ADDITIONAL INFORMATION: Copies are available from the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852, 301–827–6860.

FOR FURTHER INFORMATION CONTACT: Teresa L. Hays, Advisory Committee and Oversight Management Staff, Food and Drug Administration, 10903 New Hampshire Ave., Silver Spring, MD 20993, 301–796–8220.

SUPPLEMENTARY INFORMATION: Under section 10(d) of the Federal Advisory Committee Act (5 U.S.C. app.) and 21 CFR 14.60(d), FDA has filed with the Library of Congress the annual reports for the following FDA advisory committees that held closed meetings during the period October 1, 2010 through September 30, 2011:

Center for Biologics Evaluation and Research
Allergenic Products Advisory Committee
Blood Products Advisory Committee
Cellular, Tissue and Gene Therapies Advisory Committee
Vaccines and Related Biological Products Advisory Committee

Center for Drug Evaluation and Research
Cardiovascular and Renal Drugs Advisory Committee
Gastrointestinal Drug Advisory Committee

National Center for Toxicological Research
Science Board to the National Center for Toxicological Research

Center for Tobacco Products
Tobacco Products Scientific Advisory Committee

Annual reports are available for public inspections between 9 a.m. and 4 p.m., Monday through Friday.
- The Library of Congress, Madison Bldg., Newspaper and Current Periodical Reading Room, 101 Independence Ave. SE., Rm. 133, Washington, DC; and
- The Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.


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

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Proposed Collection: Comment Request Post-Award Reporting Requirements Including New Research Performance Progress Report Collection

SUMMARY: In compliance with the requirement of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, for opportunity for public comment on proposed data collection projects, the Office of the Director, National Institutes of Health (NIH) will publish periodic summaries of proposed projects to be submitted to the Office of Management and Budget (OMB) for review and approval.

Proposed Collection: Title: Public Health Service (PHS) Post-award Reporting Requirements. Type of Information Collection Request: Revision. This collection represents a consolidation of post-award reporting requirements under the PRA, and includes the new Research Performance Progress Report (RPPR). Need and Use of Information Collection: The RPPR will replace existing interim performance reports used by all NIH, Food and Drug Administration, Centers for Disease Control and Prevention, and Agency for Healthcare Research and Quality (AHRQ) grantees. Interim progress reports are required to continue support of a PHS grant for each budget year within a competitive segment. The phased transition to the RPPR requires the maintenance of dual reporting processes for a period of time. Thus this information collection is for the new use of the RPPR, and continued use of the PHS Non-Competing Continuation Progress Report (PHS 2590, currently approved under 0925–0001), and the NIH AHRQ Ruth L. Kirschstein National Research Service Award (NRSA) Individual Fellowship Progress Report for Continuation Support (PHS 416–9, currently approved under 0925–0002). Only one interim progress report (RPPR or PHS2590/416–9) will be utilized for any given award. This collection also includes other PHS post-award reporting requirements: PHS 416–7 NRSA Termination Notice, PHS 2271 Statement of Appointment, 6031–1 NRSA Annual Payback Activities Certification, (currently approved under 0925–0002, expiration 6/30/2012); and HHS 568 Final Invention Statement and Certification, Final Progress Report instructions, and iEdison, and PHS 3734 Statement Relinquishing Interests and Rights in a PHS Research Grant