and, if known, an estimate of the percentage of total U.S. imports of *Subject Merchandise* from the *Subject Country* accounted for by your firm's(s') imports;

(b) the quantity and value (f.o.b. U.S. port, including antidumping countervailing duties) of U.S. commercial shipments of *Subject Merchandise* imported from the *Subject Country*; and

(c) the quantity and value (f.o.b. U.S. port, including antidumping countervailing duties) of U.S. internal consumption/company transfers of *Subject Merchandise* imported from the *Subject Country*.

(11) If you are a producer, an exporter, or a trade/business association of producers or exporters of the *Subject Merchandise* in the *Subject Country*, provide the following information on your firm's(s') operations on that product during calendar year 2011 (report quantity data in pieces and value data in U.S. dollars, landed and dutypaid at the U.S. port but not including antidumping duties). If you are a trade/ business association, provide the information, on an aggregate basis, for the firms which are members of your association.

(a) Production (quantity) and, if known, an estimate of the percentage of total production of *Subject Merchandise* in the *Subject Country* accounted for by your firm's(s') production;

(b) Capacity (quantity) of your firm to produce the *Subject Merchandise* in the *Subject Country* (i.e., the level of production that your establishment(s) could reasonably have expected to attain during the year, assuming normal operating conditions (using equipment and machinery in place and ready to operate), normal operating levels (hours per week/weeks per year), time for downtime, maintenance, repair, and cleanup, and a typical or representative product mix); and

(c) The quantity and value of your firm's(s') exports to the United States of *Subject Merchandise* and, if known, an estimate of the percentage of total exports to the United States of *Subject Merchandise* from the *Subject Country* accounted for by your firm's(s') exports.

(12) Identify significant changes, if any, in the supply and demand conditions or business cycle for the *Domestic Like Product* that have occurred in the United States or in the market for the *Subject Merchandise* in the *Subject Country* after 2005, and significant changes, if any, that are likely to occur within a reasonably foreseeable time. Supply conditions to consider include technology; production methods; development

efforts; ability to increase production (including the shift of production facilities used for other products and the use, cost, or availability of major inputs into production); and factors related to the ability to shift supply among different national markets (including barriers to importation in foreign markets or changes in market demand abroad). Demand conditions to consider include end uses and applications; the existence and availability of substitute products; and the level of competition among the Domestic Like Product produced in the United States, Subject Merchandise produced in the Subject Country, and such merchandise from other countries.

(13) (Optional) A statement of whether you agree with the above definitions of the *Domestic Like Product* and *Domestic Industry;* if you disagree with either or both of these definitions, please explain why and provide alternative definitions.

Authority: This review is being conducted under authority of title VII of the Tariff Act of 1930; this notice is published pursuant to section 207.61 of the Commission=s rules.

By order of the Commission. Issued: March 27, 2012.

James R. Holbein,

Secretary to the Commission. [FR Doc. 2012–7794 Filed 3–30–12; 8:45 am] BILLING CODE 7020–02–P

DEPARTMENT OF JUSTICE

Notice of Filing of Consent Decree Pursuant to the Clean Air Act, CERCLA and EPCRA

Notice is hereby given that on March 26, 2012, a proposed Consent Decree in United States and State of Kansas v. National Cooperative Refinery Association, No. 6:12-cv-01110-EFM-JPO, was filed with the United States District Court for the District of Kansas. The Consent Decree settles the claims of the United States' and the State of Kansas set forth in the complaint for civil penalties and injunctive relief against the National Cooperative Refinery Association relating to its refinery in McPherson, Kansas, and its related storage facility in Conway, Kansas, based on violations of the refinery's Clean Air Act Title V permit, the CAA's New Source Performance Standards (NSPS), and Risk Management Plan regulations, and CERCLA and EPCRA release notification requirements. Under the terms of the Consent Decree NCRA shall pay \$700,000 in penalty, spend approximately \$730,000 in performing

Supplemental Environmental Projects and implement injunctive relief directed primarily at insuring future compliance with the Risk Management Program requirements for these facilities. \$225,000 of this penalty will be paid to the State of Kansas.

The Department of Justice will receive for a period of thirty (30) days from the date of this publication comments relating to the Consent Decree. Comments should be addressed to the Assistant Attorney General, **Environment and Natural Resources** Division, and either emailed to pubcomment-ees.enrd@usdoj.gov or mailed to P.O. Box 7611, U.S. Department of Justice, Washington, DC 20044-7611, and should refer to United States and State of Kansas v. National Cooperative Refinerv Association, No. 6:12-cv-01110-EFM-JPO (D. Kansas), Department of Justice Case Number 90-5-1-1-06025/3.

During the public comment period, the Consent Decree may be examined at the Office of the United States Attorney, District of Kansas, 301 North Main St. Wichita, Kansas, 67212. The Settlement Agreement may also be examined on the following Department of Justice Web site, http://www.usdoj.gov/enrd/ *Consent Decrees.html*. A copy of the Consent Decree may also be obtained by mail from the Consent Decree Library, P.O. Box 7611, U.S. Department of Justice, Washington, DC 20044-7611 or by faxing or emailing a request to "Consent Decree Copy" (eescdcopy.enrd@usdoj.gov), fax no. (202) 514-0097, phone confirmation number (202) 514–5271. In requesting a copy from the Consent Decree Library, please enclose a check in the amount of \$22.75 (25 cents per page reproduction cost) payable to the U.S. Treasury.

Maureen Katz,

Assistant Section Chief. [FR Doc. 2012–7714 Filed 3–30–12; 8:45 am] BILLING CODE 4410–15–P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

Importer of Controlled Substances; Notice of Application Meridian Medical Technologies

Pursuant to 21 U.S.C. 958(i), the Attorney General shall, prior to issuing a registration under this Section to a bulk manufacturer of a controlled substance in schedule I or II, and prior to issuing a regulation under 21 U.S.C. 952(a)(2) authorizing the importation of such a substance, provide manufacturers holding registrations for the bulk manufacture of the substance an opportunity for a hearing.

Therefore, in accordance with 21 CFR 1301.34(a), this is notice that on January 4, 2012, 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 (9300), 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, including in Europe. 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 may 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, and 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), 8701 Morrissette Drive, Springfield, Virginia 22152; and must be filed no later than May 2, 2012.

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 substance in schedule 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 23, 2012. Joseph T. Rannazzisi, Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration. [FR Doc. 2012–7755 Filed 3–30–12; 8:45 am]

BILLING CODE 4410-09-P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

Importer of Controlled Substances; Notice of Registration Mallinckrodt LLC

By Notice dated January 23, 2012, and published in the **Federal Register** on January 31, 2012, 77 FR 4831, Mallinckrodt LLC, 3600 North Second Street, St. Louis, Missouri 63147, made application by renewal to the Drug Enforcement Administration (DEA) to be registered as an importer of the following basic classes of controlled substances:

Drug	Schedule
Phenylacetone (8501) Coca Leaves (9040) Opium, raw (9600) Poppy Straw Concentrate (9670)	= = =

The company plans to import the listed controlled substances to manufacture bulk controlled substances for distribution to its customers, and for research and analytical standards.

The company has withdrawn its application for registration to import the following drug codes: Methylphenidate (1724), Oxycodone (9143), Hydromorphone (9150), Hydrocodone (9193), Morphine (9300), and Fentanyl (9801).

Comments and requests for hearings on applications to import narcotic raw material are not appropriate, 72 FR 3417 (2007). Regarding all other basic classes of controlled substances, 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 Mallinckrodt LLC, to import the basic classes of controlled substances is consistent with the public interest, and with United States obligations under international treaties, conventions, or protocols in effect on May 1, 1971. DEA has investigated Mallinckrodt LLC 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. 952(a) and 958(a), and in accordance with 21 CFR 1301.34, the above named company is granted registration as an importer of the basic classes of controlled substances listed.

Dated: March 23, 2012.

Joseph T. Rannazzisi,

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

[FR Doc. 2012–7758 Filed 3–30–12; 8:45 am] BILLING CODE 4410–09–P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

Manufacturer of Controlled Substances; Notice of Application; Cerilliant Corporation

Pursuant to § 1301.33(a), Title 21 of the Code of Federal Regulations (CFR), this is notice that on February 17, 2012, Cerilliant Corporation, 811 Paloma Drive, Suite A, Round Rock, Texas 78665–2402, made application by letter to the Drug Enforcement Administration (DEA) to be registered as a bulk manufacturer of the following basic classes of controlled substances:

Drug	Schedule
4-methyl-N-methylcathinone (1248).	I
3,4-methylenedioxypyrovalerone (7535).	1
3,4-methylenedioxy-N- methylcathinone (7540).	I
Desomorphine (9055)	1

The company plans to manufacture small quantities of the listed controlled substances to make reference standards which will be distributed to their customers.

Any other such applicant, and any person who is presently registered with DEA to manufacture such substances, may file comments or objections to the issuance of the proposed registration pursuant to 21 CFR 1301.33(a).

Any such written comments or objections should be addressed, in quintuplicate, to the Drug Enforcement Administration, Office of Diversion Control, **Federal Register** Representative (ODL), 8701 Morrissette Drive, Springfield, Virginia 22152; and must be filed no later than June 1, 2012.