Bishop’s Lodge Road, Tesuque, New Mexico, 87574.
• Pojoaque—Pojoaque Valley School District Campus, West Wing Conference Room, 1574 State Road 502 West, Santa Fe, New Mexico 87506.
• Santa Fe—Santa Fe Community College, Jemez Rooms 1&2, 6401 Richards Avenue, Santa Fe, New Mexico 87508.
• Espanola—Northern New Mexico College, Cafeteria, 921 N. Paseo de Oñate, Española, New Mexico 87532.
• Taos—Taos Convention Center, Rio Grande Hall, Room A, 120 Civic Plaza, Taos, New Mexico 87571.

FOR FURTHER INFORMATION CONTACT: Ms. Molly Thrash, Bureau of Reclamation, Albuquerque Area Office, 555 Broadway NE., Suite 100, Albuquerque, New Mexico, 87102; telephone (505) 462–3702; facsimile (505) 462–3780; email stthrash@usbr.gov. Persons who use a telecommunications device for the deaf may call the Federal Information Relay Service (FIRS) at 1–800–877–8339 to contact the above individual during normal business hours. The FIRS is available 24 hours a day, 7 days a week to leave a message or question with the above individual. You will receive a reply during normal business hours.

SUPPLEMENTARY INFORMATION: The Bureau of Reclamation is the lead Federal agency for preparation of the Pojoaque Basin Regional Water System Environmental Impact Statement (EIS). As such, Reclamation published a Notice of Intent to Prepare an EIS on February 24, 2012 (77 FR 11155). The U.S. Army Corps of Engineers, Bureau of Indian Affairs, U.S. Fish and Wildlife Service, Indian Health Service, New Mexico Department of Transportation, New Mexico Office of the State Engineer, County of Santa Fe, City of Santa Fe, and the Pueblos of Nambé, Pojoaque, San Ildefonso, and Tesuque have been invited to participate as cooperating agencies. Other entities may be considered as necessary during the EIS process.

Reclamation is proposing to plan, design, and construct the Pojoaque Basin Regional Water System in accordance with the Aamodt Litigation Settlement Act, which is Title VI of the Claims Resolution Act of 2010 (Pub. L. 111–291, Title VI; 124 Stat. 3065). The proposed project would divert, treat, and distribute potable water to the Pueblo and non-Pueblo residents of the Pojoaque Basin. The Regional Water System would consist of surface water diversion and water treatment facilities within the boundaries of San Ildefonso Pueblo on the Rio Grande and storage tanks, transmission and distribution pipelines, and aquifer storage and recovery well fields that would supply up to 4,000 acre-feet of water annually to customers within the Pojoaque Basin.

Additional Information on the project is available at the project Web site at PojoaqueBasinEIS.com.

Public Scoping
Scoping is an early, ongoing, and open public process for determining the relevant issues to be addressed in the EIS and for identifying any significant issues and suggested alternatives related to the proposed Federal action. Public comments on the scope and content of the EIS may be provided at the public meetings, submitted online through the project Web site, sent via email or facsimile, or mailed to the address shown below. To be most effectively considered, comments should be submitted by May 3, 2013.

Public comments and/or requests to be added to the project mailing list will be accepted at all of the public scoping meetings or by any of the methods shown below:
• Email: PojoaqueBasinEIS@usbr.gov.
• Facsimile: (505) 462–3780.
• Web site: PojoaqueBasinEIS.com.
• Address: Bureau of Reclamation, Albuquerque Area Office, Suite 100 (ALB–842), 555 Broadway NE., Suite 100, Albuquerque, New Mexico, 87102.

In addition to the public scoping meetings described above, Reclamation may host additional scoping meetings with Pueblo members at or near each of the four Pueblos. Government-to-government consultation will continue with the Pueblo governments and coordination will continue with other Federal and State agencies.

Special Assistance for Public Meetings
If special assistance is required to participate in a particular scoping meeting, please contact Ms. Molly Thrash at (505) 462–3702, or via email at stthrash@usbr.gov. A telephone device for the hearing impaired is available at 1–800–877–8339. Please provide notification far in advance as possible to enable Reclamation to secure the needed services. If a request cannot be honored, the requester will be notified.

Public Disclosure
Before including your address, phone number, email address, or other personal identifying information in any communication, you should be aware that your entire comments—including your personal identifying information—may be made publicly available at any time. We may ask you in your communication to withhold your personal identifying information from public review, we cannot guarantee that we will be able to do so.

Dated: January 16, 2013.
Larry Walkoviak,
Regional Director—Upper Colorado Region, Bureau of Reclamation.

DEPARTMENT OF JUSTICE
Drug Enforcement Administration
Importer of Controlled Substances, Notice of Application; Meridian Medical Technologies

Pursuant to Title 21 Code of Federal Regulations 1301.34 (a), this is notice that on January 8, 2013, Meridian Medical Technologies, 2555 Hermelin Drive, St. Louis, Missouri 63144, made application by renewal to the Drug Enforcement Administration (DEA) to be registered as an importer of Morphine (93900), a basic class of controlled substance listed in schedule II.

The company manufactures a product containing morphine in the United States. The company exports this product to customers around the world. The company has been asked to ensure that its product sold to European customers meets standards established by the European Pharmacopeia, which is administered by the Directorate for the Quality of Medicines (EDQM) in order to ensure that its product will meet European specifications, the company seeks to import morphine supplied by EDQM to use as reference standards. This is the sole purpose for which the company will be authorized by DEA to import morphine.

Any bulk manufacturer who is presently, or is applying to be, registered with DEA to manufacture such basic class of controlled substance listed in schedule II, which falls under the authority of section 1002(a)(2)(B) of the Act (21 U.S.C. 952(a)(2)(B)) may, in the circumstances set forth in 21 U.S.C. 958(i), file comments or objections to the issuance of the proposed registration and may, at the same time, file a written request for a hearing on such application pursuant to 21 CFR 1301.43 in such form as prescribed by 21 CFR 1316.47.

Any such written comments or objections should be addressed, in quintuplicate, to the Drug Enforcement Administration, Office of Diversion Control, Federal Register Representative (ODL), 555 Broadway NE., Suite 100, Albuquerque, New Mexico, 87102.

Notices
This procedure is to be conducted simultaneously with, and independent of, the procedures described in 21 CFR 1301.34(b), (c), (d), (e), and (f). As noted in a previous notice published in the Federal Register on September 23, 1975, 40 FR 43745–46, all applicants for registration to import a basic class of any controlled substances in schedules I or II are, and will continue to be, required to demonstrate to the Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration, that the requirements for such registration pursuant to 21 U.S.C. 958(a); 21 U.S.C. 823(a); and 21 CFR 1301.34(b), (c), (d), (e), and (f) are satisfied.

Dated: March 7, 2013.

Joseph T. Rannazzisi,
Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration.

[FR Doc. 2013–05794 Filed 3–12–13; 8:45 am]
BILLING CODE 4410–09–P

DEPARTMENT OF JUSTICE
Drug Enforcement Administration

Drug Enforcement Administration

Importer of Controlled Substances, Notice of Registration: Hospira Inc.

By Notice dated December 14, 2012, and published in the Federal Register on December 21, 2012, 77 FR 75670, Hospira Inc., 1776 North Centennial Drive, McPherson, Kansas 67460–1247, made application by renewal to the Drug Enforcement Administration (DEA) to be registered as an importer of Remifentanil (9739), a basic class of controlled substance listed in schedule II.

The company plans to import Remifentanil for use in dosage form manufacturing.

No comments or objections have been received. DEA has considered the factors in 21 U.S.C. 823(a) and 952(a), and determined that the registration of Hospira Inc. to import the basic class of controlled substance is consistent with the public interest, and with United States obligations under international treaties, conventions, or protocols in effect on May 1, 1971, at this time. DEA has investigated Hospira Inc. to ensure that the company's registration is consistent with the public interest. The investigation has included inspection and testing of the company's physical security systems, verification of the company's compliance with state and local laws, and a review of the company's background and history. Therefore, pursuant to 21 U.S.C. 958(a), and in accordance with 21 CFR 1301.34, the above named company is granted registration as an importer of the basic class of controlled substance listed.

Dated: March 7, 2013.

Joseph T. Rannazzisi,
Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration.

[FR Doc. 2013–05794 Filed 3–12–13; 8:45 am]
BILLING CODE 4410–09–P

DEPARTMENT OF JUSTICE
Drug Enforcement Administration

Manufacturer of Controlled Substances, Notice of Registration; Johnson Matthey, Inc., Pharmaceuticals Materials

By Notice dated November 1, 2012, and published in the Federal Register on November 9, 2012, 77 FR 67397, Johnson Matthey Inc., Pharmaceuticals Materials, 900 River Road, Conshohocken, Pennsylvania 19428, made application by renewal to the Drug Enforcement Administration (DEA) to be registered as a bulk manufacturer of the following basic classes of controlled substances:

<table>
<thead>
<tr>
<th>Drug</th>
<th>Schedule</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gamma Hydroxybutyric Acid</td>
<td>I</td>
</tr>
<tr>
<td>Amphetamine (1100)</td>
<td>II</td>
</tr>
<tr>
<td>Methylphenidate (1724)</td>
<td>II</td>
</tr>
<tr>
<td>Codeine (9050)</td>
<td>II</td>
</tr>
<tr>
<td>Oxycodone (9143)</td>
<td>II</td>
</tr>
<tr>
<td>Diphenoxylate (9170)</td>
<td>II</td>
</tr>
<tr>
<td>Hydrocodone (9193)</td>
<td>II</td>
</tr>
<tr>
<td>Meperidine (9230)</td>
<td>II</td>
</tr>
<tr>
<td>Methadone (9250)</td>
<td>II</td>
</tr>
<tr>
<td>Methadone intermediate (9224)</td>
<td>II</td>
</tr>
<tr>
<td>Morphine (9300)</td>
<td>II</td>
</tr>
<tr>
<td>Thebaine (9333)</td>
<td>II</td>
</tr>
</tbody>
</table>

The company plans to manufacture the listed controlled substances in bulk for distribution and sale to its customers. The Thebaine (9333) will also be used to manufacture other controlled substances for sale in bulk to its customers.

No comments or objections have been received. DEA has considered the factors in 21 U.S.C. 823(a) and 952(a), and determined that the registration of Johnson Matthey Inc., Pharmaceuticals Materials to manufacture the listed basic classes of controlled substances is consistent with the public interest at this time. DEA has investigated Johnson Matthey Inc., Pharmaceuticals Materials to ensure that the company's registration is consistent with the public interest. The investigation has included inspection and testing of the company's physical security systems, verification of the company's compliance with state and local laws, and a review of the company's background and history.

Therefore, pursuant to 21 U.S.C. 823(a), and in accordance with 21 CFR 1301.33, the above named company is granted registration as a bulk manufacturer of the basic classes of controlled substances listed.

Dated: February 27, 2013.

Joseph T. Rannazzisi,
Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration.

[FR Doc. 2013–05799 Filed 3–12–13; 8:45 am]
BILLING CODE 4410–09–P