Times and Dates: 10:00 a.m.–2:00 p.m., February 26, 2015.

The meeting will be webcast live via the World Wide Web; for instructions and more information on ACIP please visit the ACIP Web site: http:// www.cdc.gov/vaccines/acip/index.html.

Matters To Be Discussed: The shortened agenda will include discussions and votes on: Influenza LAIV use, serogroup B meningococcal (MenB) vaccines use in high risk groups, including outbreaks, and the use of 9vHPV vaccine.

Agenda items are subject to change as priorities dictate.

Contact Person for More Information: Stephanie B. Thomas, National Center for Immunization and Respiratory Diseases, CDC, 1600 Clifton Road NE., MS–A27, Atlanta, Georgia 30333, telephone: (404) 639–8836; Email ACIP@CDC.GOV.

The Director, Management Analysis and Services Office, has been delegated the authority to sign **Federal Register** notices pertaining to announcements of meetings and other committee management activities for both the Centers for Disease Control and

Prevention and the Agency for Toxic Substances and Disease Registry.

Catherine Ramadei,

Acting Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 2015–04330 Filed 3–2–15; 8:45 am] BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Comment Request

Title: Grant Reviewer Recruitment Form.

OMB No.: NEW.

Description: The Administration for Children and Families' Children's Bureau (CB) is responsible for administering the review of eligible grant applications submitted in response to funding opportunity announcements issued by CB. CB ensures that the objective review process is independent, efficient, effective, economical, and complies with the applicable statutes, regulations,

and policies. Applications are reviewed by subject experts knowledgeable in child welfare and related fields. Review findings are advisory to CB; CB is responsible for making award decisions.

This announcement is a request for approval of the proposed information collection system, the Reviewer Recruitment Module (RRM). CB will use a web-based data collection form and database to gather critical reviewer information in drop down menu format for data such as: degree, occupation, affiliations with organizations and institutions that serve special populations, and demographic information that may be voluntarily provided by a potential reviewer.

These data elements will help CB find and select expert grant reviewers for objective review committees. The webbased system will permit reviewers to access and update their information at will and as needed. The RRM will be accessible by the general public via https://rrm.grantsolutions.gov/AgencyPortal/cb.aspx.

Respondents: Generally, our reviewers are current or retired professionals with backgrounds in child welfare and related fields and in some instances current or former foster care parents or clients.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
Reviewer recruitment module	500	1	.25	125

Estimated Total Annual Burden Hours: 125.

Additional Information

Copies of the proposed collection may be obtained by writing to the Administration for Children and Families, Office of Planning, Research and Evaluation, 370 L'Enfant Promenade SW., Washington, DC 20447, Attn: ACF Reports Clearance Officer. All requests should be identified by the title of the information collection. Email address: infocollection@acf.hhs.gov.

OMB Comment

OMB is required to make a decision concerning the collection of information between 30 and 60 days after publication of this document in the **Federal Register**. Therefore, a comment is best assured of having its full effect if OMB receives it within 30 days of publication. Written comments and recommendations for the proposed information collection should be sent

directly to the following: Office of Management and Budget, Paperwork Reduction Project, Email: OIRA_ SUBMISSION@OMB.EOP.GOV, Attn: Desk Officer for the Administration for Children and Families.

Robert Sargis,

BILLING CODE 4184-01-P

Reports Clearance Officer. [FR Doc. 2015–04320 Filed 3–2–15; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Proposed Information Collection Activity; Comment Request

Proposed Projects

Title: ORR–2 Quarterly Report on Expenditures and Obligations.

OMB No.: 0970-0407.

Description: The Office of Refugee Resettlement (ORR) reimburses, to the extent of available appropriations, certain non-federal costs for the provision of cash and medical assistance to refugees, along with allowable expenses for the administration of the refugee resettlement program at the State level. States (and Wilson/Fish projects; i.e., alternative projects for the administration of the refugee resettlement program) currently submit the ORR-2 Quarterly Report on Expenditures and Obligations, which provides aggregate expenditure and obligation data. This proposed data collection collects expenditures and obligations data separately for each of the four CMA program components: Refugee cash assistance, refugee medical assistance, cash and medical assistance administration, and services for unaccompanied minors. This breakdown of financial status data allows ORR to track program

expenditures in greater detail to anticipate any funding issues and to meet the requirements of ORR regulations at CFR 400.211 to collect these data for use in estimating future costs of the refugee resettlement program. ORR must implement the methodology at CFR 400.211 each year after receipt of its annual appropriation to ensure that appropriated funds will be adequate for reimbursement to States

of the costs for assistance provided to entering refugees. The estimating methodology prescribed in the regulations requires the use of actual past costs by program component. In the event that the methodology indicates that appropriated funds are inadequate, ORR must take steps to reduce federal expenses, such as by limiting the number of months of eligibility for Refugee Cash Assistance and Refugee

Medical Assistance. This proposed single-page financial report allows ORR to collect the necessary data to ensure that funds are adequate for the projected need and thereby meet the requirements of both the Refugee Act and ORR regulations.

Respondents: State governments, Wilson/Fish Alternative Projects.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
ORR Financial Status Report	58	4	0.50	116

Estimated Total Annual Burden Hours: 116.

In compliance with the requirements of Section 506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Administration for Children and Families is soliciting public comment on the specific aspects of the information collection described above. Copies of the proposed collection of information can be obtained and comments may be forwarded by writing to the Administration for Children and Families, Office of Planning, Research and Evaluation, 370 L'Enfant Promenade SW., Washington, DC 20447, Attn: ACF Reports Clearance Officer. Email address: infocollection@ acf.hhs.gov. All requests should be identified by the title of the information collection.

The Department specifically requests comments on: (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency's estimate of the burden of the proposed collection of information; (c) the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology. Consideration will be given to comments and suggestions submitted within 60 days of this publication.

Robert Sargis,

Reports Clearance Officer. [FR Doc. 2015–04365 Filed 3–2–15; 8:45 am] BILLING CODE 4184–01–P DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2011-N-0535]

Agency Information Collection
Activities; Submission for Office of
Management and Budget Review;
Comment Request; Notification of a
Health Claim or Nutrient Content Claim
Based on an Authoritative Statement
of a Scientific Body

AGENCY: Food and Drug Administration, HHS

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that a proposed collection of information has been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995. DATES: Fax written comments on the collection of information by April 2,

ADDRESSES: To ensure that comments on the information collection are received, OMB recommends that written comments be faxed to the Office of Information and Regulatory Affairs, OMB, Attn: FDA Desk Officer, FAX: 202–395–7285, or emailed to oira_submission@omb.eop.gov. All comments should be identified with the OMB control number 0910–0374. Also include the FDA docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: FDA PRA Staff, Office of Operations, Food and Drug Administration, 8455 Colesville Rd., COLE–14526, Silver Spring, MD 20993–0002 PRAStaff@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: In compliance with 44 U.S.C. 3507, FDA has submitted the following proposed collection of information to OMB for review and clearance.

Notification of a Health Claim or Nutrient Content Claim Based on an Authoritative Statement of a Scientific Body—(OMB Control Number 0910– 0374)—Extension

Section 403(r)(2)(G) and (r)(3)(C) of the Federal Food, Drug and Cosmetic Act (the FD&C Act) (21 U.S.C. 343(r)(2)(G) and (r)(3)(C), as amended by the FDA Modernization Act of 1997, provides that any person may market a food product whose label bears a nutrient content claim or a health claim that is based on an authoritative statement of a scientific body of the U.S. Government or the National Academy of Sciences (NAS). Under this section of the FD&C Act, a person that intends to use such a claim must submit a notification of its intention to use the claim 120 days before it begins marketing the product bearing the claim. In the **Federal Register** of June 11, 1998 (63 FR 32102), we announced the availability of a guidance entitled "Guidance for Industry: Notification of a Health Claim or Nutrient Content Claim Based on an Authoritative Statement of a Scientific Body." The guidance provides the Agency's interpretation of terms central to the submission of a notification and the Agency's views on the information that should be included in the notification. We believe that the guidance will enable persons to meet the criteria for notifications that are established in section 403(r)(2)(G) and (r)(3)(C) of the FD&C Act. In addition to the information specifically required by the FD&C Act to be in such notifications, the guidance states that the notifications