presentations should attend a prehearing conference to be held at 9:30 a.m. on May 28, 2009 at the U.S. International Trade Commission Building. Oral testimony and written materials to be submitted at the hearing are governed by sections 201.6(b)(2) and 201.13(f) of the Commission's rules.

Written submissions.—Each party is encouraged to submit a prehearing brief to the Commission. The deadline for filing prehearing briefs is May 28, 2009. Parties may also file posthearing briefs. The deadline for filing posthearing briefs is June 8, 2009. In addition, any person who has not entered an appearance as a party to the investigation may submit a written statement of information pertinent to the consideration of market disruption or threat thereof and/or remedy on or before June 8, 2009. Parties may submit final comments on market disruption on or before June 16, 2009 and on remedy on or before June 24, 2009. Final comments shall contain no more than ten (10) double-spaced and single-sided pages of textual material. All written submissions must conform with the provisions of section 201.8 of the Commission's rules; any submissions that contain confidential business information must also conform with the requirements of section 201.6 of the Commission's rules. The Commission's rules authorize filing submissions with the Secretary by facsimile or electronic means only to the extent permitted by section 201.8 of the rules (see Handbook for Electronic Filing Procedures, http:// www.usitc.gov/secretary/ fed reg notices/rules/documents/ handbook_on_electronic filing.pdf). Persons with questions regarding

In accordance with section 201.16(c) of the Commission's rules, each document filed by a party to the investigation must be served on all other parties to the investigation (as identified by the service list), and a certificate of service must be timely filed. The Secretary will not accept a document for filing without a certificate of service.

electronic filing should contact the

Secretary (202-205-2000).

Remedy.—No separate hearing on the issue of remedy will be held. Those parties wishing to present arguments on the issue of remedy may do so orally at the hearing or in their prehearing or posthearing briefs or other written submissions.

Authority: This investigation is being conducted under the authority of section 421 of the Trade Act of 1974; this notice is published pursuant to section 206.3 of the Commission's rules.

By order of the Commission.

Marilyn R. Abbott,

Secretary to the Commission. Issued: April 24, 2009.

William R. Bishop,

Acting Secretary to the Commission.
[FR Doc. E9–9760 Filed 4–28–09; 8:45 am]
BILLING CODE P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

Importer of Controlled Substances; Notice of Application

This is notice that on March 10, 2009, Penick Corporation, 33 Industrial Park Road, Pennsville, New Jersey 08070, made application by renewal to the Drug Enforcement Administration (DEA) for registration as an importer of the basic classes of controlled substances listed in schedule II.

Drug	Schedule
Coca Leaves (9040)	
(9670)	ll II

The company plans to import the listed controlled substances to manufacture bulk controlled substance intermediates for sale to its customers.

As noted in a previous notice published in the **Federal Register** on September 23, 1975 (40 FR 43745), all applicants for registration to import a basic class of any controlled substances 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: April 17, 2009.

Joseph T. Rannazzisi,

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

[FR Doc. E9–9778 Filed 4–28–09; 8:45 am] BILLING CODE 4410–09–P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

Importer of Controlled Substances; Notice of Application

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 Title 21 Code of Federal Regulations § 1301.34(a), this is notice that on February 20, 2009, 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 plans to import products for research experimentation or clinical use and analytical testing.

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 comments or objections being sent via regular mail 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 29, 2009.

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 listed 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 USC § 823(a), and 21 CFR 1301.34(b), (c), (d), (e), and (f) are satisfied.

Dated: April 17, 2009.

Joseph T. Rannazzisi,

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

[FR Doc. E9–9782 Filed 4–28–09; 8:45 am]

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

Importer of Controlled Substances; Notice of Application

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 Title 21 Code of Federal Regulations § 1301.34(a), this is notice that on March 10, 2009, Almac Clinical Services Inc. (ACSI), 2661 Audubon Road, Audubon, Pennsylvania 19403, has made application by renewal to the Drug Enforcement Administration (DEA) to be registered as an importer of the basic classes of controlled substances listed in schedule II:

Drug	Schedule
Oxycodone (9143) Fentanyl (9801)	11

The company plans to import small quantities of the listed controlled substances in dosage form to conduct clinical trials.

Any bulk manufacturer who is presently, or is applying to be, registered with DEA to manufacture such basic classes of controlled substances 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 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 29, 2009.

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 the 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: April 17, 2009.

Joseph T. Rannazzisi,

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

[FR Doc. E9–9780 Filed 4–28–09; 8:45 am]

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

Importer of Controlled Substances; Notice of Registration

By Notice dated January 30, 2009, and published in the **Federal Register** on February 6, 2009 (74 FR 6309), Mallinckrodt Inc., 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 basic classes of controlled substances listed in schedule II:

Drug	Schedule
Phenylacetone (8501)	

The company plans to import the listed controlled substances for the manufacture of controlled substances in bulk for distribution 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 Mallinckrodt Inc. 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, at this time. DEA has investigated Mallinckrodt 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. 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: April 17, 2009.

Joseph T. Rannazzisi,

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

[FR Doc. E9–9789 Filed 4–28–09; 8:45 am] BILLING CODE 4410–09–P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

Manufacturer of Controlled Substances; Notice of Registration

By Notice dated December 22, 2008, and published in the Federal Register on December 31, 2008 (73 FR 80431), Norac Inc., 405 S. Motor Avenue, P.O. Box 577, Azusa, California 91702–3232, made application by renewal to the Drug Enforcement Administration (DEA) to be registered as a bulk manufacturer of Tetrahydrocannabinols (7370), a basic class of controlled substance listed in schedule I.

The company plans to manufacture the listed controlled substance in bulk for formulation into the pharmaceutical controlled substance Marinol® for sale to its customers.

No comments or objections have been received. DEA has considered the factors in 21 U.S.C. 823(a) and determined that the registration of Norac Inc. to manufacture the listed basic class of controlled substance is consistent with the public interest at this time. DEA has investigated Norac 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. 823, and in accordance with 21 CFR 1301.33, the above named company is granted registration as a bulk manufacturer of the basic class of controlled substance listed.