must be resolved through the Disputes clause. Any audits requested by the commercial supplier or licensor will be performed at supplier or licensor's expense.

This class deviation will apply to all new awards for GSA acquisitions for commercial supplies or services. Existing contracts will be required to incorporate the new terms whenever an option period is exercised or the contract is otherwise modified.

This effort will reduce risk by uniformly addressing common unacceptable Commercial Supplier Agreement terms, facilitate efficiency and effectiveness in the contracting process by reducing the administrative burden for the Government and industry, and promote competition by reducing barriers to industry, particularly small businesses.

Dated: March 17, 2015.

#### Jeffrey A. Koses,

Senior Procurement Executive, Office of Acquisition Policy, Office of Governmentwide Policy.

[FR Doc. 2015-06422 Filed 3-19-15; 8:45 am]

BILLING CODE 6820-61-P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

#### **National Institutes of Health**

Submission for OMB Review; 30-Day Comment Request: The Genetic Testing Registry

**SUMMARY:** Under the provisions of 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. This proposed information collection was previously published in the Federal Register on November 25, 2014 (79 FR 70194), and allowed 60days for public comment. No public comments were received. The purpose of this notice is to allow an additional 30 days for public comment. The Office of the Director (OD), National Institutes of Health, may not conduct or sponsor, and the respondent is not required to respond to, an information collection that has been extended, revised, or implemented on or after October 1, 1995, unless it displays a currently valid OMB control number.

Direct Comments to OMB: Written comments and/or suggestions regarding the item(s) contained in this notice, especially regarding the estimated public burden and associated response time, should be directed to the: Office of Management and Budget, Office of Regulatory Affairs, OIRA\_submission@omb.eop.gov or by fax to 202–395–6974, Attention: NIH Desk Officer.

**DATES:** Comment Due Date: Comments regarding this information collection are best assured of having their full effect if received within 30-days of the date of this publication.

**FOR FURTHER INFORMATION CONTACT:** To obtain a copy of the data collection plans and instruments or request more information on the proposed project contact: Ms. Sarah Carr, Acting Director,

Office of Clinical Research and Bioethics Policy, Office of Science Policy, NIH, 6705 Rockledge Dr., Suite 750, Bethesda, MD 20892, or call nontoll-free number (301) 496–9838, or Email your request, including your address to: OCRBP-OSP@od.nih.gov. Formal requests for additional plans and instruments must be requested in writing.

## Proposed Collection: The Genetic Testing Registry, 0925–0651, Reinstatment Without Change,—Office of the Director (OD), National Institutes of Health (NIH)

Need and Use of Information
Collection: Clinical laboratory tests are
available for more than 5,000 genetic
conditions. The Genetic Testing Registry
(GTR) provides a centralized, online
location for test developers,
manufacturers, and researchers to
voluntarily submit detailed information
about the availability and scientific
basis of their genetic tests. The GTR is
of value to clinicians by providing
information about the accuracy,
validity, and usefulness of genetic tests.
The GTR also highlights evidence gaps
where additional research is needed.

OMB approval is requested for 3 years. There are no costs to respondents other than their time. The total estimated annualized burden hours are 5,536.

### **ESTIMATED ANNUALIZED BURDEN HOURS**

Type of respondent	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total annual burden hours
Laboratory Personnel Using Bulk Submission.	Minimal Fields	190	29	18/60	1,653
Laboratory Personnel Not Using	Optional Fields	159 116	29 29	14/60 30/60	1,076 1,682
Bulk Submission.	Optional Fields	97	29	24/60	1,125

Dated: March 13, 2015. Lawrence A. Tabak,

Deputy Director, National Institutes of Health. [FR Doc. 2015–06370 Filed 3–19–15; 8:45 am]

BILLING CODE 4140-01-P