Dated: June 14, 2007.

Robert Sargis,

Reports Clearance Officer.
[FR Doc. 07–3049 Filed 6–20–07; 8:45 am]
BILLING CODE 4184–01–M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Statement of Organization, Functions and Delegation of Authority

Notice is hereby given that I have delegated to the Principal Deputy Assistant Secretary, Deputy Assistant Secretaries, Program Directors, Program Commissioners, Deputy Director/ Commissioner, Office of Child Support Enforcement, and Staff Office Directors the following authority vested in me by the Secretary of Health and Human Services in the memorandum dated August 20, 1991, Delegations of Authority for Social Security Act Programs; 31 U.S.C. 1535; and HHS General Administrative Manual, Chapter 8–77.

(a) Authorities Delegated

- 1. Authority to administer approved cooperative research, experimental, pilot or demonstration projects under the provisions of sections 1110 and 1115 of the Social Security Act.
- 2. Authority to approve interagency agreements to procure, provide or exchange services, supplies or equipment.

(b) Limitations

- 1. The authority listed in #1 above shall be exercised under the condition that projects may be administered by the Office of Planning, Research and Evaluation (OPRE), by the program/staff office or jointly by OPRE with the program/staff office.
- 2. Where all or any part of an experimental, pilot, demonstration, or other project is wholly financed with Federal funds made available under sections 1110 or 1115 of the Social Security Act, without any State, local or other non-Federal financial participation, that project must be approved by the Secretary of Health and Human Services.
- 3. This delegation of authority does not include the authority to approve/ disapprove projects under section 1115 of the Social Security Act or approve/ disapprove waivers of State Plan requirements or costs that would not otherwise be included as expenditures under the provisions of section

1115(a)(1) and (2) of the Social Security Act.

4. The authority to approve interagency agreements to procure, provide, or exchange services, supplies, or equipment requires the concurrence of the ACF Chief Financial Officer if it exceeds \$250,000 (including amendments) within a fiscal year or if it requires the signature of the Assistant Secretary, ACF, or the Secretary of HHS.

(c) Effective Date

This delegation is effective upon the date of signature.

(d) Effect on Existing Delegations

As related to this delegation of authority, this delegation supersedes all previous delegations of authority involving the administration of the cross-program authorities delegated herein.

I hereby ratify and affirm any actions taken by the Principal Deputy Assistant Secretary, Deputy Assistant Secretaries, Program Directors, Program Commissioners, Deputy Director/Commissioner, Office of Child Support Enforcement, and Staff Office Directors, which involved the exercise of the authority delegated herein prior to the effective date of this delegation.

Dated: June 13, 2007.

Daniel C. Schneider.

Acting Assistant Secretary for Children and Families.

[FR Doc. E7–12019 Filed 6–20–07; 8:45 am] **BILLING CODE 4184–01–P**

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 2007N-0091]

Agency Information Collection
Activities; Submission for Office of
Management and Budget Review;
Comment Request; Preparing a Claim
of Categorical Exclusion or an
Environmental Assessment for
Submission to the Center for Food
Safety and Applied Nutrition

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 July 23, 2007.

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: 202–395–6974. All comments should be identified with the OMB control number 0910–0541. Also include the FDA docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: Jonna Capezzuto, Office of the Chief Information Officer (HFA–250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301–827– 4659

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.

Preparing a Claim of Categorical Exclusion or an Environmental Assessment for Submission to the Center for Food Safety and Applied Nutrition (OMB Control Number 0910– 0541)—Extension

As an integral part of its decisionmaking process, FDA is obligated under the National Environmental Policy Act of 1969 (NEPA) to consider the environmental impact of its actions, including allowing notifications for food contact substances to become effective and approving food additive petitions, color additive petitions, generally recognized as safe affirmation petitions, requests for exemption from regulation as a food additive, and actions on certain food labeling citizen petitions, nutrient content claims petitions, and health claims petitions. In 1997, FDA amended its regulations in part 25 (21 CFR part 25) to provide for categorical exclusions for additional classes of actions that do not individually or cumulatively have a significant effect on the human environment (62 FR 40570, July 29, 1997). As a result of that rulemaking, FDA no longer routinely requires submission of information about the manufacturing and production of FDAregulated articles. FDA also has eliminated the previously required Environmental Assessment (EA) and abbreviated EA formats from the amended regulations. Instead, FDA has provided guidance that contains sample formats to help industry submit a claim of categorical exclusion or an EA to CFSAN. The guidance document entitled "Preparing a Claim of

Categorical Exclusion or an Environmental Assessment for Submission to the Center for Food Safety and Applied Nutrition'' identifies, interprets, and clarifies existing requirements imposed by statute and regulation, consistent with the Council on Environmental Quality regulations (40 CFR 1507.3). It consists of recommendations that do not themselves create requirements; rather, they are explanatory guidance for FDA's own procedures in order to ensure full compliance with the purposes and provisions of NEPA.

The guidance provides information to assist in the preparation of claims of categorical exclusion and EAs for

submission to CFSAN. The following questions are covered in this guidance: (1) What types of industry-initiated actions are subject to a claim of categorical exclusion? (2) What must a claim of categorical exclusion include by regulation? (3) What is an EA? (4) When is an EA required by regulation and what format should be used? (5) What are extraordinary circumstances? and (6) What suggestions does CFSAN have for preparing an EA? Although CFSAN encourages industry to use the EA formats described in the guidance because standardized documentation submitted by industry increases the efficiency of the review process, alternative approaches may be used if

these approaches satisfy the requirements of the applicable statutes and regulations.

FDA is requesting the extension of OMB approval for the information collection provisions in the guidance.

Description of Respondents: The likely respondents include businesses engaged in the manufacture or sale of food, food ingredients, and substances used in materials that come into contact with food.

In the **Federal Register** of March 28, 2007 (72 FR 14581), FDA published a 60-day notice requesting public comment on the information collection provisions. No comments were received.

TABLE 1.—ESTIMATED ANNUAL REPORTING BURDEN¹

21 CFR Section	No. of Respondents	Annual Frequency per Response	Total Annual Responses	Hours per Response	Total Burden Hours
25.32(i)	52	3	156	1	156
25.32(o)	1	1	1	1	1
25.32(q)	7	2	14	1	14
Total			171		171

¹There are no capital costs or operating and maintenance costs associated with this collection of information.

The estimates in table 1 of this document for respondents and numbers of responses are based on the annualized numbers of petitions and notifications qualifying for § 25.32(i) and (q) that the agency has received in the past 3 years. Please note that, in the past 3 years, there have been no submissions that requested an action that would have been subject to the categorical exclusion in § 25.32(o). To avoid counting this burden as zero, FDA has estimated the burden for this categorical exclusion at one respondent making one submission a year for a total of one annual submission.

To calculate the estimate for the hours per response values, we assumed that the information requested in this guidance for each of these three categorical exclusions is readily available to the submitter. For the information requested for the exclusion in § 25.32(i), we expect that submitter will need to gather information from appropriate persons in the submitter's company and to prepare this information for attachment to the claim for categorical exclusion. We believe that this effort should take no longer than 1 hour per submission. For the information requested for the exclusions in § 25.32(o) and (q), the submitters will almost always merely need to copy existing documentation and attach it to the claim for categorical exclusion. We

believe that collecting this information should also take no longer than 1 hour per submission.

Dated: June 14, 2007.

Jeffrey Shuren,

Assistant Commissioner for Policy. [FR Doc. E7–11969 Filed 6–20–07; 8:45 am]

BILLING CODE 4160-01-S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 2007N-0230]

Agency Information Collection Activities; Proposed Collection; Comment Request; Information From United States Processors That Export to the European Community

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing an opportunity for public comment on the proposed collection of certain information by the agency. Under the Paperwork Reduction Act of 1995 (the PRA), Federal agencies are required to publish notice in the Federal Register concerning each proposed collection of information, including each proposed

extension of an existing collection of information, and to allow 60 days for public comment in response to the notice. This notice solicits comments on reporting requirements in implementing the lists of U.S. firms/processors exporting shell eggs, dairy products, game meat and game meat products to the European Community (the EC).

DATES: Submit written or electronic comments on the collection of information by August 20, 2007.

ADDRESSES: Submit electronic comments on the collection of information to: http://www.fda.gov/dockets/ecomments. Submit written comments on the collection of information to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. All comments should be identified with the docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT:

Jonna Capezzuto, Office of the Chief Information Officer (HFA–250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301–827– 4659.

SUPPLEMENTARY INFORMATION: Under the PRA (44 U.S.C. 3501–3520), Federal agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of