

of 1989 (FIRREA) agencies¹ conduct the FR 29a survey jointly. The FR 29b is collected by the Board only.

Legal authorization and confidentiality: The Board's Legal Division has determined that the FR 29a and FR29b surveys are voluntary and authorized by sections 10(4) and 11(1) of the Federal Reserve Act (12 U.S.C. 244 and 248(1)), which authorize the Board to determine employees' compensation. The FR 29a survey is completed by an outside consultant that submits to the Board a report of the survey containing only aggregate data. Because the Board does not collect or have access to the individual respondent data, no confidentiality issue arises with respect to the individual responses to the FR 29a. The Board does not consider the report containing aggregate data to be confidential. The FR 29b consists of ad hoc surveys conducted by the Board during the year to collect information on specific salary and non-salary matters that affect Board employees. The ability of the Board to maintain the confidentiality of information provided by respondents to the FR 29b surveys will have to be determined on a case by case basis depending on the data collected under a particular survey. Some of the information collected on the surveys may be protected from Freedom of Information Act (FOIA) disclosure by FOIA exemptions 4 and 6. (5 U.S.C. 552 (b)(4) and (6)). Exemption 4 protects from disclosure trade secrets and commercial or financial information, while Exemption 6 protects information "the disclosure of which would constitute a clearly unwarranted invasion of personal privacy."

Consultation outside the agency: Towers Watson and the Board work together to review and update the FR 29a survey instrument.

Current actions: On June 29, 2017 the Board published a notice in the **Federal Register** (82 FR 29564) requesting public comment for 60 days on the extension, without revision, of the Compensation and Salary Surveys. The comment period for this notice expired on August 28, 2017. The Board did not receive any comments.

¹ For purposes of this proposal, the FIRREA agencies consist of: The Federal Reserve Board, the Office of the Comptroller of the Currency, the Federal Deposit Insurance Corporation, the National Credit Union Administration, the Commodities Futures Trading Commission, the Farm Credit Administration, and the Securities and Exchange Commission.

Board of Governors of the Federal Reserve System, September 6, 2017.

Ann E. Misback,

Secretary of the Board.

[FR Doc. 2017-19168 Filed 9-8-17; 8:45 am]

BILLING CODE 6210-01-P

FEDERAL RETIREMENT THRIFT INVESTMENT BOARD

Sunshine Act; Notice of Board Member Meeting: Federal Retirement Thrift Investment Board

AGENDA: Federal Retirement Thrift Investment Board Members' Meeting, September 18, 2017, 8:30 a.m. (In-Person).

Open Session

1. Approval of the Minutes of the August 28, 2017 Board Members' Meeting
2. Monthly Reports
 - (a) Participant Activity Report
 - (b) Investment Performance Report
 - (c) Legislative Report
3. Calendar Review 2017/2018
4. FY 18 Budget Review and Approval
5. Vendor Financials
6. The Office of Participant Services' Annual Report
7. Blended Retirement Update
8. IT Update

Closed Session

Information covered under 5 U.S.C. 552b(c)(4) and (c)(9)(B).

CONTACT PERSON FOR MORE INFORMATION: Kimberly Weaver, Director, Office of External Affairs, (202) 942-1640.

Dated: September 7, 2017.

Megan Grumbine,

General Counsel, Federal Retirement Thrift Investment Board.

[FR Doc. 2017-19268 Filed 9-7-17; 4:15 pm]

BILLING CODE 6760-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Meeting of the Community Preventive Services Task Force (Task Force)

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

ACTION: Notice of meeting.

SUMMARY: The Centers for Disease Control and Prevention (CDC) within the Department of Health and Human Services announces the next meeting of the Community Preventive Services

Task Force (Task Force) on October 18-19, 2017 in Atlanta, Georgia.

DATES: The meeting will be held on Wednesday, October 18, 2017 from 8:30 a.m. to 6:00 p.m. EDT and Thursday, October 19, 2017 from 8:30 a.m. to 1:00 p.m. EDT.

ADDRESSES: The Task Force Meeting will be held at the CDC Edward R. Roybal Campus, Centers for Disease Control and Prevention Headquarters (Building 19), 1600 Clifton Road NE., Atlanta, GA 30329. You should be aware that the meeting location is in a Federal government building; therefore, Federal security measures are applicable. For additional information, please see Roybal Campus Security Guidelines under **SUPPLEMENTARY INFORMATION**. Information regarding meeting logistics will be available on the Community Guide Web site (www.thecommunityguide.org) closer to the date of the meeting.

FOR FURTHER INFORMATION CONTACT: Onslow Smith, Center for Surveillance, Epidemiology and Laboratory Services; Centers for Disease Control and Prevention, 1600 Clifton Road NE., MS-E-69, Atlanta, GA 30329, phone: (404) 498-6778, email: CPSTF@cdc.gov.

SUPPLEMENTARY INFORMATION:

Meeting accessibility: This space-limited meeting is open to the public. All meeting attendees must register. To ensure completion of required security procedures and access to the CDC's Global Communications Center, U.S. citizens intending to attend in person must register by October 11, 2017. Non-U.S. citizens intending to attend in person must register by September 20, 2017. Failure to register by the dates identified could result in the inability to attend the Task Force meeting in person.

Those unable to attend the meeting in person are able to do so via Webcast. CDC will send the Webcast URL to registrants upon receipt of their registration. All meeting attendees must register by October 13, 2017 to receive the webcast information. CDC will email webcast information from the CPSTF@cdc.gov mailbox.

Public comment: A public comment period, limited to three minutes per person, will follow the Task Force's discussion of each systematic review. Individuals wishing to make public comments must indicate their desire to do so with their registration by providing their name, organizational affiliation, and the topic to be addressed (if known). Public comments will become part of the meeting summary. Public comment is not possible via Webcast.

Background on the Task Force: The Task Force is an independent, nonfederal panel whose members are appointed by the CDC Director. Task Force members represent a broad range of research, practice, and policy expertise in prevention, wellness, health promotion, and public health. The Task Force was convened in 1996 by the Department of Health and Human Services (HHS) to identify community preventive programs, services, and policies that increase healthy longevity, save lives and dollars, and improve Americans' quality of life. CDC is mandated to provide ongoing administrative, research, and technical support for the operations of the Task Force. During its meetings, the Task Force considers the findings of systematic reviews on existing research and practice-based evidence and issues recommendations. Task Force recommendations are not mandates for compliance or spending. Instead, they provide information about evidence-based options that decision makers and stakeholders can consider when they are determining what best meet the specific needs, preferences, available resources, and constraints of their jurisdictions and constituents. The Task Force's recommendations, along with the systematic reviews of the evidence on which they are based, are compiled in the *Guide to Community Preventive Services (The Community Guide)*.

Matters proposed for discussion: Cardiovascular Disease Prevention (Mobile Health Interventions for Cardiovascular Disease Prevention); Diabetes Prevention and Control (Lifestyle Interventions to Reduce Risk of Gestational Diabetes); Nutrition (Gardening-Based Interventions to Increase Fruit and Vegetable Intake); Obesity Prevention and Control (School-based Diet and Physical Activity Interventions); and Women's Health (Primary Prevention of Intimate Partner Violence and Sexual Violence Among Youth). The agenda is subject to change without notice.

Roybal Campus Security Guidelines: The Edward R. Roybal Campus is the headquarters of the CDC and is located at 1600 Clifton Road NE., Atlanta, Georgia. The meeting is being held in a Federal government building; therefore, Federal security measures are applicable.

All meeting attendees must register by the dates outlined under *Meeting Accessibility*. In planning your arrival time, please take into account the need to park and clear security. All visitors must enter the Edward R. Roybal Campus through the front entrance on Clifton Road. Vehicles may be searched,

and the guard force will then direct visitors to the designated parking area. Upon arrival at the facility, visitors must present government-issued photo identification (e.g., a valid federal identification badge, state driver's license, state non-driver's identification card, or passport). Non-U.S. citizens must complete the required security paperwork prior to the meeting date and must present a valid passport, visa, Permanent Resident Card, or other type of work authorization document upon arrival at the facility. All persons entering the building must pass through a metal detector. CDC Security personnel will issue a visitor's ID badge at the entrance to Building 19. Visitors may receive an escort to the meeting room. All items brought to HHS/CDC are subject to inspection.

Dated: September 6, 2017.

Sandra Cashman,

Executive Secretary, Centers for Disease Control and Prevention.

[FR Doc. 2017-19203 Filed 9-8-17; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2017-N-4851]

Vaccines and Related Biological Products Advisory Committee; Notice of Meeting

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) announces a forthcoming public advisory committee meeting of the Vaccines and Related Biological Products Advisory Committee (VRBPAC). The general function of the committee is to provide advice and recommendations to the Agency on FDA's regulatory issues. The meeting will be open to the public. Members will participate via teleconference.

DATES: The meeting will be held on October 4, 2017, from 1 p.m. to 4:30 p.m.

ADDRESSES: FDA White Oak Campus, 10903 New Hampshire Ave., Bldg. 31 Conference Center, the Great Room (Rm. 1503), Silver Spring, MD 20993-0002. For those unable to attend in person, the meeting will also be webcast and will be available at the following link: <https://collaboration.fda.gov/cbervrpbac2017>. Answers to commonly asked questions including information regarding special

accommodations due to a disability, visitor parking, and transportation may be accessed at: <https://www.fda.gov/AdvisoryCommittees/AboutAdvisoryCommittees/ucm408555.htm>.

FOR FURTHER INFORMATION CONTACT:

Serina Hunter-Thomas or Rosanna Harvey, Center for Biologics Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 71, Rm. 6307C, Silver Spring, MD 20993-0002, 240-402-5771 serina.hunter-thomas@fda.hhs.gov and 240-402-8072, rosanna.harvey@fda.hhs.gov, or FDA Advisory Committee Information Line, 1-800-741-8138 (301-443-0572 in the Washington, DC area). A notice in the **Federal Register** about last minute modifications that impact a previously announced advisory committee meeting cannot always be published quickly enough to provide timely notice. Therefore, you should always check the Agency's Web site at <https://www.fda.gov/AdvisoryCommittees/default.htm> and scroll down to the appropriate advisory committee meeting link, or call the advisory committee information line to learn about possible modifications before coming to the meeting.

SUPPLEMENTARY INFORMATION:

Agenda: On October 4, 2017, the VRBPAC will meet in an open session to discuss and make recommendations on the selection of strains to be included in an influenza virus vaccine for the 2018 southern hemisphere influenza season. FDA intends to make background material available to the public no later than 2 business days before the meeting. If FDA is unable to post the background material on its Web site prior to the meeting, the background material will be made publicly available at the location of the advisory committee meeting, and the background material will be posted on FDA's Web site after the meeting. Background material is available at <https://www.fda.gov/AdvisoryCommittees/Calendar/default.htm>. Scroll down to the appropriate advisory committee meeting link.

Procedure: Interested persons may present data, information, or views, orally or in writing, on issues pending before the committee. Written submissions may be made to the contact person on or before September 27, 2017. Oral presentations from the public will be scheduled between approximately 1:15 p.m. and 2:15 p.m. Those individuals interested in making formal oral presentations should notify the contact person and submit a brief