that the draft reports should be available on the website by June 5, 2019. Additional information will be posted when available or may be requested in hardcopy from Susan Blaine by phone: (703) 225–2471 or email: NTP-Meetings@icf.com. Individuals are encouraged to access the meeting web page to stay abreast of current information regarding the meeting.

Following the meeting, a report of the peer-review will be prepared and made available on the NTP website.

**Background Information on NTP Peer-Review Panels:** NTP panels are technical, scientific advisory bodies established on an “as needed” basis to provide independent scientific peer-review and advise NTP on agents of public health concern, new/revised toxicological test methods, or other issues. These panels help ensure transparent, unbiased, and scientifically rigorous input to the program for its use in making credible decisions about human hazard, setting research and testing priorities, and providing information to regulatory agencies about alternative methods for toxicity screening. NTP welcomes nominations of scientific experts for upcoming panels. Scientists interested in serving on an NTP panel should provide their current curriculum vitae to Susan Blaine by email: NTP-Meetings@icf.com.

The authority for NTP panels is provided by 42 U.S.C. 217a; section 222 of the Public Health Service Act, as amended. The panel is governed by the Federal Advisory Committee Act, as amended (5 U.S.C. Appendix 2), which sets forth standards for the formation and use of advisory committees.

This peer review is being conducted by a panel in attendance at NIEHS. Peer-review of future draft reports will be conducted in accordance with Department of Health and Human Services peer-review policies (https://aspe.hhs.gov/hhs-information-quality-peer-review) and Office of Management and Budget’s Final Information Quality Bulletin for Peer Review (70 FR 2664, January 4, 2005).

Dated: May 21, 2019.

Brian R. Berridge,
Associate Director, National Toxicology Program

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**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**National Institutes of Health**

Eunice Kennedy Shriver National Institute of Child Health & Human Development; Notice of Closed Meetings

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended, 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 552(b)(6) and 552(b)(4), 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:** National Institute of Child Health and Human Development Special Emphasis Panel; ZH01 DSR–A (50).
**Date:** July 12, 2019.
**Time:** 8:00 a.m. to 5:00 p.m.
**Agenda:** To review and evaluate grant applications.
**Place:** Residence Inn Bethesda Downtown, 7335 Wisconsin Ave., Bethesda, MD 20814.
**Contact Person:** Rita Anand, Ph.D., Scientific Review Officer, Division of Scientific Review, National Institute of Child Health and Human Development, NIH, 6710B Rockledge Drive, Bethesda, MD 20892, 301–496–1487, anandr@mail.nih.gov.

**Name of Committee:** National Institute of Child Health and Human Development Special Emphasis Panel; NIH Infant and Toddler Toolbox.
**Date:** July 25, 2019.
**Time:** 1:00 p.m. to 5:00 p.m.
**Agenda:** To review and evaluate contract proposals.
**Place:** National Institutes of Health, 6710B Rockledge Drive, Bethesda, MD 20817.
**Contact Person:** Sathasiva B. Kandasamy, Ph.D., Scientific Review Officer, Division of Scientific Review, National Institute of Child Health and Human Development, 6710B Rockledge Drive, Bethesda, MD 20892–9304, (301) 435–6680, skandas@nih.gov.

**Catalogue of Federal Domestic Assistance Program Nos.** 93.864, Population Research; 93.865, Research for Mothers and Children; 93.929, Center for Medical Rehabilitation Research; 93.209, Contraception and Infertility Loan Repayment Program, National Institutes of Health, HHS


Ronald J. Livingston, Jr.,
Program Analyst, Office of Federal Advisory Committee Policy.
In compliance with Section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the National Institutes of Health (NIH) has submitted to the Office of Management and Budget (OMB) a request for review and approval of the information collection listed below.

Proposed Collection: Center for Cancer Training (CCT) Application Form for electronic Individual Development Plan (eIDP), 0925-XXXX, Exp., Date XX/XXXX, NEW, National Cancer Institute (NCI), National Institutes of Health (NIH).

SUMMARY:

Training (CCT) supports NCI's goal of Institute's (NCI) Center for Cancer Testing for Federal Agencies Standards To Engage in Urine Drug Testing Facilities Which Meet Minimum Laboratories and Instrumented Initial Test Collection: The National Cancer Institute's (NCI) Center for Cancer Training (CCT) supports NCI's goal of training cancer researchers with various educational levels (postbaccalaureate, graduate students, postdoctoral fellows) and for varying periods of time (3 months to 5 years). The eIDP is an online, detailed questionnaire focused on responses to career and professional goals and expectations while the trainee works at the NCI. The eIDP ensures the NCI trainees are receiving proper career and professional guidance, making appropriate progress, and determining activities to achieve their goals. The eIDP is also used to track trainees' career and professional goals and to ensure trainees receive the tools needed to achieve those goals. It is expected the trainees will complete the eIDP annually and that the eIDP process could be improved by their responses. The effectiveness of training could also be enhanced by the reports received by the trainees completing the eIDP. Individual Development Plans have been collected by paper and pencil from trainees since 2001. With the implementation of the electronic system, a pilot of the eIDP was approved by OMB (#0925-0046) and implemented in December 2018. The pilot improved the clarity of the instructions for the eIDP system, and incorporated feedback from the trainees to improve the overall trainee IDP experience, which advances the effectiveness of training.

OMB approval is requested for 3 years. There are no costs to respondents other than their time. The total estimated annualized burden are 1,209 hours.

ESTIMATED ANNUALIZED BURDEN HOURS

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<th>Type of respondent</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average time per response (in hours)</th>
<th>Total annual burden hours</th>
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Patricia M. Busche, Project Clearance Liaison, National Cancer Institute, National Institutes of Health.

[FR Doc. 2019–11460 Filed 5–31–19; 8:45 am]
BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Substance Abuse and Mental Health Services Administration

Current List of HHS-Certified Laboratories and Instrumented Initial Testing Facilities Which Meet Minimum Standards To Engage in Urine Drug Testing for Federal Agencies

AGENCY: Substance Abuse and Mental Health Services Administration, HHS.

ACTION: Notice

SUMMARY: The Department of Health and Human Services (HHS) notifies federal agencies of the laboratories and Instrumented Initial Testing Facilities (IITF) currently certified to meet the standards of the Mandatory Guidelines for Federal Workplace Drug Testing Programs (Mandatory Guidelines). The Mandatory Guidelines were first published in the Federal Register on April 11, 1988 (53 FR 11970), and subsequently revised in the Federal Register on June 9, 1994 (59 FR 29908); September 30, 1997 (62 FR 51118); April 13, 2004 (69 FR 19644); November 25, 2008 (73 FR 71858); December 10, 2008 (73 FR 75122); April 30, 2010 (75 FR 22809); and on January 23, 2017 (82 FR 7920).

The Mandatory Guidelines were initially developed in accordance with Executive Order 12564 and section 503 of Public Law 100–71. The “Mandatory Guidelines for Federal Workplace Drug Testing Programs,” as amended in the revisions listed above, requires strict standards that laboratories and IITFs must meet in order to conduct drug and specimen validity tests on urine specimens for federal agencies.

To become certified, an applicant laboratory or IITF must undergo three rounds of performance testing plus an on-site inspection. To maintain that certification, a laboratory or IITF must participate in a quarterly performance testing program plus undergo periodic, on-site inspections.

Laboratories and IITFs in the applicant stage of certification are not to be considered as meeting the minimum requirements described in the HHS Mandatory Guidelines. A HHS-certified laboratory or IITF must have its letter of certification from HHS/SAMHSA (formerly: HHS/NIDA), which attests that it has met minimum standards.

In accordance with the Mandatory Guidelines dated January 23, 2017 (82 FR 7920), the following HHS-certified laboratories and IITFs meet the minimum standards to conduct drug