health care professionals and other stakeholders about the public health risks posed by counterfeit and unapproved drugs, in addition to safe purchasing practices, and how FDA can evaluate that communication and its impact.

FDA intends to make background material available to the public no later than 2 business days before the meeting. If FDA is unable to post the background material on its Web site prior to the meeting, the background material will be made publicly available at the location of the advisory committee meeting, and the background material will be posted on FDA’s Web site after the meeting. Background material is available at http://www.fda.gov/AdvisoryCommittees/Calendar/default.htm. Scroll down to the appropriate advisory committee meeting link.

Procedure: Interested persons may present data, information, or views, orally or in writing, on issues pending before the committee. Written submissions may be made to the contact person on or before August 8, 2013. Oral presentations from the public will be scheduled between approximately 1 p.m. and 2 p.m. Those individuals interested in making formal oral presentations should notify the contact person and submit a brief statement of the general nature of the evidence or arguments they wish to present, the names and addresses of proposed participants, and an indication of the approximate time requested to make their presentation on or before July 31, 2013. Time allotted for each presentation may be limited. If the number of registrants requesting to speak is greater than can be reasonably accommodated during the scheduled open public hearing session, FDA may conduct a lottery to determine the speakers for the scheduled open public hearing session. The contact person will notify interested persons regarding their request to speak by August 1, 2013. Persons attending FDA’s advisory committee meetings are advised that the Agency is not responsible for providing access to electrical outlets.

FDA welcomes the attendance of the public at its advisory committee meetings and will make every effort to accommodate persons with physical disabilities or special needs. If you require special accommodations due to a disability, please contact Luis G. Bravo at least 7 days in advance of the meeting.

FDA is committed to the orderly conduct of its advisory committee meetings. Please visit our Web site at http://www.fda.gov/AdvisoryCommittees/AboutAdvisoryCommittees/ucm111462.htm for procedures on public conduct during advisory committee meetings.

Notice of this meeting is given under the Federal Advisory Committee Act (5 U.S.C. app. 2).

Dated: July 9, 2013.

Leslie Kux,
Assistant Commissioner for Policy.

[FR Doc. 2013–16831 Filed 7–12–13; 8:45 am]

BILLING CODE 4160–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Notice]

Anesthetic and Analgesic Drug Products Advisory Committee; Cancellation

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The meeting of the Anesthetic and Analgesic Drug Products Advisory Committee scheduled for July 17, 2013, is cancelled. The meeting was announced in the Federal Register of May 17, 2013 (78 FR 29142 to 29143). This meeting has been cancelled due to new information submitted to the application. The Agency intends to continue evaluating the application and, as needed, will announce future meeting dates in the Federal Register.

FOR FURTHER INFORMATION CONTACT: Caleb Briggs, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 31, Rm. 2417, Silver Spring, MD 20993–0002, 301–796–9001, FAX: 301–847–8533, email: AADPAC@fda.hhs.gov, or FDA Advisory Committee Information Line, 1–800–741–8138 (301–443–0572 in the Washington, DC area), and follow the prompts to the desired center or product area. Please call the Information Line for up-to-date information on this meeting.

Dated: July 9, 2013.

Leslie Kux,
Assistant Commissioner for Policy.

[FR Doc. 2013–16823 Filed 7–12–13; 8:45 am]

BILLING CODE 4160–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Services Administration

Agency Information Collection Activities: Proposed Collection: Public Comment Request

AGENCY: Health Resources and Services Administration, HHS.

ACTION: Notice.

SUMMARY: In compliance with the requirement for opportunity for public comment on proposed data collection projects (Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995), the Health Resources and Services Administration (HRSA) announces plans to submit an Information Collection Request (ICR), described below, to the Office of Management and Budget (OMB). Prior to submitting the ICR to OMB, HRSA seeks comments from the public regarding the burden estimate, below, or any other aspect of the ICR.

DATES: Comments on this Information Collection Request must be received within 60 days of this notice.

ADDRESSES: Submit your comments to paperwork@hrsa.gov or mail the HRSA Information Collection Clearance Officer, Room 10–29, Parklawn Building, 5600 Fishers Lane, Rockville, MD 20857.

FOR FURTHER INFORMATION CONTACT: To request more information on the proposed project or to obtain a copy of the data collection plans and draft instruments, email paperwork@hrsa.gov or call the HRSA Information Collection Clearance Officer at (301) 443–1984.

SUPPLEMENTARY INFORMATION: When submitting comments or requesting information, please include the information request collection title for reference.

Information Collection Request Title: National Hospital Organ Donation Campaign’s Activity Scorecard. OMB No. 0915–xxxx—New.

Need and Proposed Use of the Information: HRSA’s Healthcare Systems Bureau, Division of Transplantation administers the Workplace Partnership for Life program under the authority of Section 377A(a) of the Public Health Service (PHS) Act, (42 U.S.C. 274f–1). The Workplace Partnership for Life program seeks to increase the number of registered organ, eye, and tissue donors and to increase awareness about organ donation. HRSA launched a challenge to hospitals nationwide to assist in this effort by conducting donor education and donor
registry enrollment events in their hospitals and communities. The nation’s 58 organ procurement organizations (OPOs), who already work with hospitals on clinical aspects of transplantation, are invited to participate in HRSA’s National Hospital Organ Donation Campaign to increase the number of enrollments in state donor registries. The Campaign supports OPOs by providing fresh communications materials, facilitating the sharing of best practices, leveraging the influence of national associations and organizations related to hospitals and organ donation, and offering the additional incentive of national-level recognition to hospitals.

The National Hospital Organ Donation Campaign’s Activity Scorecard is one piece of this campaign. A campaign leadership committee comprised of representatives from OPOs, Donate Life America (DLA) affiliates, and hospitals helped conceptualize the Activity Scorecard which is based on the committee’s experience of hospital receptivity to friendly competition and the opportunity to be recognized among their peers. The Activity Scorecard provides hospitals that wish to participate in the campaign with ideas for outreach activities. Each activity on the programmable PDF is assigned a particular number of points based on the activity’s potential for generating registrations.

Hospitals can complete the Activity Scorecard and submit it by email or fax it to HRSA or to their OPO or DLA. This is a voluntary activity. Hospitals can participate in the campaign without using the Activity Scorecard. HRSA anticipates that most hospitals enrolled in the campaign (currently 802) will submit a completed Activity Scorecard once a year.

Most importantly, the Activity Scorecard provides incentive for hospitals to conduct activities that will increase the number of registered donors throughout the nation. A list of hospitals that reach these levels will be shared with all campaign participants during monthly webinars, in monthly campaign e-newsletters from HRSA, and in communications pieces sent out by the campaign’s ten national partners, which include the American Hospital Association, the Association of Organ Procurement Organizations, and the American Society of Transplant Surgeons. In addition, OPOs, DLA affiliates, participating state hospital associations, HRSA, and the national partners can use the results to recognize hospital participation and successes. The “write-in” option that allows hospital participation and successes.

Likely Respondents: A hospital representative, most often the organ donation champion identified by the OPO, can download the form from organdonor.gov or receive it from their OPO or Donate Life America (DLA) affiliate.

Burden Statement: Burden in this context means the time expended by persons to generate, maintain, retain, disclose or provide the information requested. This includes the time needed to review instructions; to develop, acquire, install and utilize technology and systems for the purpose of collecting, validating and verifying information, processing and maintaining information, and disclosing and providing information; to train personnel and to be able to respond to a collection of information; to search data sources; to complete and review the collection of information; and to transmit or otherwise disclose the information. The total annual burden hours estimated for this Information Collection Request are summarized in the table below.

Total Estimated Annualized burden hours:

<table>
<thead>
<tr>
<th>Form Name</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Total responses</th>
<th>Average burden per response (in hours)</th>
<th>Total burden hours</th>
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<td>1</td>
<td>802</td>
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<td>802</td>
</tr>
<tr>
<td>Total ..............................................................................</td>
<td>802</td>
<td>1</td>
<td>802</td>
<td>1</td>
<td>802</td>
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</tbody>
</table>

HRSA specifically requests comments on (1) the necessity and utility of the proposed information collection for the proper performance of the agency’s functions, (2) the accuracy of the estimated burden, (3) ways to enhance the quality, utility, and clarity of the information to be collected, and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

Dated: July 8, 2013.

Bahar Niakan,
Director, Division of Policy and Information Coordination.

[FR Doc. 2013–16894 Filed 7–12–13; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Services Administration

Statement of Organization, Functions and Delegations of Authority

This notice amends Part R of the Statement of Organization, Functions and Delegations of Authority of the Department of Health and Human Services (HHS), Health Resources and Services Administration (HRSA) (60 FR 56605, as amended November 6, 1995; as last amended at 78 FR 38720–38723 dated June 27, 2013).

This notice reflects organizational changes in the Health Resources and Services Administration (HRSA). This notice updates the functional statements for the Office of Communications and the Office of Management. Specifically, this notice: (1) Transfers the Freedom of Information Act function from the Office of Communications (RA6) to the Office of Management (RB4), Division of Policy and Information Coordination (RB41); and (2) updates the functional statements for the Office of Communications, the Office of Management, and the Division of Policy and Information Coordination.

Chapter RA6—Office of Communications

Section RA6–20, Functions

Delete the functional statement for the Office of Communications (RA6) and replace in its entirety with the following:

The Office of Communications (RA6) provides leadership and general policy and program direction, and conducts and coordinates communications and public affairs activities of the agency.