products or services. These collections will allow for ongoing, collaborative and actionable communications between the Agency and its customers and stakeholders. It will also allow feedback to contribute directly to the improvement of program management.

Feedback collected under this generic clearance will provide useful information, but it will not yield data that can be generalized to the overall population. This type of generic clearance for qualitative information will not be used for quantitative information collections that are designed to yield reliably actionable results, such as monitoring trends over time or documenting program performance. Such data uses require more rigorous designs that address: the target population to which generalizations will be made, the sampling frame, the sample design (including stratification and clustering), the precision requirements or power calculations that justify the proposed sample size, the expected response rate, methods for assessing potential non-response bias, the protocols for data collection, and any testing procedures that were or will be undertaken prior fielding the study. Depending on the degree of influence the results are likely to have, such collections may still be eligible for submission for other generic mechanisms that are designed to yield quantitative results.

The Agency received 0 comments were received in response to the 60-day notice published in the Federal Register of March 14, 2017 (79 FR 18692).


Type of Review: New Collection.

Affected Public: Individuals, households, professionals, public/private sector.

Average Expected Annual Number of Activities: 600.

Respondents per Activity: 50.

Annual Responses: 30,000.

Frequency of Response: Once per request.

Average Minutes per Response: 30.

Burden Hours: 500,000 hours annually.

An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid Office of Management and Budget control number.

Darius Taylor,

Deputy Information Collection Officer.

[FR Doc. 2017–12046 Filed 6–9–17; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Center for Scientific Review; Notice of Closed Meetings

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meetings.

The meetings will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: Center for Scientific Review Special Emphasis Panel; Member Conflict: Epidemiology.

Date: June 19, 2017.

Time: 1:00 p.m. to 4:00 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892

[Telephone Conference Call].

Contact Person: Heidi B Friedman, PhD., Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 1012A, MSC 7770, Bethesda, MD 20892, 301–370–5632, hfriedman@csr.nih.gov.

This notice is being published less than 15 days prior to the meeting due to the timing limitations imposed by the review and funding cycle.

Name of Committee: Center for Scientific Review Special Emphasis Panel; Experimental and Bioinformatic Approaches in the Druggable Genome.

Date: June 26, 2017.

Time: 10:00 a.m. to 5:00 p.m.

Agenda: To review and evaluate grant applications and/or proposals.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892

[Virtual Meeting].

Contact Person: Luis Dettin, PhD., Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 2208, Bethesda, MD 20892, 301 451 1327, dettinle@csr.nih.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel; Mentored Training in Comparative and Veterinary Medicine.

Date: June 27, 2017.

Time: 2:00 p.m. to 4:00 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892

[Telephone Conference Call].

Contact Person: Amy Kathleen Wernimont, PhD., Scientific Review Officer, Center for Scientific Review, 6701 Rockledge Drive, Bethesda, MD 20892, 301–827–6427, amy.wernimont@nih.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel; Immigrant Women’s Health.

Date: June 30, 2017.

Time: 8:00 a.m. to 5:00 p.m.

Agenda: To review and evaluate grant applications.

Place: The Dupont Hotel, 1500 New Hampshire Avenue NW., Washington, DC 20036.

Contact Person: Martha L Hare, RN, PhD., Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 3154, MSC 7770, Bethesda, MD 20892, (301) 451–8504, hareml@email.nih.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel; Member Conflict: Acute Brain Injury and Regeneration.

Date: July 3, 2017.

Time: 2:00 p.m. to 5:00 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892

[Telephone Conference Call].

Contact Person: Alexander Yakovlev, PhD., Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 5206, MSC 7846, Bethesda, MD 20892–7846, 301–435–1254, yakovleva@csr.nih.gov.


Dated: June 6, 2017.

Sylvia L. Neal,

Program Analyst, Office of Federal Advisory Committee Policy.

[FR Doc. 2017–12030 Filed 6–9–17; 8:45 am]

BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Substance Abuse and Mental Health Services Administration

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

Periodically, the Substance Abuse and Mental Health Services Administration (SAMHSA) will publish a summary of information collection requests under OMB review, in compliance with the Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these documents, call the SAMHSA Reports Clearance Officer on (240) 276–1243.

[FR Doc. 2017–12046 Filed 6–9–17; 8:45 am]
Project: Project—Division of State Programs—Management Reporting Tool (DSP–MRT) (OMB No. 0930–0354)—Revision

The Substance Abuse and Mental Health Services Administration (SAMHSA)’s Center for Substance Abuse Prevention (CSAP) aims to address two of SAMHSA’s top substance abuse prevention priorities: Underage drinking (UAD; age 12 to 20) and prescription drug misuse and abuse (PDM; age 12 to 25) through the Division of State Program—Monitoring and Reporting Tool. This data collection will allow all DSP programs to report into a standard tool that aligns with the Strategic Prevention Framework model. This request for data collection includes a revision from a previously approved OMB instrument formally known as Partnerships for Success-Management and Reporting Tool.

Monitoring data on SPF model will allow SAMHSA project officers to systematically collect data to monitor their grant program performance and outcomes along with grantee technical assistance needs. In addition to assessing activities related to the SPF steps, the performance monitoring instruments covered in this statement collect data to assess the following:

- Number of training and technical assistance activities per funded community provided by the grantee to support communities;
- Reach of training and technical assistance activities (numbers served) provided by the grantee;
- Percentage of subrecipient communities that submit data to the grantee data system;
- Number of sub-recipient communities that improved on one or more targeted NOMs indicators (Outcome);
- Number of grantees who integrate Prescription Drug Monitoring Data into their program needs assessment.

Changes to this package include the following:

- Standard language for all DSP–MRT questions;
- New disparities module to align with SAMHSA’s monitoring requirements;
- Updated technical assistance section;
- Deletion of cost questions specific to funding amounts and in-kind resources;
- Deletion of advisory council and other workgroup sub-committee questions;
- Addition of Section A specific to SPF-Rx questions;
- Addition of Section B specific to PDO questions;

### ANNUALIZED DATA COLLECTION BURDEN

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Written comments and recommendations concerning the proposed information collection should be sent by July 12, 2017 to the SAMHSA Desk Officer at the Office of Information and Regulatory Affairs, Office of Management and Budget (OMB). To ensure timely receipt of comments, and to avoid potential delays in OMB’s receipt and processing of mail sent through the U.S. Postal Service, commenters are encouraged to submit their comments to OMB via email to: OIRA_Submission@omb.eop.gov. Although commenters are encouraged to send their comments via email, commenters may also fax their comments to: (202) 395–7285.

Commenters may also mail them to: Office of Management and Budget, Office of Information and Regulatory Affairs, New Executive Office Building, Room 10102, Washington, DC 20503.

Summer King, Statistician.

[FR Doc. 2017–12090 Filed 6–9–17; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Substance Abuse and Mental Health Services Administration

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

Periodically, the Substance Abuse and Mental Health Services Administration (SAMHSA) will publish a summary of information collection requests under OMB review, in compliance with the Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these documents, call the SAMHSA Reports Clearance Officer on (240) 276–1243.

Project: Mental Health Block Grant Ten Percent Set Aside Evaluation of First Episode Psychosis—NEW

The Substance Abuse and Mental Health Services Administration (SAMHSA) is directed by Congress through its FY 2016 Omnibus bill, Public Law 114–113, to set aside ten percent of the Mental Health Block Grant (MHBG) allocation for each state to support evidence-based programs that provide treatment for those with early serious mental illness (SMI) and a first episode psychosis (FEP)—an increase from the previous five percent set aside.

The purpose of this 3-year evaluation is to assess the relationship between fidelity of selected coordinated specialty care (CSC) programs supported with Mental Health Block Grant (MHBG) Ten Percent Set Aside funding and participant outcomes. There are approximately 250 sites implementing CSC programs with MHBG ten percent set aside funding. All 250 sites will be asked to report on their implementation through an online survey. Up to 32 CSC sites across the nation will be recruited to participate in a process and outcome evaluation. The data collection activities for the Mental Health Block Grant Ten Percent Set Aside Evaluation will include the following six data collection tools:

- Site Survey: This is a one-time online survey with site directors of all 250 centers using MHBG ten percent set aside funding (not just those included in the evaluation). The survey focuses on how centers across the U.S. are providing services to individuals with First Episode Psychosis (FEP) in their communities.