

recommendations on the use of the two *in vitro* NRU test methods, as well as recommended test method protocols, recommendations for future studies to further characterize the usefulness and limitations of *in vitro* methods for assessing acute systemic toxicity, recommended performance standards for tests with similar scientific principles and that measure or predict acute oral systemic toxicity, the peer panel report and **Federal Register** notices. The final BRD, which provides the supporting documentation for this report, is available as a separate document. The ICCVAM Test Method Evaluation Report and the supporting final BRD were forwarded to U.S. Federal agencies for their consideration for regulatory acceptance as required by the ICCVAM Authorization Act of 2000 (42 U.S.C. 285l-3). Agencies' responses to the test method recommendations will be posted on the ICCVAM/NICEATM Web site as they are received.

Background Information on ICCVAM, NICEATM, and SACATM

ICCVAM is an interagency committee composed of representatives from 15 Federal regulatory and research agencies that use, generate, or disseminate toxicological information. ICCVAM conducts technical evaluations of new, revised, and alternative methods with regulatory applicability and promotes the scientific validation and regulatory acceptance of toxicological test methods that more accurately assess the safety and hazards of chemicals and products and that refine, reduce, and replace animal use. The ICCVAM Authorization Act of 2000 established ICCVAM as a permanent interagency committee of the NIEHS under NICEATM. NICEATM administers ICCVAM and provides scientific and operational support for ICCVAM-related activities. NICEATM and ICCVAM work collaboratively to evaluate new and improved test methods applicable to the needs of U.S. Federal agencies. Additional

information about ICCVAM and NICEATM can be found on their Web site (<http://iccvam.niehs.nih.gov>).

SACATM was established January 9, 2002, and is composed of scientists from the public and private sectors (**Federal Register**, Vol. 67, No. 49, page 11358). SACATM provides advice to the Director of the NIEHS, to ICCVAM, and to NICEATM regarding the statutorily mandated duties of ICCVAM and activities of NICEATM. Additional information about SACATM, including the charter, roster, and records of past meetings, can be found at <http://ntp.niehs.nih.gov>/ see "Advisory Board & Committees" (or directly at <http://ntp.niehs.nih.gov/go/167>).

Dated: March 14, 2008.

Samuel H. Wilson,

Acting Director, National Institute of Environmental Health Sciences and National Toxicology Program.

[FR Doc. E8-5936 Filed 3-24-08; 8:45 am]

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FOR FURTHER INFORMATION CONTACT: <http://www.hhs.gov/healthit/ahic/confidentiality/>.

SUPPLEMENTARY INFORMATION: The Workgroup Members will continue discussing and evaluating the confidentiality, privacy, and security protections and requirements for participants in electronic health information exchange environments.

The meeting will be available via Web cast. For additional information, go to: http://www.hhs.gov/healthit/ahic/cps_instruct.html.

Dated: March 13, 2008.

Judith Sparrow,

Director, American Health Information Community, Office of Programs and Coordination, Office of the National Coordinator for Health Information Technology.

[FR Doc. E8-5853 Filed 3-24-08; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Office of the National Coordinator for Health Information Technology; American Health Information Community Confidentiality, Privacy, & Security Workgroup Meeting

ACTION: Announcement of meeting.

SUMMARY: This notice announces the 18th meeting of the American Health Information Community Confidentiality, Privacy, & Security Workgroup in accordance with the Federal Advisory Committee Act (Pub. L. No. 92-463, 5 U.S.C., App.)

DATES: April 17, 2008, from 1 p.m. to 5 p.m. (Eastern Time).

ADDRESSES: Mary C. Switzer Building (330 C Street, SW., Washington, DC 20201), Conference Room 4090 (please bring photo ID for entry to a Federal building).

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Proposed Information Collection Activity; Comment Request

Proposed Projects

Title: Grants to States for Access and Visitation: State Child Access Program Survey.

OMB No.: 0970-0204.

Description: On an annual basis, States must provide OCSE with data on programs that the Grants to States for Access and Visitation Program has funded. These program reporting requirements include, but are not limited to, the collection of data on the number of parents served, types of services delivered, program outcomes, client socio-economic data, referral sources, and other relevant data. OCSE is proposing revisions to the current survey.

ANNUAL BURDEN ESTIMATES

Instrument	No. of respondents	No. of responses per respondent	Average burden hours per response	Total burden hours
State Child Access Program Survey	314	1	15	4,710

Estimated Total Annual Burden Hours: 4,710.

In compliance with the requirements of Section 506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Administration for Children and

Families is soliciting public comment on the specific aspects of the information collection described above. Copies of the proposed collection of information can be obtained and comments may be forwarded by writing

to the Administration for Children and Families, Office of Administration, Office of Information Services, 370 L'Enfant Promenade, SW., Washington, DC 20447, Attn: Office of Administration, Office of Information

Services, 370 L'Enfant Promenade, SW., Washington, DC 20447, Attn: ACE Reports Clearance Officer. E-mail address: infocollection@acf.hhs.gov. All requests should be identified by the title of the information collection.

The Department specifically requests comments on: (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency's estimate of the burden of the proposed collection of information; (c) 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.

Consideration will be given to comments and suggestions submitted within 60 days of this publication.

Dated: March 19, 2008.

Janean Chambers,

Reports Clearance Officer.

[FR Doc. E8-5951 Filed 3-24-08; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Proposed Information Collection Activity; Comment Request

Proposed Projects

Title: Temporary Assistance for Needy Families (TANF) State Plan; Guidance.

OMB No.: 0970-0145.

Description: The State plan is a mandatory statement submitted to the Secretary of the Department of Health and Human Services by the State. It consists of an outline of how the State's TANF program will be administered and operated and certain required certifications by the State's Chief Executive Officer. Its submittal triggers the State's family assistance grant funding and it is used to provide the public with information about the program. If a State makes changes in its program, it must submit a State plan amendment.

Respondents: The 50 States, the District of Columbia, Guam, Puerto Rico and the Virgin Islands.

ANNUAL BURDEN ESTIMATES

Instrument	No. of respondents	No. of responses per respondent	Average burden hours per response	Total burden hours
Temporary Assistance for Needy Families (TANF) State Plan Guidance	54	0.5	33	891

Estimated Total Annual Burden Hours: 891.

In compliance with the requirements of Section 506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Administration for Children and Families is soliciting public comment on the specific aspects of the information collection described above. Copies of the proposed collection of information can be obtained and comments may be forwarded by writing to the Administration for Children and Families, Office of Administration, Office of Information Services, 370 L'Enfant Promenade, SW., Washington, DC 20447, Attn: ACF Reports Clearance Officer. E-mail address: infocollection@acf.hhs.gov. All requests should be identified by the title of the information collection.

The Department specifically requests comments on: (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency's estimate of the burden of the proposed collection of information; (c) 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.

Consideration will be given to comments and suggestions submitted within 60 days of this publication.

Dated: March 19, 2008.

Janean Chambers,

Reports Clearance Officer.

[FR Doc. E8-5952 Filed 3-24-08; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2008-D-0149] (formerly Docket No. 2007D-0031)

Global Harmonization Task Force, Study Group 4; Final Document; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of a final document that has been prepared by Study Group 4 of the Global Harmonization Task Force (GHTF). This document represents a harmonized proposal and recommendation from Study Group 4 of the GHTF that may be used by governments developing and updating their regulatory requirements for medical devices. This document is

intended to provide information only and does not describe current regulatory requirements; elements of this document may not be consistent with current U.S. regulatory requirements.

DATES: Submit written or electronic comments on this guidance at any time. General comments on agency guidance documents are welcome at any time.

ADDRESSES: Submit written requests for single copies of the guidance document to the Division of Small Manufacturers, International, and Consumer Assistance (HFZ-220), Center for Devices and Radiological Health, Food and Drug Administration, 1350 Piccard Dr., Rockville, MD 20850. Send one self-addressed adhesive label to assist that office in processing your request, or fax your request to 240-276-3151. See the **SUPPLEMENTARY INFORMATION** section for information on electronic access to the guidance. Submit written comments concerning this document to the Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments to <http://www.regulations.gov>. Identify comments with the docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: Jan Welch, GHTF, Study Group 4, Office of Compliance, Center for Devices and Radiological Health (HFZ-320), Food