

regulatory and supervisory assistance, when appropriate, to facilitate the testing of innovative and advanced technologies, products, services, systems, or activities.

The FDIC anticipates that products developed as part of innovation pilot programs will improve the efficiency and effectiveness of bank operations, and eventually, examinations, while increasing transparency and ultimately reducing the cost of regulatory compliance for participating institutions. In addition, the FDIC anticipates that proposals provided in connection with the innovation pilot programs will involve cutting-edge innovations and novel approaches or applications involving a banking product, service, system, or activity that benefits and can lead to better outcomes for consumers.

As part of an innovation pilot program, innovators may request information from banks and other members of the public outside of their normal course of business. Any information provided by banks and other members of the public will be provided on a voluntary basis. FDIC staff may similarly request information on a voluntary basis from banks or other members of the public to evaluate the products or services developed in the pilot programs. This information is intended to allow banks and the FDIC to analyze the health of the overall banking system, critical financial sectors, or national, regional or local economic conditions (*i.e.*, horizontal analysis). Additionally, bank specific information may be collected in order to allow for better insights into current and escalating risks across all aspects of banking. In particular, innovators may request from banks and other members of the public general ledger information about all products and services, or a subset of products and services, systems or activities. Information requested will not contain any personally identifiable information (PII) as defined in OMB Circular A-130 or include the disclosure of any financial records or information which is identified with or identifiable as being derived from the financial records of a particular customer.

The annual burden for this information collection is estimated to be 40,000 hours. This represents an increase of hours from the current burden estimate and also a change in focus. In particular, when this information collection was first obtained, it included the burden imposed on the innovators and partner banks. In review of this information collection, the FDIC has decided to

transfer the burden imposed on innovators to existing information collection 3064-0072 entitled, "Acquisition Services Information Requirements," which is related to the FDIC's procurement process. The remaining hours in this information collection, which have been updated and increased, reflect the burden imposed on banks and other members of the public in connection with innovation pilot programs.

Request for Comment

Comments are invited on: (a) Whether the collection of information is necessary for the proper performance of the FDIC's functions, including whether the information has practical utility; (b) the accuracy of the estimate of the burden of the information collection, including the validity of the methodology and assumptions used; (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. All comments will become a matter of public record.

Federal Deposit Insurance Corporation.

Dated at Washington, DC, on November 23, 2021.

James P. Sheesley,

Assistant Executive Secretary.

[FR Doc. 2021-25924 Filed 11-26-21; 8:45 am]

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FEDERAL ELECTION COMMISSION

Sunshine Act Meeting

TIME AND DATE: Thursday, December 2, 2021 at the conclusion of the open meeting on December 2, 2021.

PLACE: 1050 First Street NE, Washington, DC (This meeting will be a virtual meeting).

STATUS: This meeting will be closed to the public.

MATTERS TO BE CONSIDERED: Compliance matters pursuant to 52 U.S.C. 30109.

Matters relating to internal personnel decisions, or internal rules and practices.

Information the premature disclosure of which would be likely to have a considerable adverse effect on the implementation of a proposed Commission action.

Matters concerning participation in civil actions or proceedings or arbitration.

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CONTACT PERSON FOR MORE INFORMATION: Judith Ingram, Press Officer, Telephone: (202) 694-1220.

Vicktorija J. Allen,

Acting Deputy Secretary of the Commission.

[FR Doc. 2021-26000 Filed 11-24-21; 11:15 am]

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FEDERAL FINANCIAL INSTITUTIONS EXAMINATION COUNCIL

[Docket No. AS21-08]

Appraisal Subcommittee Notice of Meeting

AGENCY: Appraisal Subcommittee of the Federal Financial Institutions Examination Council.

ACTION: Notice of meeting.

SUPPLEMENTARY INFORMATION: In accordance with Section 1104(b) of Title XI of the Financial Institutions Reform, Recovery, and Enforcement Act of 1989, as amended, notice is hereby given that the Appraisal Subcommittee (ASC) will meet in open session for a special meeting:

Location: Due to the COVID-19 Pandemic, the meeting will be open to the public via live webcast only. Visit the agency's homepage (www.asc.gov) and access the provided registration link in the What's New box. You MUST register in advance to attend this Meeting.

Date: December 8, 2021.

Time: 11:00 a.m. ET.

Status: Open.

Reports

Chairman
Executive Director
Grants Director
Financial Manager

Action and Discussion Items

Approval of Minutes
September 15, 2021 Open Session
Quarterly Meeting
Notice of Proposed Rulemaking on
Temporary Waiver

How To Attend and Observe an ASC Meeting

Due to the COVID-19 Pandemic, the meeting will be open to the public via live webcast only. Visit the agency's homepage (www.asc.gov) and access the provided registration link in the What's New box. The meeting space is intended to accommodate public attendees. However, if the space will not accommodate all requests, the ASC may refuse attendance on that reasonable basis. The use of any video or audio tape recording device, photographing

device, or any other electronic or mechanical device designed for similar purposes is prohibited at ASC Meetings.

James R. Park,

Executive Director.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Board on Radiation and Worker Health (ABRWH), Subcommittee for Dose Reconstruction Reviews (SDRR), National Institute for Occupational Safety and Health (NIOSH)

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

ACTION: Notice of meeting.

SUMMARY: In accordance with the Federal Advisory Committee Act, the CDC announces the following meeting for the Subcommittee for Dose Reconstruction Reviews (SDRR) of the Advisory Board on Radiation and Worker Health (ABRWH or the Advisory Board). This meeting is open to the public, but without a public comment period. The public is welcome to submit written comments in advance of the meeting, to the contact person below. Written comments received in advance of the meeting will be included in the official record of the meeting. The public is also welcomed to listen to the meeting by joining the audio conference (information below). The audio conference line has 150 ports for callers.

DATES: The meeting will be held on January 19, 2022, from 10:30 a.m. to 4:00 p.m., EST.

Written comments must be received on or before January 12, 2022.

ADDRESSES: You may submit comments by mail to: Sherri Diana, National Institute for Occupational Safety and Health, 1090 Tusculum Avenue, MS C-34, Cincinnati, Ohio 45226. Meeting Information: Audio Conference Call via FTS Conferencing. The USA toll-free dial-in number is 1-866-659-0537; the pass code is 9933701.

FOR FURTHER INFORMATION CONTACT: Rashaun Roberts, Ph.D., Designated Federal Officer, NIOSH, CDC, 1090 Tusculum Avenue, Mailstop C-24, Cincinnati, Ohio 45226, Telephone: (513) 533-6800; Toll Free 1(800) CDC-INFO; Email: ocas@cdc.gov.

SUPPLEMENTARY INFORMATION:

Background: The Advisory Board was established under the Energy Employees Occupational Illness Compensation Program Act of 2000 to advise the President on a variety of policy and technical functions required to implement and effectively manage the new compensation program. Key functions of the Advisory Board include providing advice on the development of probability of causation guidelines that have been promulgated by the Department of Health and Human Services (HHS) as a final rule; advice on methods of dose reconstruction, which have also been promulgated by HHS as a final rule; advice on the scientific validity and quality of dose estimation and reconstruction efforts being performed for purposes of the compensation program; and advice on petitions to add classes of workers to the Special Exposure Cohort (SEC). In December 2000, the President delegated responsibility for funding, staffing, and operating the Advisory Board to HHS, which subsequently delegated this authority to CDC. NIOSH implements this responsibility for CDC.

The Advisory Board's charter was issued on August 3, 2001, renewed at appropriate intervals, and rechartered under Executive Order 13889 on March 22, 2020, and will terminate on March 22, 2022.

Purpose: The Advisory Board is charged with (a) providing advice to the Secretary, HHS, on the development of guidelines under Executive Order 13179; (b) providing advice to the Secretary, HHS, on the scientific validity and quality of dose reconstruction efforts performed for this program; and (c) upon request by the Secretary, HHS, advise the Secretary on whether there is a class of employees at any Department of Energy facility who were exposed to radiation but for whom it is not feasible to estimate their radiation dose, and on whether there is reasonable likelihood that such radiation doses may have endangered the health of members of this class. SDRR was established to aid the Advisory Board in carrying out its duty to advise the Secretary, HHS, on dose reconstruction.

Matters to be Considered: The agenda will include discussions on the following dose reconstruction program quality management and assurance activities: Dose reconstruction cases under review from Sets 29 and 30, possibly including cases involving: Nevada test Site, Oak Ridge Institute for Science Education, Rocky Flats Plant, Idaho National Laboratory, Y-12 Plant, Clarksville Modification Center, Pantex Plant, Oak Ridge National Laboratory

(X-10), Albuquerque Operations Office, General Atomics, Area IV of the Santa Susanna Field Laboratory, Oak Ridge Gaseous Diffusion Plant (K-25), Savannah River Site (SRS), Hanford, SRS, X-10, Y-12 Plant, and General Steel Industries. Agenda items are subject to change as priorities dictate.

The Director, Strategic Business Initiatives Unit, Office of the Chief Operating Officer, Centers for Disease Control and Prevention, 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.

Kalwant Smagh,

Director, Strategic Business Initiatives Unit, Office of the Chief Operating Officer, Centers for Disease Control and Prevention.

[FR Doc. 2021-25863 Filed 11-26-21; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Understanding the Value of Centralized Services Study (New Collection)

AGENCY: Office of Planning, Research, and Evaluation, Administration for Children and Families, HHS.

ACTION: Request for public comment.

SUMMARY: The Administration for Children and Families (ACF), U.S. Department of Health and Human Services (HHS) is proposing a new data collection activity as part of the Understanding the Value of Centralized Services study. The objective of this descriptive study is to understand the advantages, disadvantages, and costs of centralizing services for individuals and families with low incomes.

DATES: *Comments due within 30 days of publication.* OMB must make a decision about the collection of information between 30 and 60 days after publication of this document in the **Federal Register**. Therefore, a comment is best assured of having its full effect if OMB receives it within 30 days of publication.

ADDRESSES: Written comments and recommendations for the proposed information collection should be sent within 30 days of publication of this notice to www.reginfo.gov/public/do/PRAMain. Find this particular information collection by selecting