseminated in reports, journals and at conferences to better inform the child welfare field.)

Description: The Children's Bureau (CB), Administration for Children and Families (ACF), U.S. Department of Health and Human Services (HHS) is proposing new or expanded data collection activities as part of its Evaluation of Child Welfare Information Gateway.

Child Welfare Information Gateway (CWIG) is a service of the Children's Bureau, a component within the Administration for Children and Families, and is dedicated to the mission of connecting professionals and concerned citizens to information on programs, research, legislation, and statistics regarding the safety, permanency, and well-being of children and families. The Evaluation of Child Welfare Information Gateway was initiated in response to Executive Order 12862 issued on September 11, 1993. The Order calls for putting customers first and striving for a customer-driven government that matches or exceeds the best service available in the private sector. To that end, CWIG's evaluation is designed to better understand the kind and quality of information services that customers want, as well as customers' level of satisfaction with existing services.

A new Market Research Sub-Study is also being proposed as part of this submission to complement information obtained from the larger Evaluation of Child Welfare Information Gateway. The sub-study component seeks to learn more about how child welfare professionals and students planning to enter the child welfare workforce access and consume work-related information. This national study will focus on understanding child welfare professionals' and students' characteristics, use of technology, and preferences for obtaining information that they use in their work. The goal of the sub-study is to provide child welfare technical assistance providers and other organizations with a better understanding of their target audiences so they can design more effective

products, services, and dissemination strategies to reach these populations.

Data collection activities proposed for the Evaluation of Child Welfare Information Gateway include: ten online targeted surveys designed to evaluate CWIG's special initiative websites and other targeted website sections; ten online event surveys administered after CWIG-sponsored webinars, presentations, or other events; five focus groups (each with approximately 10 participants) with users and non-users of CWIG's special initiative websites and other CWIG products and services; and, a general customer survey delivered via multiple modes (e.g., website, email, live chat, print, and phone). The sampling plan for the CWIG general customer survey is designed to reach the various types of customers using Child Welfare Information Gateway services such as professionals, students, and customers looking for assistance with a personal situation while reducing burden for respondents by only asking relevant questions for their backgrounds.

The market research sub-study seeks to deliver surveys and conduct focus groups to gauge online information habits and preferences. The proposed market research sub-study will consist of a national online survey of child welfare professionals and students, which will be administered through four different instruments tailored for four different populations. Ten focus groups (each with 8 to 10 participants) will be used to learn more about different audiences' habits and preferences related to child welfare information access and consumption.

Respondents: The Evaluation of Child Welfare Information Gateway will target all types of possible CWIG users including: State and local governments, the territories, service providers, Tribes and tribal organizations, grantees, researchers, and the general public seeking information and resources from Child Welfare Information Gateway via the website, mail, telephone, Live Chat, and email. The Market Research Sub-Study will target child welfare professionals in state, county, tribal, and private agencies; Court Improvement Program coordinators and directors: judges and attorneys involved in child welfare-related work; and students in Bachelor's and Master's degree programs in social work that receive Title IV–E or IV–B stipends.