• Designate an employee or employees to coordinate and be accountable for the information security program;

• Identify material internal and external risks to the security, confidentiality, and integrity of personal information that could result in the unauthorized disclosure, misuse, loss, alteration, destruction, or other compromise of such information, and assess the sufficiency of any safeguards in place to control these risks;

• Design and implement reasonable safeguards to control the risks identified through risk assessment, and regularly test or monitor the effectiveness of the safeguards' key controls, systems, and

procedures:

• Develop and use reasonable steps to select and retain service providers capable of appropriately safeguarding personal information they receive from Lookout, and require service providers by contract to implement and maintain appropriate safeguards; and

• Evaluate and adjust its information security programs in light of the results of testing and monitoring, any material changes to operations or business arrangements, or any other circumstances that it knows or has reason to know may have a material impact on its information security

program

protected.

Part III of the proposed order requires Lookout to obtain within the first one hundred eighty (180) days after service of the order, and on a biennial basis thereafter for a period of twenty (20) years, an assessment and report from a qualified, objective, independent thirdparty professional, certifying, among other things, that: (1) It has in place a security program that provides protections that meet or exceed the protections required by Part II of the proposed order; and (2) its security program is operating with sufficient effectiveness to provide reasonable assurance that the security, confidentiality, and integrity of sensitive consumer, employee, and job applicant information has been

Parts IV through VIII of the proposed order are reporting and compliance provisions. Part IV requires Lookout to retain documents relating to its compliance with the order. For most records, the order requires that the documents be retained for a five-year period. For the third-party assessments and supporting documents, Lookout must retain the documents for a period of three years after the date that each assessment is prepared. Part V requires dissemination of the order now and in the future to all current and future

subsidiaries, current and future principals, officers, directors, and managers, and to persons with responsibilities relating to the subject matter of the order. Part VI ensures notification to the FTC of changes in corporate status.

Part VII mandates that Lookout submit a compliance report to the FTC within 60 days, and periodically thereafter as requested. Part VIII is a provision "sunsetting" the order after twenty (20) years, with certain exceptions.

The purpose of this analysis is to facilitate public comment on the proposed order. It is not intended to constitute an official interpretation of the proposed order or to modify its terms in any way.

By direction of the Commission.

#### Donald S. Clark,

Secretary.

[FR Doc. 2011–11182 Filed 5–9–11; 8:45 am] BILLING CODE 6750–01–P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Submission for OMB Review; Comment Request

Request; OMB No. 0925–0177 "Special Volunteer and Guest Researcher Assignment," Form 590

**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 August 25, 2010, page 52351 and allowed 60 days for public comment. No public comments were received. The purpose of this notice is to allow an additional 30 days for public comment. The 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 July 31, 2005, unless it displays a currently valid OMB control number.

Proposed Collection: Title: Special Volunteer and Guest Researcher Assignment for use in NIH facilities. Type of Information Collection Request: Reinstatement, 0MB 0925–0177, Expiration Date July 31, 2005. Need and Use of Information Collection Request: Form Number: NIH–590. A single Form NIH–590 is completed by an NIH official for each Guest Researcher or

Special Volunteer prior to his/her arrival at NIH. The information on the form is necessary for the approving official to reach a decision on whether to allow a Guest Researcher to use NIH facilities, or whether to accept volunteer services offered by a Special Volunteer. If the original assignment is extended, another form notating the extension is completed to update the file. Frequency of Response: once. Affected Public: Individuals. Type of Respondents: Nonfederal scientific professionals and/or individuals. The annual Reporting burden is as follows: Estimated Number of Respondents: 1660; Estimated Number of Responses per Respondent: 1.0; Average Burden Hours Per Response: 0.1; and Estimated Total Annual Burden Hours Requested: 166. The estimated annualized cost to respondents is \$2,275. There are no Capital Costs to report. There are no Operating or Maintenance Costs to report.

Request for Comments: Written comments and/or suggestions from the public and affected agencies are invited on one or more of the following points: (1) Whether the proposed collection of information is necessary for the proper performance of the function of the agency, including whether the information will have practical utility; (2) The accuracy of the agency's estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) Ways to enhance the quality, utility, and clarity of the information to be collected; and (4) Ways to minimize the burden of the collection of information on those who are to respond, including the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology.

### FOR FURTHER INFORMATION CONTACT:

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: Desk Officer for NIH. To request more information on the proposed project or to obtain a copy of the data collection plans and instruments, contact Mrs. Wanda Darwin, Office of Human Resources, Office of The Director, NIH, Building 31, Room 1C31E, One Center Drive, Bethesda, MD 20892-2269, or call non-toll-free number 301-4022820, or e-mail your request, including your address, to: [darwinw@od.nih.gov].

Comments 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.

Dated: May 4, 2011.

#### Wanda R. Darwin,

Human Resources Specialist, Office of Human Resources, National Institutes of Health.

[FR Doc. 2011-11406 Filed 5-9-11; 8:45 am]

BILLING CODE 4140-01-P

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

Solicitation of Written Comments on the Draft Report and Draft Recommendations of the Vaccine Safety Working Group for Consideration by the National Vaccine Advisory Committee on the Federal Vaccine Safety System

**AGENCY:** National Vaccine Program Office, Office of the Assistant Secretary for Health, Office of the Secretary, Department of Health and Human Services.

**ACTION:** Notice.

**SUMMARY:** The National Vaccine Advisory Committee (NVAC) was established in 1987 to comply with Title XXI of the Public Health Service Act (Pub. L. 99-660) (Section 2105) (42 U.S. Code 300aa-5 (PDF-78 KB)). Its purpose is to advise and make recommendations to the Director of the National Vaccine Program on matters related to program responsibilities. The Assistant Secretary for Health (ASH) has been designated by the Secretary of Health and Human Services as the Director of the National Vaccine Program. The ASH has charged the NVAC "To review the current federal vaccine safety system and develop a White Paper describing the infrastructure needs for a federal vaccine safety system to fully characterize the safety profile of vaccines in a timely manner, reduce adverse events whenever possible, and maintain and improve public confidence in vaccine safety." On behalf of the NVAC, the Vaccine Safety Working Group (VSWG) has developed a draft report and draft recommendations for the consideration by the NVAC in developing the NVAC's final recommendations to the ASH. The National Vaccine Program Office (NVPO) is soliciting public comment on the National Vaccine Advisory Committee (NVAC) Vaccine Safety

Working Group draft report and draft recommendations for the federal vaccine safety system to be considered by the NVAC. Individuals and organizations are encouraged to submit their comments on the draft report and draft recommendations. It is anticipated that the draft report and draft recommendations, as revised with consideration given to public comment and stakeholder input, will be presented in mid to late 2011 to the NVAC for deliberation and decision on their final recommendations.

**DATES:** To receive consideration comments should be received no later than 5 p.m. EST on June 6, 2011.

**ADDRESSES:** 1. The draft report and draft recommendations are available on the Web at http://www.hhs.gov/nvpo/nvac/subgroups/vaccinesafety.html.

2. Electronic responses are preferred and may be addressed to *vaccinesafetvRFI@hhs.gov*.

3. Written responses should be addressed to: National Vaccine Program Office, U.S. Department of Health and Human Services, 200 Independence Avenue, SW., Room 739G.5, Washington, DC 20201, Attention: Vaccine Safety c/o Kristin Goddard.

FOR FURTHER INFORMATION CONTACT:
Kristin Goddard, National Vaccine
Program Office, Department of Health
and Human Services, Hubert H.
Humphrey Building, 200 Independence
Avenue, SW., Room 439G.5,
Washington, DC 20201, Attn: NVAC
Vaccine Safety Working Group,
telephone (202) 205–5317; fax 202–260–
1165; e-mail vaccinesafetyRFI@hhs.gov.
SUPPLEMENTARY INFORMATION:

### I. Background

The National Vaccine Program Office (NVPO) is located within the Office of the Assistant Secretary for Health (OASH), Office of the Secretary, Department of Health and Human Services and has the responsibility for coordinating and fostering collaborations among the many Federal agencies involved in vaccine and immunization activities. NVPO also has responsibility for the management of the National Vaccine Advisory Committee, a chartered federal advisory committee that reports to the Assistant Secretary for Health in his role as the Director of the National Vaccine Program (NVP).

Recognizing the importance of vaccine safety in the NVP, the ASH charged NVAC to "review the current federal vaccine safety system and develop a White Paper describing the infrastructure needs for a federal vaccine safety system to fully characterize the safety profile of

vaccines in a timely manner, reduce adverse events whenever possible, and maintain and improve public confidence in vaccine safety." On behalf of the NVAC the Vaccine Safety Working Group (VSWG) has developed a draft report and draft recommendations for the consideration by the NVAC in developing the NVAC's final recommendations to the ASH. The VSWG membership represents a broad range of expertise including pediatric and adult infectious diseases, genomics, immunology, epidemiology, public health, maternal and child health, pharmacoepidemiology, and biostatistics. Through review of previous recommendations on improvement to the vaccine safety system, input from an array of experts and stakeholders, and identification of gaps in the current federal vaccine safety system the VSWG developed draft recommendations for the consideration of the NVAC to achieve the charge as noted above.

The draft report describes relative benefits and risks of vaccines, current vaccine coverage levels, successes and challenges of the current system, methodology for VSWG recommendation development, and the conclusions of the VSWG from these findings. From these conclusions the VSWG has developed draft recommendations in eight categories: Leadership, Coordination, Research, Post Licensure Surveillance, Clinical Practice, Communications, Stakeholder and Public Engagement, and Assurance and Accountability.

Through this request for comment HHS is seeking comments from everyone, including stakeholders and the broad public, on the NVAC Vaccine Safety Working Group draft report and draft recommendations to be submitted to the NVAC for consideration in their final recommendations to the ASH. Comments received will be available for public viewing on the NVAC Vaccine Safety Working Group section of the NVPO Web site (http://www.hhs.gov/nvpo/nvac/subgroups/vaccinesafety.html).

### **II. Request for Comment**

NVPO, on behalf of the NVAC Vaccine Safety Working Group, requests input on the draft report and draft recommendations. (http://www.hhs.gov/nvpo/nvac/subgroups/vaccinesafety.html). In addition to general comments, NVPO is seeking input on any additional gaps not addressed in the NVAC Vaccine Safety Working Group draft report, and/or prioritization criteria and its application