

AboutAdvisoryCommittees/ucm111462.htm for procedures on public conduct during advisory committee meetings.

Notice of this meeting is given under the Federal Advisory Committee Act (5 U.S.C. app. 2).

Dated: October 3, 2012.

Jill Hartzler Warner,
Acting Associate Commissioner for Special Medical Programs.

[FR Doc. 2012-24861 Filed 10-9-12; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Services Administration

Agency Information Collection Activities: Proposed Collection; Comment Request

Periodically, the Health Resources and Services Administration (HRSA) publishes abstracts of information collection requests under review by the

Office of Management and Budget (OMB), in compliance with the Paperwork Reduction Act of 1995 (44 U.S.C. Chapter 35). To request a copy of the clearance requests submitted to OMB for review, email *paperwork@hrsa.gov* or call the HRSA Reports Clearance Office on (301) 443-1984.

The following request has been submitted to the Office of Management and Budget for review under the Paperwork Reduction Act of 1995:

Proposed Project: Maternal and Child Health Bureau Performance Measures for Discretionary Grants (OMB No. 0915-0298): Revision

The Maternal and Child Health Bureau (MCHB) intends to continue to collect performance data for Special Projects of Regional and National Significance (SPRANS), Community Integrated Service Systems (CISS), and other grant programs administered by MCHB.

The Health Resources and Services Administration (HRSA) proposes to continue using reporting requirements

for SPRANS projects, CISS projects, and other grant programs administered by MCHB, including national performance measures previously approved by OMB, and in accordance with the “Government Performance and Results Act (GPRA) of 1993” (Pub. L. 103-62). This Act requires the establishment of measurable goals for federal programs that can be reported as part of the budgetary process, thus linking funding decisions with performance. Performance measures for MCHB discretionary grants were initially approved in January 2003. Approval from OMB is being sought to continue the use of these measures. Some of these measures are specific to certain types of programs and will not apply to all grantees. Through the experience of utilizing these measures, we are enhancing them to better reflect program goals. Specifically, additional outcome measures that can be utilized by grantees that predominantly provide infrastructure services are being developed for submission to OMB.

The estimated response burden is as follows:

Form	Number of respondents	Responses per respondent	Total responses	Burden hours per response	Total burden hours
Grant Report	900	1	900	41	36,900

Written comments and recommendations concerning the proposed information collection should be sent within 30 days of this notice to the desk officer for HRSA, either by email to *OIRA_submission@omb.eop.gov* or by fax to (202) 395-5806. Please direct all correspondence to the “attention of the desk officer for HRSA.”

Dated: October 3, 2012.

Bahar Niakan,
Director, Division of Policy and Information Coordination.

[FR Doc. 2012-24889 Filed 10-9-12; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Interagency Coordinating Committee on the Validation of Alternative Methods Evaluation Report and Recommendations for Identifying Chemical Eye Hazards With Fewer Animals; Availability of Report; Notice of Transmittal to Federal Agencies

SUMMARY: The NTP Interagency Center for the Evaluation of Alternative Toxicological Methods (NICEATM) announces availability of an Interagency Coordinating Committee on the Validation of Alternative Methods (ICCVAM) test method evaluation report (TMER) that provides recommendations for identifying chemical eye hazards with fewer animals.

ICCVAM concludes that using a classification criterion of one or more positive animals in a three-animal test to identify chemicals and products that are eye hazards will maintain hazard classification equivalent to that provided by current testing procedures, while using up to 50% to 83% fewer animals. ICCVAM recommends

consideration of this classification criterion together with eye safety testing procedures that use a maximum of three animals per test substance. This recommendation also harmonizes the number of animals used for eye safety testing across U.S. regulatory agencies and international test guidelines.

The report and recommendations have been transmitted to Federal agencies for their review and response to ICCVAM.

FOR FURTHER INFORMATION CONTACT: Dr. William S. Stokes, Director, NICEATM, National Institute of Environmental Health Sciences (NIEHS), P.O. Box 12233, Mail Stop: K2-16, Research Triangle Park, NC 27709. Phone: 919-541-2384, Fax: 919-541-0947, Email: *niceatm@niehs.nih.gov*. Hand Deliver/Courier address: NICEATM, NIEHS, Room 2034, 530 Davis Drive, Morrisville, NC 27560.

SUPPLEMENTARY INFORMATION:

Background: Eye safety testing procedures vary among U.S. agencies. Current testing procedures specified in the U.S. Code of Federal Regulations (16 CFR 1500.42) provide criteria and procedures for identifying eye hazards based on rabbit eye test results (CPSC,