

**DEPARTMENT OF HEALTH AND
HUMAN SERVICES**
Food and Drug Administration
[Docket No. FDA-2019-D-3592]
**Agency Information Collection
Activities; Submission for Office of
Management and Budget Review;
Comment Request; Certificates of
Confidentiality**
AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that a proposed collection of information has been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995.

DATES: Submit written comments (including recommendations) on the collection of information by July 17, 2020.

ADDRESSES: To ensure that comments on the information collection are received, OMB recommends that written comments be submitted to www.reginfo.gov/public/do/PRAMain. Find this particular information collection by selecting “Currently under Review—Open for Public Comments” or by using the search function. All comments should be identified with the OMB control number 0910–0130. Also include the FDA docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: Domini Bean, Office of Operations, Food and Drug Administration, Three White Flint North, 10A–12M, 11601 Landsdown St., North Bethesda, MD 20852, 301–796–5733, PRASStaff@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: In compliance with 44 U.S.C. 3507, FDA has submitted the following proposed collection of information to OMB for review and clearance.

**Protection of Human Subjects and
Institutional Review Boards;
Certificates of Confidentiality**
**OMB Control Number 0910–0130—
Revision**

This information collection supports Agency guidance regarding the issuance of Certificates of Confidentiality (CoCs). The 21st Century Cures Act (Cures Act) (Pub. L. 114–255, section 2012) amended the Public Health Service Act, section 301(d) (42 U.S.C. 241(d)), to help protect the privacy of human subject research participants from whom identifiable, sensitive information is being collected or used in furtherance of the research. Historically, a CoC generally protected a researcher from being compelled in a legal proceeding (such as by subpoena or court order) to disclose identifiable and sensitive information about the research participant, created or compiled for purposes of the human subject research. The Cures Act broadened the protections of the statutory provision by affirmatively prohibiting holders of CoCs from disclosing such information unless a specific exception applies. For efficiency of Agency operations, we are revising information collection currently approved under OMB Control No. 0910–0130 pertaining to the protection of human subjects and institutional review boards to include information collection pertaining to the issuance of CoCs. As information collection activity is planned and undertaken by FDA, we find consolidating related collection elements better utilizes our resources. We have developed guidance to assist respondents to the information collection with this topic and are including it in the information collection accordingly.

The Cures Act simplified certain aspects of the issuance of CoCs by requiring that CoCs be issued for federally funded human subject research that collects or uses identifiable, sensitive information (referred to in the draft guidance as mandatory CoCs). For non-federally funded research, issuance of CoCs is not required but may be issued at the discretion of FDA (referred to in the

draft guidance as discretionary CoCs) when the study involves a product subject to FDA’s jurisdiction and regulatory authority. FDA intends to continue receiving such requests and will issue discretionary CoCs as appropriate.

To assist respondents with information collection attendant to CoCs, we developed the draft guidance document entitled “Certificates of Confidentiality; Draft Guidance for Sponsors, Sponsor-Investigators, Researchers, Industry, and Food and Drug Administration Staff.” The draft guidance is intended to provide information on how to request a discretionary CoC, the statutory requirements for requesting such a CoC, and the statutory responsibilities associated with possessing a CoC. Although the mandatory CoC and the discretionary CoC are issued under different processes, the protections afforded by the issuance of either CoC are identical and the statutory responsibilities are applicable to both. The draft guidance was developed and issued consistent with our Good Guidance Practice regulations at 21 CFR 10.115 which provide for comment at any time. The draft guidance is available at: <https://www.fda.gov/media/132966/download>. We intend to finalize the draft guidance and are revising the associated information collection accordingly.

In the **Federal Register** of November 25, 2019 (84 FR 64906), we published a notice announcing the availability of the draft guidance entitled “Certificates of Confidentiality; Draft Guidance for Sponsors, Sponsor-Investigators, Researchers, Industry, and Food and Drug Administration Staff; Availability,” including an analysis of burden that may be attributable to the information collection recommendations. Although we received some comments requesting that clarifying discussion be included with regard to topics covered in the guidance, no comments were received in response to the four information collection topics solicited under the PRA.

We estimate the burden of this collection of information as follows:

TABLE 1—ESTIMATED ANNUAL REPORTING BURDEN ¹

Draft guidance for sponsors, sponsor-investigators, researchers, industry, and FDA staff on CoCs	Number of respondents	Number of responses per respondent	Total annual responses	Average burden per response	Total hours
Submissions of CoC Requests From Sponsors, Sponsor-Investigators, or Authorized Representatives	150	1	150	2	300

¹ There are no capital costs or operating and maintenance costs associated with this collection of information.

Based on the number of CoC requests we have received prior to the Cures Act, we estimate receiving approximately 150 discretionary CoC requests annually. We estimate that approximately 150 sponsors, sponsor-investigators, or authorized representatives will submit requests. Preparing and sending each request would take approximately 2 hours

Dated: June 9, 2020.

Lowell J. Schiller,
Principal Associate Commissioner for Policy.
[FR Doc. 2020-13081 Filed 6-16-20; 8:45 am]

BILLING CODE 4164-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Renewal of the President's Council on Sports, Fitness and Nutrition's Charter

AGENCY: Office of Disease Prevention and Health Promotion, Office of the Assistant Secretary for Health, Office of the Secretary, Department of Health and Human Services.

ACTION: Notice.

SUMMARY: The U.S. Department of Health and Human Services (HHS) is hereby giving notice that the charter for the President's Council on Sports, Fitness, and Nutrition (hereafter referred to as the Council) has been renewed.

FOR FURTHER INFORMATION CONTACT:

Jennifer Anne Bishop, ScD, MPH, Designated Federal Official, U.S. Department of Health and Human Services, Office of Disease Prevention and Health Promotion; 1101 Wootton Parkway, Suite 420, Rockville, Maryland 20852; telephone: (240) 276-9567; email address jennifer.bishop@hhs.gov. Additional information is available on the Council's website at: <https://www.hhs.gov/fitness/about-pcsfn/executive-order/index.html#charter>.

SUPPLEMENTARY INFORMATION: The Council is charged with advising the President, through the Secretary of Health and Human Services (Secretary), concerning progress made in carrying out the provisions of Executive Order 13265, as amended by Executive Order 13824, which aims to expand and encourage youth sports participation and to promote the overall physical fitness, health, and nutrition of all Americans. The Council promotes this goal through external outreach, raising public awareness, and recommending to the President, through the Secretary, actions to accelerate such progress. Executive Order 13824 directs the Secretary to carry out the

responsibilities for public health and human services and to recognize the benefits of youth sports participation, physical activity, and a nutritious diet in helping create habits that support a healthy lifestyle. The Executive Order also directs the Secretary to develop a national strategy to expand children's participation in youth sports, encourage regular physical activity, including active play, and promote good nutrition for all Americans. Published in September 2019, the National Youth Sport Strategy (NYSS) focuses on children and youth in communities with below-average sports participation and communities with limited access to athletic facilities or recreational areas. The Council is tasked to provide additional support in promoting, shaping, and executing elements of the NYSS.

On May 13, 2020, the Secretary of Health and Human Services approved the renewal of the Council's charter. The new charter was executed and filed with the appropriate Congressional committees and the Library of Congress on May 17, 2020. The renewal of the Council's charter gives the Council authorization to operate until May 16, 2022.

A copy of the Council's charter is available on the Council's website at: <https://www.hhs.gov/fitness/about-pcsfn/executive-order/index.html#charter>.

A copy of the charter can also be obtained by accessing the Federal Advisory Committee Act Database that is managed by the Committee Management Secretariat under the General Services Administration. The website for the FACA database is <https://facadatabase.gov>.

Dated: June 10, 2020.

Carter Blakey,

Acting Director, Office of Disease Prevention and Health Promotion.

[FR Doc. 2020-12972 Filed 6-16-20; 8:45 am]

BILLING CODE 4150-28-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Cancer Institute; Notice of Closed Meetings

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended, notice is hereby given of meetings of the Board of Scientific Counselors for Basic Sciences, National Cancer Institute and the Board of Scientific Counselors for Clinical

Sciences and Epidemiology, National Cancer Institute.

The meetings will be closed to the public as indicated below in accordance with the provisions set forth in section 552b(c)(6), Title 5 U.S.C., as amended for the review, discussion, and evaluation of individual intramural programs and projects conducted by the National Cancer Institute, including consideration of personnel qualifications and performance, and the competence of individual investigators, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: Board of Scientific Counselors for Basic Sciences, National Cancer Institute.

Date: July 13, 2020.

Time: 10:00 a.m. to 11:45 a.m.

Agenda: To review and evaluate personnel qualifications and performance, and competence of individual investigators.

Place: National Institutes of Health, Shady Grove, 9609 Medical Center Drive, Rockville, MD 20850 (Virtual Meeting).

Contact Person: Mehrdad M. Tondravi, Ph.D., Chief, Institute Review Office, Office of the Director, National Cancer Institute, National Institutes of Health, 9609 Medical Center Drive, Room 3W302, Rockville, MD 20852, 240-276-5664, tondravim@mail.nih.gov.

Name of Committee: Board of Scientific Counselors for Clinical Sciences and Epidemiology, National Cancer Institute.

Date: July 13, 2020.

Time: 10:00 a.m. to 11:45 a.m.

Agenda: To review and evaluate personnel qualifications and performance, and competence of individual investigators.

Place: National Institutes of Health, Shady Grove, 9609 Medical Center Drive, Rockville, MD 20850 (Virtual Meeting).

Contact Person: Brian E. Wojcik, Ph.D., Executive Secretary, Institute Review Office, Office of the Director, National Cancer Institute, National Institutes of Health, 9609 Medical Center Drive, Room 3W414, Rockville, MD 20850, 240-276-5660, wojcikb@mail.nih.gov.

Name of Committee: Board of Scientific Counselors for Basic Sciences, National Cancer Institute.

Date: July 13, 2020.

Time: 11:45 a.m. to 2:10 p.m.

Agenda: To review and evaluate personnel qualifications and performance, and competence of individual investigators.

Place: National Institutes of Health, Shady Grove, 9609 Medical Center Drive, Rockville, MD 20850 (Virtual Meeting).

Contact Person: Mehrdad M. Tondravi, Ph.D., Chief, Institute Review Office, Office of the Director, National Cancer Institute, National Institutes of Health, 9609 Medical Center Drive, Room 3W302, Rockville, MD 20852, 240-276-5664, tondravim@mail.nih.gov.

(Catalogue of Federal Domestic Assistance Program Nos. 93.392, Cancer Construction; 93.393, Cancer Cause and Prevention