The last notification was filed with the Department on October 22, 2015. A notice was published in the **Federal Register** pursuant to Section 6(b) of the Act on December 7, 2015 (80 FR 76042).

Patricia A. Brink,

Director of Civil Enforcement, Antitrust Division.

[FR Doc. 2016–08584 Filed 4–13–16; 8:45 am] BILLING CODE P

DEPARTMENT OF JUSTICE

Antitrust Division

Notice Pursuant to the National Cooperative Research and Production Act of 1993—Cooperative Research Group on Chede–VII

Notice is hereby given that, on March 15, 2016, pursuant to Section 6(a) of the National Cooperative Research and Production Act of 1993, 15 U.S.C. 4301 et seq. ("the Act"), Southwest Research Institute—Cooperative Research Group on CHEDE-VII ("CHEDE-VII") has filed written notifications simultaneously with the Attorney General and the Federal Trade Commission disclosing changes in its membership. The notifications were filed for the purpose of extending the Act's provisions limiting the recovery of antitrust plaintiffs to actual damages under specified circumstances. Specifically, Detroit Diesel Corporation, Detroit, MI; Mahle Engine Components USA, Inc., Farmington Hills, MI; Volvo Powertrain North America, Hagerstown, MD; Tata Motors Ltd., Mumbai, INDIA; Convergent Science, Madison, WI; and PACCAR C/O DAF Trucks N.V., Eindhoven, NETHERLANDS, have been added as parties to this venture.

No other changes have been made in either the membership or planned activity of the group research project. Membership in this group research project remains open, and CHEDE-VII intends to file additional written notifications disclosing all changes in membership.

On January 6, 2016, CHEDE–VII filed its original notification pursuant to Section 6(a) of the Act. The Department of Justice published a notice in the **Federal Register** pursuant to Section 6(b) of the Act on February 2, 2016, (81 FR 5484).

The last notification was filed with the Department on February 10, 2016. A notice was published in the **Federal** **Register** pursuant to section 6(b) of the Act on March 9, 2016, (81 FR 12529).

Patricia A. Brink,

Director of Civil Enforcement, Antitrust Division.

[FR Doc. 2016–08580 Filed 4–13–16; 8:45 am]

DEPARTMENT OF JUSTICE

Antitrust Division

Notice Pursuant to the National Cooperative Research and Production Act of 1993—National Armaments Consortium

Notice is hereby given that, on March 15, 2016, pursuant to Section 6(a) of the National Cooperative Research and Production Act of 1993, 15 U.S.C. 4301 et seq. ("the Act"), National Armaments Consortium ("NAC") has filed written notifications simultaneously with the Attorney General and the Federal Trade Commission disclosing changes in its membership. The notifications were filed for the purpose of extending the Act's provisions limiting the recovery of antitrust plaintiffs to actual damages under specified circumstances. Specifically, Akita Innovations, LLC, North Billerica, MA; American Engineering & Manufacturing, Inc., Elvria, OH; American Rheinmetall Munitions, Inc., Stafford, VA; Aria Microwave Systems, Inc., Teaneck, NJ; BEAM Engineering for Advanced Measurements, Orlando, FL; Bradshaw Engineering and Technical Services, LLC, Union Grove, AL; C-2 Innovations, Inc., Stow, MA; CIRTEMO, LLC, Cayce, SC; Cummings Aerospace, Inc., Huntsville, AL; Elbit Systems of America, LLC, Fort Worth, TX; Electronics & Manufacturing Co., LLC, Columbia, MO; Elmet Technologies, LLC, Lewiston, ME; Evigia Systems, Inc., Ann Arbor, MI; Fairlead Precision Manufacturing & Integration, LLC, Portsmouth, VA; Fiocchi of America, Ozark, MO; GECO, Inc., Mesa, AZ; General Dynamics C4 Systems, Inc., Scottsdale, AZ; Hardigg Industries, Inc., South Deerfield, MA; Integrated Global Insights, LLC, Burke, VA; Integration Innovation, Inc. (i3), Huntsville, AL; Kratos Defense & Rocket Support Services, Inc., King George, VA; L-3 Electron Devices, Williamsport, PA; Lancer Systems, LP, Quakertown, PA; Leigh Aerosystems Corporation, Carlsbad, CA; Logikos, Inc., Fort Wayne, IN; Magnesium Elektron North American, Inc., Madison, IL; Materion Brush, Inc., Elmore, OH; MBDA Incorporated, Arlington, VA; Molex, LLC, Lisle, IL; OMNI Consulting

Solutions, LLC, El Segundo, CA; Orion Munitions Development, LLC, Gladstone, MO; pH Matter, LLC, Columbus, OH; PolyPlus Battery Company, Berkeley, CA; Polysciences, Inc., Warrington, PA; Pyrolink International Inc., Alexandria, VA; Quantum Dimension, Inc., Huntington Beach, CA; Radiation Monitoring Devices, Inc., Watertown, PA; Saab Defense and Security USA, LLC, East Syracuse, NY; SimVentions, Inc., Fredericksburg, VA; Southern Research Institute, Birmingham, AL; STS Technologies, LLC, Mahwah, NJ; The Regents of the University of California, Irvine, Irvine, CA; Vadum, Inc., Raleigh, NC; ViaSat, Inc., Gilbert, AZ; and Wilcox Industries Corp, Newintong, PA, have been added as parties to this venture.

Also, Applied Thin Films, Inc., Skokie, IL; Fluorochem, Inc., Azusa, CA; Prime Photonics, LC, Blacksburg, VA; Safety Consulting Engineers, Schauraburg, IL; Saint-Gobain Ceramics & Plastics, Inc., Milford, NH; and Soligie, Inc., Savage, MN, have withdrawn as parties to this venture.

No other changes have been made in either the membership or planned activity of the group research project. Membership in this group research project remains open, and NAC intends to file additional written notifications disclosing all changes in membership.

On May 2, 2000, NAC filed its original notification pursuant to Section 6(a) of the Act. The Department of Justice published a notice in the **Federal Register** pursuant to Section 6(b) of the Act on June 30, 2000 (65 FR 40693).

The last notification was filed with the Department on November 10, 2015. A notice was published in the **Federal Register** pursuant to Section 6(b) of the Act on December 23, 2015 (80 FR 79931).

Patricia A. Brink,

Director of Civil Enforcement, Antitrust Division.

[FR Doc. 2016–08577 Filed 4–13–16; 8:45 am]

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

[Docket No. DEA-392]

Manufacturer of Controlled Substances Registration: Stepan Company

ACTION: Notice of registration.

SUMMARY: Stepan Company applied to be registered as a manufacturer of

certain basic classes of controlled substances. The Drug Enforcement Administration (DEA) grants Stepan Company registration as a manufacturer of those controlled substances.

SUPPLEMENTARY INFORMATION: By notice dated April 14, 2015, and published in the Federal Register on April 22, 2015, 80 FR 22555, Stepan Company, Natural Products Department, 100 W. Hunter Avenue, Maywood, New Jersey 07607 applied to be registered as a manufacturer of certain basic classes of controlled substances. No comments or objections were submitted for this notice.

The DEA has considered the factors in 21 U.S.C. 823(a) and determined that the registration of Stepan Company to manufacture 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. The DEA investigated the company's maintenance of effective controls against diversion by inspecting and testing the company's physical security systems, verifying the company's compliance with state and local laws, and reviewing 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 following basic classes of controlled substances:

Controlled substance	Schedule
Cocaine (9041)	II
Ecgonine (9180)	II

The company plans to manufacture the listed controlled substances in bulk for distribution to its customers.

Dated: April 4, 2016 Louis J. Milione,

Deputy Assistant Administrator. [FR Doc. 2016–08576 Filed 4–13–16; 8:45 am]

BILLING CODE 4410-09-P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

[Docket No. DEA-392]

Bulk Manufacturer of Controlled Substances Application: Patheon API Manufacturing, Inc.

ACTION: Notice of application.

DATES: Registered bulk manufacturers of the affected basic classes, and applicants therefore, may file written comments on or objections to the

issuance of the proposed registration in accordance with 21 CFR 1301.33(a) on or before June 13, 2016.

ADDRESSES: Written comments should be sent to: Drug Enforcement Administration, Attention: DEA Federal Register Representative/ODW, 8701 Morrissette Drive, Springfield, Virginia 22152.

SUPPLEMENTARY INFORMATION: The Attorney General has delegated her authority under the Controlled Substances Act to the Administrator of the Drug Enforcement Administration (DEA), 28 CFR 0.100(b). Authority to exercise all necessary functions with respect to the promulgation and implementation of 21 CFR part 1301, incident to the registration of manufacturers, distributors, dispensers, importers, and exporters of controlled substances (other than final orders in connection with suspension, denial, or revocation of registration) has been redelegated to the Deputy Assistant Administrator of the DEA Office of Diversion Control ("Deputy Assistant Administrator") pursuant to section 7 of 28 CFR part 0, appendix to subpart R.

In accordance with 21 CFR 1301.33(a), this is notice that on January 13, 2016, Patheon API Manufacturing, Inc., 309 Delaware Street, Building 1106, Greenville, South Carolina 29605 applied to be registered as a bulk manufacturer of the following basic classes of controlled substances:

Controlled substance	Schedule
Marihuana (7360) Tetrahydrocannabinols (7370) Noroxymorphone (9668)	

The company plans to manufacture the above-listed controlled substances as Active Pharmaceutical Ingredients (API) for clinical trials.

In reference to drug codes 7360 (marihuana), and 7370 (THC), the company plans to bulk manufacture these drugs as synthetics. No other activities for these drug codes are authorized for this registration.

Dated: March 29, 2016.

Louis J. Milione,

Deputy Assistant Administrator. [FR Doc. 2016–08569 Filed 4–13–16; 8:45 am]

BILLING CODE 4410-09-P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration [Docket No. 16–7]

Rezik A. Saqer, M.D.; Decision and Order

On October 1, 2015, the Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration, issued an Order to Show Cause to Rezik A. Sager, M.D., (Respondent). The Show Cause Order proposed the revocation of Respondent's DEA Certificates of Registration BS4072637 and FS1975359, pursuant to which he is authorized to dispense controlled substances in schedules II through V, as a practitioner, at the respective registered locations of 11037 FM 1960 West, Suite B1, Houston, Texas, and 3074 College Park Drive, Conroe, Texas. Show Cause Order, at 1. The Show Cause Order further proposed the denial of any applications to renew or modify either registration, as well as the denial of any other application for a DEA registration.

More specifically, the Show Cause Order alleged that "[e]ffective September 28, 2015, the Texas Medical Board issued an Order of Temporary Suspension . . . which suspended [Respondent's] medical license," and therefore, he is currently "without authority to handle controlled substances in Texas, the State in which [he is] registered with" DEA. Id. at 2. The Show Cause Order thus advised Respondent that "DEA must revoke [his] registrations based upon [his] lack of authority to handle controlled substances in the State of Texas." Id. (citing 21 U.S.C. 802(21), 823(f), and 824(a)(3)).

On October 2, 2015, a Diversion Investigator served the Show Cause Order by travelling to Respondent's registered location in Houston, and leaving it with a medical assistant, who provided a signed receipt for the Order. Affidavit of DI, at 1. On November 5, 2015, Respondent, through his counsel, requested a hearing on the allegations of the Show Cause Order.¹ The matter was then placed on the docket of the Office of Administrative Law Judges, and

¹ While Respondent's request was untimely, Respondent's counsel subsequently filed a motion which established that his secretary had attempted to file the hearing request by UPS overnight delivery, but had provided an incorrect address. DEA has previously held that this type of inadvertence may establish "good cause" to excuse an untimely hearing request, at least when the party promptly moves to rectify the omission. *Tony Bui*, 75 FR 49979, 49980 (2010).