requirement. The bulletin provides agencies with information on the reporting instructions and training information in the Simplified Mail Accountability Reporting Tool (SMART). FMR Bulletin G–06 and all other FMR bulletins are located at http://www.gsa.gov/frmbulletins.

DATES: Effective Date: October 7, 2015.

FOR FURTHER INFORMATION CONTACT: Ms. Cynthia Patterson, Office of Government-wide Policy (MAF), Office of Asset and Transportation Management, General Services Administration at 703–589–2641 or via email at cynthia.patterson@gsa.gov. Please cite FMR Bulletin G–06.

SUPPLEMENTARY INFORMATION: FMR Bulletin G–06 provides large Federal agencies with reporting instructions and training information in SMART and is consistent with the Federal Management Regulation.


Christine Harada,
Associate Administrator, Office of Government-wide Policy, General Services Administration.

[FR Doc. 2015–25460 Filed 10–6–15; 8:45 am]

BILLING CODE 6820–14–P

DEPARTMENT OF DEFENSE

GENERAL SERVICES ADMINISTRATION

NATIONAL AERONAUTICS AND SPACE ADMINISTRATION

[OMB Control No. 9000–0061; Docket 2015–0055; Sequence 9]

Federal Acquisition Regulation; Submission for OMB Review; Transportation Requirements

AGENCIES: Department of Defense (DOD), General Services Administration (GSA), and National Aeronautics and Space Administration (NASA).

ACTION: Notice of request for public comments regarding an extension to an existing OMB clearance.

SUMMARY: Under the provisions of the Paperwork Reduction Act, the Regulatory Secretariat Division will be submitting to the Office of Management and Budget (OMB) a request to review and approve an extension of a previously approved information collection requirement concerning Transportation Requirements. A notice was published in the Federal Register at 80 FR 34159, on June 15, 2015. No Comments were received.

DATES: Submit comments on or before November 6, 2015.

ADDRESS: Submit comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to: Office of Information and Regulatory Affairs of OMB, Attention: Desk Officer for GSA, Room 10236, NEOB, Washington, DC 20503. Additionally submit a copy to GSA by any of the following methods:

- Regulations.gov: http://www.regulations.gov. Submit comments via the Federal eRulemaking portal by inputting the OMB Control number. Select the link “Submit a Comment” that corresponds with “Information Collection 9000–0061, Transportation Requirements”. Follow the instructions provided at the “Submit a Comment” screen. Please include your name, company name (if any), and “Information Collection 9000–0061, Transportation Requirements” on your attached document.
- Mail: General Services Administration, Regulatory Secretariat Division (MVCB), 1800 F Street NW., Washington, DC 20405. ATTN: Ms. Flowers/IC 9000–0061, Transportation Requirements. Instructions: Please submit comments only and cite Information Collection 9000–0061, Transportation Requirements, in all correspondence related to this collection. All comments received will be posted without change to http://www.regulations.gov, including any personal and/or business confidential information provided.

FOR FURTHER INFORMATION CONTACT: Mr. Curtis E. Glover, Sr., Procurement Analyst, Office of Governmentwide Acquisition Policy, GSA 202–501–1448 or via email at curtis.glover@gsa.gov.

SUPPLEMENTARY INFORMATION:

A. Purpose

FAR Part 47 contains policies and procedures for applying transportation and traffic management considerations in the acquisition of supplies. The FAR part also contains policies and procedures when acquiring transportation or transportation-related services. Generally, contracts involving transportation require information regarding the nature of the supplies, method of shipment, place and time of shipment, applicable charges, marking of shipments, shipping documents and other related items. Contractors are required to provide the information in accordance with the following FAR Part 47 clauses: 52.247–29 through 52.247–44, 52.247–50, 52.247–52, and 52.247–64. The information is used to ensure that: (1) Acquisitions are made on the basis most advantageous to the Government and; (2) supplies arrive in good order and condition, and on time at the required place.

B. Annual Reporting Burden

Respondents: 65,000.

Responses per Respondent: 22.

Annual Responses: 1,430,000.

Hours per Response: 0.5.

Total Burden Hours: 71,500.

C. Public Comments

Public comments are particularly invited on: Whether this collection of information is necessary; whether it will have practical utility; whether our estimate of the public burden of this collection of information is accurate, and based on valid assumptions and methodology; ways to enhance the quality, utility, and clarity of the information to be collected; and ways in which we can minimize the burden of the collection of information on those who are to respond, through the use of appropriate technological collection techniques or other forms of information technology.

Obtaining Copies of Proposals:

Requesters may obtain a copy of the information collection documents from the General Services Administration, Regulatory Secretariat Division (MVCB), 1800 F Street NW., Washington, DC 20405, telephone 202–501–4755. Please cite OMB Control No. 9000–0061, Transportation Requirements, in all correspondence.

Edward Loeb,
Acting Director, Federal Acquisition Policy Division, Office of Governmentwide Acquisition Policy, Office of Acquisition Policy, Office of Governmentwide Policy.

[FR Doc. 2015–25456 Filed 10–6–15; 8:45 am]

BILLING CODE 6820–EP–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Community Living

Agency Information Collection Activities: Submission for OMB Review; Comment Request; Developmental Disabilities Protection and Advocacy Statement of Goals and Priorities

AGENCY: Administration for Community Living, HHS.

ACTION: Notice.

SUMMARY: The Administration on Intellectual and Developmental Disabilities (AIDD), Administration for Community Living (ACL) is announcing an opportunity to comment on the
proposed collection of 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 for public comment in response to the notice. This notice collects comments on the information collection requirements relating to an existing collection: Developmental Disabilities Protection and Advocacy Statement of Goals and Priorities (0985–0034).

DATES: Submit written comments on the collection of information by November 6, 2015.

ADDRESSES: Submit written comments on the collection of information by fax 202.395.5806 or by email to OIRA_submission@omb.eop.gov, Attn: Desk Officer for ACL.


SUPPLEMENTARY INFORMATION: Federal statute and regulation require each State Protection and Advocacy (P&A) System annually prepare for public comment a Statement of Goals and Priorities (SGP) for the P&A for Developmental Disabilities (PADD) program for each coming fiscal year. Following the required public input for the coming fiscal year, the P&A is required by Federal statute and regulation to submit the final version of the SGP to the Administration on Intellectual and Developmental Disabilities (AIDD). AIDD reviews the SGP for compliance and will aggregate the information in the SGPs into a national profile of programmatic emphasis for P&A Systems in the coming year to provide an overview of program direction, and permit AIDD to track accomplishments against goals and formulate areas of technical assistance and compliance with Federal requirements. ACL estimates the burden of this collection of information as follows:

### ANNUAL BURDEN ESTIMATES

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden hours per response</th>
<th>Total burden hours</th>
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<tr>
<td>PADD SGP</td>
<td>57</td>
<td>1</td>
<td>44</td>
<td>2,508</td>
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</tbody>
</table>

**Estimated Total Annual Burden Hours:** 2,508.

Dated: October 1, 2015.

Kathy Greenlee, Administrator & Assistant Secretary for Aging.

[FR Doc. 2015–25592 Filed 10–6–15; 8:45 am]

BILLING CODE 4154–01–P

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**

[Docket No. FDA–2015–N–0001]

**Gastroenterology and Urology Devices Panel of the Medical Devices Advisory Committee; Notice of Meeting**

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice.

This notice announces a forthcoming meeting of a public advisory committee of the Food and Drug Administration (FDA). The meeting will be open to the public.

**Name of Committee:** Gastroenterology and Urology Devices Panel of the Medical Devices Advisory Committee.

**General Function of the Committee:** To provide advice and recommendations to the Agency on FDA’s regulatory issues.

**Dates and Times:** The meeting will be held on November 18, 2015, from 8 a.m. to 6 p.m. and November 19, 2015, from 8 a.m. to 11 a.m.

**Location:** Hilton Washington DC North/Gaithersburg, Salons A, B, and C, 620 Perry Pkwy., Gaithersburg, MD 20877. The hotel’s telephone number is 301–977–8900.

**Contact Person:** Patricia G. Garcia, Center for Devices and Radiological Health, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 66, Rm. 1535, Silver Spring, MD 20993–0002, Patricia.Garcia@fda.hhs.gov, 301–796–6875, or FDA Advisory Committee Information Line, 1–800–741–8138 (301–443–0572 in the Washington, DC area). A notice in the Federal Register about last minute modifications that impact a previously announced advisory committee meeting cannot always be published quickly enough to provide timely notice. Therefore, you should always check the Agency’s Web site at http://www.fda.gov/AdvisoryCommittees/default.htm and scroll down to the appropriate advisory committee meeting link, or call the advisory committee information line to learn about possible modifications before coming to the meeting.

**Agenda:** On November 18, 2015, the committee will discuss, make recommendations and vote on information regarding the premarket approval application (PMA) for the TransMedics® Organ Care System™ (OCS)—Heart, by TransMedics, Inc. The proposed Indication for Use for the TransMedics® Organ Care System™ (OCS)—Heart, as stated in the PMA, is as follows:

The TransMedics® Organ Care System™ (OCS)—Heart is a portable, ex vivo organ perfusion system intended to preserve a donor heart in a near-normothermic and beating state from retrieval until the eventual transplantation into a suitable recipient.

On November 19, 2015, the committee will discuss and make recommendations regarding the classification of the product code “LKK”, and the associated device classification name, “Device, Thermal, Hemorrhoids”. The product code LKX represents a category of devices intended to apply controlled cooling and conductive heating to hemorrhoids. These devices are considered preamendments devices since they were in commercial distribution prior to May 28, 1976, when the Medical Devices Amendments became effective. Some examples of the means by which these devices perform these functions and their respective Indications for Use (IFU)/Intended Use (IU) statements are as follows:

- Uses an aluminum probe that contains a temperature sensitive element to regulate temperature within 2 degrees (between 37 and 46 degrees centigrade).
- IFU/IU: The apparatus is intended to apply controlled, conductive heating to hemorrhoids.
- Uses a heat applicator inserted into the rectum, applicator contains a battery...