2. Form CMS–20017 with complete contact information that includes name, address, phone number, and email addresses for all presenters and commenters and a contact person that can answer any questions, and provide revisions that are requested, for the presentation. Presenters and commenters must clearly explain the actions that they are requesting CMS to take in the appropriate section of the form. A presenter’s or commenter’s relationship with the organization that they represent must also be clearly listed.

- We encourage presenters to make efforts to ensure that their presentations and comments are 508 compliant.

IV. Oral Comments

In addition to formal oral presentations, which are limited to 5 minutes total per presentation, there will be an opportunity during the meeting for public oral comments, which will be limited to 1 minute for each individual and a total of 3 minutes per organization.

V. Meeting Attendance

The meeting is open to the public; however, attendance is limited to space available. Priority will be given to those who pre-register and attendance may be limited based on the number of registrants and the space available. Persons wishing to attend this meeting, which is located on federal property, must register by following the instructions in the DATES section of this notice under “Meeting Registration Timeframe”. A confirmation email will be sent to the registrants shortly after completing the registration process.

VI. Security, Building, and Parking Guidelines

The following are the security, building, and parking guidelines:
- Persons attending the meeting, including presenters, must be pre-registered and on the attendance list by the prescribed date.
- Individuals who are not pre-registered in advance may not be permitted to enter the building and may be unable to attend the meeting.
- Attendees must present a government-issued photo identification to the Federal Protective Service or Guard Service personnel before entering the building. Without a current, valid photo ID, persons may not be permitted entry to the building.
- Security measures include inspection of vehicles, inside and out, at the entrance to the grounds.
- All persons entering the building must pass through a metal detector.
- All items brought into CMS, including personal items, for example, laptops and cell phones, are subject to physical inspection.
- The public may enter the building 30 to 45 minutes before the meeting convenes each day.
- All visitors must be escorted in areas other than the lower and first-floor levels in the Central Building.
- The main-entrance guards will issue parking permits and instructions upon arrival at the building.
- Foreign nationals visiting any CMS facility require prior approval. If you are a foreign national and wish to attend the meeting onsite, in addition to registering for the meeting, you must also send a separate email to APCPanel@cms.hhs.gov prior to the close of registration to request authorization to attend as a foreign national.

Note: As of March 30, 2015, the “Real ID Act” requires a second form of identification from those whose government issued photo identification or government issued driver’s license was issued by American Samoa, Arizona, Louisiana, Maine, Minnesota, and New York. Attendees with a government issued photo identification or driver’s license issued by the states previously mentioned may need to provide alternative or additional approved proof of identification in order to comply with the “Real ID Act.”

VII. Special Accommodations

Individuals requiring special accommodations must include the request for these services during registration.

VIII. Panel Recommendations and Discussions

The Panel’s recommendations at any Panel meeting generally are not final until they have been reviewed and approved by the Panel on the last day of the meeting, before the final adjournment. These recommendations will be posted to our Web site after the meeting.

IX. Collection of Information Requirements

This document does not impose information collection requirements, that is, reporting, recordkeeping, or third-party disclosure requirements. Consequently, there is no need for review by the Office of Management and Budget under the authority of the Paperwork Reduction Act of 1995 (44 U.S.C. 3501 et seq.).

Dated: May 18, 2017.

Seema Verma,
Administrator, Centers for Medicare & Medicaid Services.

[FR Doc. 2017–10683 Filed 5–24–17; 8:45 am]
BILLING CODE 4120–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Food and Drug Administration

Food and Drug Administration


Agency Information Collection Activities; Announcement of Office of Management and Budget Approvals

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is publishing a list of information collections that have been approved by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995.

FOR FURTHER INFORMATION CONTACT: Ila S. Mizrahi, Office of Operations, Food and Drug Administration, Three White Flint North, 11601 Landsdown Street, North Bethesda, MD 20852, 301–796–7726, PRAStaff@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: The following is a list of FDA information collections recently approved by OMB under section 3507 of the Paperwork Reduction Act of 1995 (44 U.S.C. 3507). The OMB control number and expiration date of OMB approval for each information collection are shown in table 1. Copies of the supporting statements for the information collections are available on the Internet at https://www.reginfo.gov/public/do/ PRAMain. 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.
### TABLE 1—LIST OF INFORMATION COLLECTIONS APPROVED BY OMB

<table>
<thead>
<tr>
<th>Title of Collection</th>
<th>OMB Control No.</th>
<th>Date Approval Expires</th>
</tr>
</thead>
<tbody>
<tr>
<td>Procedures for the Safe Processing and Importing of Fish and Fishery Products</td>
<td>0910–0354</td>
<td>2/29/2020</td>
</tr>
<tr>
<td>Biological Products: Reporting of Biological Product Deviations, and Tissue-Based Product Deviations</td>
<td>0910–0458</td>
<td>2/29/2020</td>
</tr>
<tr>
<td>Designation of New Animal Drugs for Minor Use or Minor Species</td>
<td>0910–0605</td>
<td>2/29/2020</td>
</tr>
<tr>
<td>Unique Device Identification System</td>
<td>0910–0720</td>
<td>2/29/2020</td>
</tr>
<tr>
<td>Animal Feed Regulatory Program Standards</td>
<td>0910–0760</td>
<td>2/29/2020</td>
</tr>
<tr>
<td>Premarket Approval of Medical Devices—21 CFR Part 814</td>
<td>0910–0762</td>
<td>2/29/2020</td>
</tr>
<tr>
<td>Human Tissue Intended for Transplantation</td>
<td>0910–0231</td>
<td>3/31/2020</td>
</tr>
<tr>
<td>General Licensing Provisions: Biological License Application, Changes to an Approved Application, Labeling, Revocation and Suspension, and Form FDA 356H</td>
<td>0910–0302</td>
<td>3/31/2020</td>
</tr>
<tr>
<td></td>
<td>0910–0338</td>
<td>3/31/2020</td>
</tr>
</tbody>
</table>

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**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 June 26, 2017.

**ADDRESS:** 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 0910–0520. Also include the FDA docket number found in brackets in the heading of this document.

**FOR FURTHER INFORMATION CONTACT:** Ila S. Mizrachi, Office of Operations, Food and Drug Administration, Three White Flint North, 10A63, 11601 Landsdown St., North Bethesda, MD 20852, 301–796–7726, PRAsstaff@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.

**Prior Notice of Imported Food Under the Public Health Security and Bioterrorism Preparedness and Response Act of 2002—21 CFR 1.278 to 1.285, OMB Control Number 0910–0520**

The Public Health Security and Bioterrorism Preparedness and Response Act of 2002 (the Bioterrorism Act) added section 801(m) of the Federal Food, Drug, and Cosmetic Act (the FD&C Act) (21 U.S.C. 381[m]), which requires that FDA receives prior notice for food, including food for animals, that is imported or offered for import into the United States. Sections 1.278 to 1.282 of FDA regulations (21 CFR 1.278 to 1.282) set forth the requirements for submitting prior notice; §§ 1.283(d) and 1.285(j) (21 CFR 1.283(d) and 1.285(j)) set forth the procedure for requesting the Agency review after FDA has refused admission of an article of food under section 801(m) of the FD&C Act or placed an article of food under hold under section 801(l); and § 1.285(i) sets forth the procedure for post-hold submissions.

**Section 304 of the FDA Food Safety Modernization Act (FSMA) (Pub. L. 111–353) amended section 801(m) of the FD&C Act to require a person submitting prior notice of imported food, including food for animals, to report, in addition to other information already required, “any country to which the article has been refused entry.”**

Advance notice of imported food allows FDA, with the support of the U.S. Customs and Border Protection (CBP), to target import inspections more effectively and help protect the nation’s food supply against terrorist acts and other public health emergencies. By requiring that a prior notice contain additional information that indicates prior refusals by any country and also identifies the country or countries, the Agency may better identify imported food shipments that may pose safety and security risks to U.S. consumers. This additional knowledge can further help FDA to make better informed decisions in managing the potential risks of imported food shipments into the United States.

Any person with knowledge of the required information may submit prior notification of an article of food. Thus, the respondents to this information collection may include importers, owners, ultimate consignees, shippers, and carriers.

FDA regulations require that prior notice of imported food be submitted electronically using CBP’s Automated Broker Interface of the Automated Commercial System (ABI/ACS) (§ 1.280(a)(1)) or the FDA Prior Notice Submission Interface (PNSI) (Form FDA 3540) (§ 1.280(a)(2)). PNSI is an electronic submission system available on the FDA Industry Systems page at https://www.access.fda.gov/.

Information the Agency collects in the prior notice submission includes: (1) The submitter and transmitter (if different from the submitter); (2) entry type and CBP identifier; (3) the article of food, including complete FDA product code; (4) the manufacturer, for an article of food no longer in its natural state; (5) the grower, if known, for an article of food that is in its natural state; (6) the FDA Country of Production; (7) the name of any country that has refused entry of the article of food; (8) the shipper, except for food imported by international mail; (9) the country from which the article of food is shipped or, if the food is imported by international mail, the anticipated date of mailing and country from which the food is mailed; (10) the anticipated arrival information or, if the food is imported by