

Cambridge, UK; Clear Scientific, LLC, Cambridge, MA; Cornerstone Government Affairs, Inc., Washington, DC; DePuy Synthes Products, Inc., Raynham, MA; Dignitas Technologies, Orlando, FL; ECM Therapeutics, Inc., Warrendale, PA; ElMindA Ltd., Herzliya, Israel; Family Health International DBA FHI 360, Durham, NC; Hypatia Project, Reston, VA; Institutes for Behavior Resources, Inc. (IBR), Baltimore, MD; Integrated MicroSciences, LLC, Frederick, MD; J. Craig Venter Institute (JCVI), Rockville, MD; KaloCyte, Inc., St. Louis, MO; Lieber Institute, Inc., Baltimore, MD; NanoDirect, LLC, Baltimore, MD; OXYVITA, Inc., Middletown, NY; Parsons Government Services, Inc., Pasadena, CA; Protocentral, Inc., Woburn, MA; Q2Pharma, Haifa, Israel; RAIN Scientific, Inc., Asheville, NC; Rehat, LLC, Pittsburgh, PA; Research Foundation for Mental Hygiene, Inc. (NYSPI), New York, NY; Responde2 Corporation, Mountain View, CA; Saint Barnabas Medical Center (SBMC), Livingston, NJ; San Diego Blood Bank, San Diego, CA; SmartMD Systems, Inc., Manchester Center, VT; Sonica LLC, Evanston, IL; Spire, San Francisco, CA; Syracuse University, Syracuse, NY; The Charles Stark Draper Laboratory, Inc. (Draper), Cambridge, MA; The University of Arizona, Defense and Security Research Institute (DSRI), Tucson, AZ; Trauma Insight, LLC, San Antonio, TX; Trustees of Boston University, Boston, MA; University of Central Florida Research Foundation, Inc., Orlando, FL; University of Houston—Cullen College of Engineering, Houston, TX; Vivonics, Inc., Bedford, MA; and Washington State University, Pullman, WA; 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 MTEC intends to file additional written notifications disclosing all changes in membership.

On May 9, 2014, MTEC 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 9, 2014 (79 FR 32999).

The last notification was filed with the Department on May 3, 2018. A notice was published in the **Federal**

Register pursuant to Section 6(b) of the Act on June 19, 2018 (83 FR 28448).

Suzanne Morris,
*Chief, Premerger and Division Statistics Unit
Antitrust Division.*

[FR Doc. 2018–23989 Filed 11–1–18; 8:45 am]

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DEPARTMENT OF JUSTICE

Drug Enforcement Administration

[Docket No. DEA–392]

Bulk Manufacturer of Controlled Substances Registration

ACTION: Notice of registration.

SUMMARY: The registrants listed below have applied for and been granted registration by the Drug Enforcement Administration (DEA) as bulk manufacturers of various classes of schedule I and II controlled substances.

SUPPLEMENTARY INFORMATION: The companies listed below applied to be registered as bulk manufacturers of various basic classes of controlled substances. Information on previously published notices is listed in the table below. No comments or objections were submitted for these notices.

Company	FR citation	Published
Chemtos, LLC.	83 FR 37520	August 1, 2018.
Johnson Matthey Inc.	83 FR 34880	July 23, 2018.
AMRI Rensselaer, Inc.	83 FR 38179	August 3, 2018.

The DEA has considered the factors in 21 U.S.C. 823(a) and determined that the registration of these registrants to manufacture the applicable 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 each of the company’s maintenance of effective controls against diversion by inspecting and testing each company’s physical security systems, verifying each company’s compliance with state and local laws, and reviewing each company’s background and history.

Therefore, pursuant to 21 U.S.C. 823(a), and in accordance with 21 CFR 1301.33, the DEA has granted a registration as a bulk manufacturer to the above listed companies.

Dated: October 24, 2018.

John J. Martin,

Assistant Administrator.

[FR Doc. 2018–24005 Filed 11–1–18; 8:45 am]

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DEPARTMENT OF JUSTICE

Drug Enforcement Administration

[Docket No. DEA–392]

Bulk Manufacturer of Controlled Substances Application: Janssen Pharmaceuticals, 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 on or before January 2, 2019.

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

SUPPLEMENTARY INFORMATION: The Attorney General has delegated his 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 Assistant Administrator of the DEA Diversion Control Division (“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 September 13, 2018, Janssen Pharmaceuticals, Inc., Buildings 1–5 & 7–14, 1440 Olympic Drive, Athens, Georgia 30601 applied to be registered as a bulk manufacturer of the following basic classes of controlled substances:

Controlled substance	Drug code	Schedule
Methylphenidate	1724	II
Hydromorphone	9150	II
Hydrocodone	9193	II
Oripavine	9330	II
Thebaine	9333	II
Tapentadol	9780	II

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

Dated: October 24, 2018.

John J. Martin,

Assistant Administrator.

[FR Doc. 2018-24006 Filed 11-1-18; 8:45 am]

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DEPARTMENT OF JUSTICE

Notice of Lodging of Proposed Amended Consent Decree Under the Clean Air Act

On October 25, 2018, the Department of Justice lodged a proposed Amended Consent Decree with the United States District Court of the Virgin Islands in the lawsuit entitled *United States of America v. Virgin Islands Water and Power Authority*, Civil Action No. 3:14-cv-00086.

The original Consent Decree resolved the Clean Air Act violations as alleged in the Complaint filed by the United States on October 30, 2014. The violations alleged in the Complaint with respect to VIWAPA's St. Thomas facility include VIWAPA's failure to properly operate and/or maintain its water injection systems on its gas turbine units, failure to operate in compliance with NO_x, sulfuric acid mist, particulate matter and VOC emission limits, failure to operate in compliance with opacity limits, failure to perform required audits and maintain required quality data availability, failure to properly operate and calibrate the continuous emission monitoring systems (CEMS) for NO_x and CO, failure to conduct stack testing every 30 months, and failure to properly report non-compliance. The violations alleged in the Complaint with respect to VIWAPA's St. John facility concern VIWAPA's failure to comply with the RICE NESHAP regulations, failure to timely submit a Title V renewal application and operation without a Title V permit, and failure to conduct stack testing every 30 months.

The Consent Decree, entered by the Court on September 30, 2016, requires VIWAPA to generate a high percentage of its KWh from liquid propane gas or liquid natural gas and renewables, to implement a spare parts inventory program, to control NO_x emissions through improved operation of its water injection system, to maintain and operate continuous emissions monitoring systems on specified units, to operate a video camera system for visible emissions, to perform stack testing, and to conduct targeted self-audits and third party audits given its long term compliance problems. The

Consent Decree also required a \$1,300,000 penalty, which VIWAPA has paid. The proposed Amended Consent Decree makes certain changes to the Consent Decree, including: Updating references to current operating units; adding new units called reciprocating internal combustion engines to the requirements of Paragraph 13 and any requirements associated with the requirements of Paragraph 13; updating aspects of the Consent Decree that have become outdated and are no longer relevant to its enforcement; addressing the current status of the St. John Unit; edits to Paragraph 21 regarding the Atomizer on Unit 14; and adding a date certain for the performance of a stack test.

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 for the Environmental and Natural Resources Division, and should refer to *United States v. Virgin Islands Water and Power Authority*, DOJ Ref. # 90-5-2-1-10424. All comments must be submitted no later than thirty days after the publication date of this notice. Comments may be submitted either by email or by mail:

<i>To submit comments:</i>	<i>Send them to:</i>
By e-mail	<i>pubcomment-ees.enrd@usdoj.gov.</i>
By mail	Assistant Attorney General, U.S. DOJ—ENRD, P.O. Box 7611, Washington, DC 20044-7611.

During the public comment period, the Consent Decree may be examined and downloaded at this Justice Department website: <http://www.justice.gov/enrd/consent-decrees>. We will provide a paper copy of the Consent Decree upon written request and payment of reproduction costs. Please mail your request and payment to: Consent Decree Library, U.S. DOJ—ENRD, P.O. Box 7611, Washington, DC 20044-7611.

Please enclose a check or money order for \$22.75 (25 cents per page reproduction cost) payable to the United States Treasury.

Robert Maher,

Assistant Section Chief, Environmental Enforcement Section, Environment & Natural Resources Division.

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DEPARTMENT OF JUSTICE

U.S. Marshals Service

[OMB Number 1105-0096]

Agency Information Collection Activities; Proposed eCollection eComments Requested; Extension With No Changes, of a Previously Approved Collection; Sequestered Juror Information Form

AGENCY: U.S. Marshals Service, Department of Justice.

ACTION: 30-Day notice.

SUMMARY: The Department of Justice (DOJ), U.S. Marshals Service (USMS), will submit the following information collection request to the Office of Management and Budget (OMB) for review and approval in accordance with the Paperwork Reduction Act of 1995. The proposed information collection was previously published in the **Federal Register** on August 29, 2018, allowing for a 60-day comment period.

DATES: Comments are encouraged and will be accepted for an additional 30 days until December 3, 2018.

FOR FURTHER INFORMATION CONTACT: If you have additional comments, particularly with respect to the estimated public burden or associated response time, have suggestions, need a copy of the proposed information collection instrument with instructions, or desire any other additional information, please contact Nicole Timmons either by mail at CG-3, 10th Floor, Washington, DC 20530-0001, by email at Nicole.Timmons@usdoj.gov, or by telephone at 202-236-2646. Written comments and/or suggestions can also be directed to the Office of Management and Budget, Office of Information and Regulatory Affairs, Attention Department of Justice Desk Officer, Washington, DC 20503 or sent to OIRA_submissions@omb.eop.gov.

SUPPLEMENTARY INFORMATION: Written comments and suggestions from the public and affected agencies concerning the proposed collection of information are encouraged. Your comments should address one or more of the following four points:

- Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility;
- Evaluate the accuracy of the agency's estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used;