analyzing the redacted credit reports and related scores of the nonrespondents, we obtain a final check on the degree to which the enhanced procedures were effective in achieving a nationally representative sample of credit reports.

## 2. Estimated Hours Burden

Consumer participation in the proposed national study would involve an initial preparation for the in-depth interview and time spent by participants to understand, review, and if deemed necessary, dispute information in their credit reports. Invitation letters will be sent in progressive waves in order to obtain approximately 1,000 participants. The individuals who receive these letters are drawn from the SC list discussed above and will be asked to go directly to a designated Web site for enrollment if they wish to participate; registration is expected to take at most 15 minutes per participant. 19 The registration process thus comes to approximately 250 hours (reckoned at 1/ 4 hour for each of 1,000 consumers).

For the purpose of calculating burden under the PRA regarding the review process of the credit reports, FTC staff submits the following estimates that are based on the contractor's experience with the second pilot study. Some participants prepare thoroughly in advance of the in-depth interview of their credit reports. In such situations, even complicated reports may generally be finished under 30 minutes. Other consumers may not find time for significant preparation in advance of the in-depth review, and in such cases the interview could take up to an hour. The participants in the second pilot study reported taking an average of 69 minutes (median 53 minutes) to prepare for the interview, with 90% taking between 10 and 180 minutes. The interviews themselves took an average of 19 minutes (median 15 minutes) with 90% taking between 5 and 45 minutes. Overall, the average combined time for preparation and the interview was about

90 minutes (1.5 hours). For a national study involving 1,000 consumers, FTC staff thus estimates the burden hours for the review process to be approximately 1,500 hours (1,000 consumers x 1.5 hours). Further adding on the time spent for the registration process (0.25 hours per participant), the total burden hours come to approximately 1,750 hours.

### 3. Estimated Cost Burden

The cost per consumer for their participation should be negligible. Participation is voluntary and it will not require any start-up or capital expenditure. There is no labor time expenditure beyond the 1.75 hours per consumer estimated above. Participants may receive an honorarium to compensate them for their time. The amount will be determined by FTC staff in consultation with the contractor according to an analysis of customary procedures and a consideration of response rates within key categories, such as, response rates for consumers with impaired credit. As with the pilot studies, participants will not pay for their credit reports or credit scores.

#### Willard Tom,

General Counsel
[FR Doc. E9–17147 Filed 7–17–09: 8:45 am]
BILLING CODE: 6750 –01–S

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

## **National Institutes of Health**

## Submission for OMB Review; Comment Request; CareerTrac

Summary: Under the provisions of section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the Fogarty International Center (FIC) and National Institute of Environmental Health Sciences (NIEHS), 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 May 12, 2009, page 22172, and allowed 60-days for public comment. No comments were received from this notification regarding the cost and hour burden estimates. The purpose of this announcement 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 October 1, 1995, unless it displays a currently valid OMB control number.

Proposed Collection: Title: CareerTrac. Type of Information Collection Request: Revision (OMB No.: 0925-0568 Expiration: Aug. 31, 2009). Need and Use of Information Collection: This data collection system is being developed to track, evaluate and report short and long-term outputs, outcomes and impacts of international trainees involved in health research training programs—specifically tracking this for at least ten years following training by having Principal Investigators enter data after trainees have completed the program. The data collection system provides a streamlined, Web-based application permitting principal investigators to record career achievement progress by trainee on a voluntary basis. FIC and NIEHS management will use this data to monitor, evaluate and adjust grants to ensure desired outcomes are achieved, comply with OMB part requirements, respond to congressional inquiries, and as a guide to inform future strategic and management decisions regarding the grant program.

Frequency of Response: Annual and periodic Affected Public: none Type of Respondents: Principal Investigators and/or their administrators funded by FIC and NIEHS. The annual reporting burden is as follows: Estimated Number of Respondents: 275; Estimated Number of Responses per Respondent: 1; Average Burden Hours per Response 7.5 and Estimated Total Annual Burden Hours Requested: 2063. The annualized cost to respondents is estimated at \$82,500. 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.

above, the rescoring of the frozen files will then provide the impact of any confirmed errors on the participants' credit scores.

<sup>&</sup>lt;sup>19</sup> At the registration Web site, a person may take the time to read several disclosures, including a privacy disclosure and an outline of the various steps of the study that every participant agrees to undertake. The consumer is then asked to enter basic contact information (e.g., name, address, telephone number, best time to be contacted further about the study) and to enter an electronic signature certifying the consumer's consent to participate in the study. For those who may not have Internet access to register, the contractor would also have a procedure to mail the appropriate disclosures and study steps to the respondent and then receive back enrolment information and the consumer's signed consent in paper form.

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: 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 Dr. Linda Kupfer, Fogarty International Center, National Institutes of Health, 16 Center Drive, Building 16, Bethesda, MD 20892–6705 or call non-toll-free number 301–496–3288 or e-mail your request, including your address to kupferl@mail.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: July 13, 2009.

#### Timothy J. Tosten,

Executive Officer, Fogarty International Center, National Institutes of Health. [FR Doc. E9–17214 Filed 7–17–09; 8:45 am] BILLING CODE 4140–01–P

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Food and Drug Administration [Docket No. FDA-2008-N-0544]

Agency Information Collection Activities; Announcement of Office of Management and Budget Approval; Record Retention Requirements for the Soy Protein and Coronary Heart Disease Health Claim

**AGENCY:** Food and Drug Administration,

**ACTION:** Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that a collection of information entitled "Record Retention Requirements for the Soy Protein and Coronary Heart Disease Health Claim" has been approved by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995.

## FOR FURTHER INFORMATION CONTACT:

Jonna Capezzuto, Office of Information Management (HFA–710), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301–796–3794.

**SUPPLEMENTARY INFORMATION:** In the **Federal Register** of January 14, 2009 (74 FR 2079), the agency announced that

the proposed information collection had been submitted to OMB for review and clearance under 44 U.S.C. 3507. An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number. OMB has now approved the information collection and has assigned OMB control number 0910–0428. The approval expires on May 31, 2012. A copy of the supporting statement for this information collection is available on the Internet at <a href="http://www.reginfo.gov/public/do/PRAMain">http://www.reginfo.gov/public/do/PRAMain</a>.

Dated: July 10, 2009.

### Jeffrey Shuren,

Associate Commissioner for Policy and Planning.

[FR Doc. E9–17094 Filed 7–17–09; 8:45 am]  $\tt BILLING$  CODE 4160–01–S

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

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

## National Center for Complementary and Alternative Medicine; Notice of Closed Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meeting.

The meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Center for Complementary and Alternative Medicine Special Emphasis Panel; Review Predoctoral and Postdoctoral Fellowship Applications. Date: July 28, 2009.

Time: 12 p.m. to 6 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, Two Democracy Plaza, 6707 Democracy Boulevard, Bethesda, MD 20892 (Telephone Conference Call).

Contact Person: Peter Kozel, PhD, Scientific Review Officer, NCCAM, 6707 Democracy Boulevard, Suite 401, Bethesda, MD 20892–5475. 301–496–8004. kozelp@mail.nih.gov.

This notice is being published less than 15 days prior to meeting due to scheduling conflicts.

(Catalogue of Federal Domestic Assistance Program Nos. 93.213, Research and Training in Complementary and Alternative Medicine, National Institutes of Health, HHS)

Dated: July 14, 2009.

### Jennifer Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. E9–17202 Filed 7–17–09; 8:45 am] **BILLING CODE 4140–01–P** 

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

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

## National Center for Complementary and Alternative Medicine; Notice of Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the National Advisory Council for Complementary and Alternative Medicine (NACCAM) meeting.

The meeting will be open to the public as indicated below, with attendance limited to space available. Individuals who plan to attend and need special assistance, such as sign language interpretation or other reasonable accommodations, should notify the Contact Person listed below in advance of the meeting.

A portion of the meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and/or contract proposals and the discussion could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications and/or contract proposals, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Advisory Council for Complementary and Alternative Medicine.

Date: September 11, 2009.
Closed: 8:30 a.m. to 10:30 a.m.
Agenda: To review and evaluate grant applications and/or proposals.
Open: 11 a.m. to 4 p.m.

Agenda: Opening remarks by the Director of the National Center for Complementary and Alternative Medicine, presentation of a new research initiative, and other business of the Council.

Place: National Institutes of Health, Neuroscience Building, 6001 Executive Boulevard, Conference Rooms C & D, Bethesda, MD 20892.

Contact Person: Martin H. Goldrosen, PhD., Executive Secretary, Director, Division of Extramural Activities, National Center for Complementary and Alternative Medicine,