VerDate Mar<15>2010 21:38 Jun 28, 2013 Jkt 229001 PO 00000 Frm 00091 Fmt 4703 Sfmt 4703 E:\FR\FM\01JYN1.SGM 01JYN1

Federal Register

Vol. 78, No. 126 / Monday, July 1, 2013 / Notices

39341

Pharmaceutical, Inc., 2110 E. Galbraith Road, Cincinnati, Ohio 45237, made application by renewal to the Drug Enforcement Administration (DEA) to be registered as a bulk manufacturer of Gamma Hydroxybutyric Acid (2010), a basic class of controlled substance listed in schedule 1.

The company plans to manufacture the listed controlled substance for clinical trials and distribution 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 Patheon Pharmaceuticals, Inc., to manufacture the listed basic class of controlled substance is consistent with the public interest at this time. DEA has investigated Patheon Pharmaceuticals, 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(a), 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.

Dated: June 18, 2013.

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


DEPARTMENT OF JUSTICE

Federal Bureau of Investigation

[OMB Number 1110–NEW]

Agency Information Collection Activities: Proposed Collection, Comments Requested: Notice of Collection of Information Relative to Customer Service Satisfaction

ACTION: 30-day Notice.

The Department of Justice, Federal Bureau of Investigation (FBI), National Center for the Analysis of Violent Crime (NCAVC), will be submitting the following information collection request to the Office of Management and Budget (OMB) for review and clearance in accordance with established review procedures of the Paperwork Reduction Act of 1995. The proposed information collection is published to obtain comments from the public and affected agencies. This proposed information collection was previously published in the Federal Register Volume 78, Number 72, page 22332, on April 15, 2013, allowing for a 60 day comment period.

The purpose of this notice is to allow for an additional 30 days for public comment until July 31, 2013. This process is conducted in accordance with 5 CFR 1320.10.

ADDRESSES: All comments, suggestions, or questions regarding additional information, to include obtaining a copy of the proposed information collection instrument with instructions, should be directed to Yvonne Muirhead, Federal Bureau of Investigation, NCAVC, Critical Incident Response Group, FBI Academy, 1 Range Road, Quantico, Virginia, 22135.

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

(1) 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;

(2) Evaluate the accuracy of the agency's estimates of the burden of the proposed collection of information, including the validity of the methodology and assumptions used;

(3) Enhance the quality, utility, and clarity of the information to be collected; and

(4) Minimize the burden of the collection of information on those who are to respond, including through the use of appropriate automated, electronic, mechanical, or other technological collection techniques of other forms of information technology, e.g., permitting electronic submission of responses.

Overview of this information collection:

(1) Type of information collection: Customer satisfaction ratings regarding the quality and value of the FBI's NCAVC services.


(3) There is no agency form number applicable to this survey.

(4) The survey will be distributed to state, local and tribal law enforcement agencies to which the NCAVC has provided investigative assistance. The survey is being proposed as a means to assess the effectiveness and efficiency with which the NCAVC serves these agencies in the execution of their missions. The survey will query respondents as to the agencies' satisfaction with NCAVC services, and concrete achievements which were furthered via NCAVC services.

(5) Time burden anticipated with this collection: It is estimated that 100 respondents per calendar year will be contacted to complete a survey consisting of 11 questions. An approximate non-response rate of 50% is anticipated. It is estimated that a burden of approximately three to five minutes, or .05 to .08 hours, will be cast upon each respondent to complete the survey, with a total estimate of five to 8.3 hours in a calendar year for all respondents combined, if all respondents complete a survey. If the expected non-response rate of 50% holds true, then the combined burden estimate drops to approximately 2.5 to 4.2 hours per calendar year. The NCAVC estimates little to no variability within this time estimate based upon on individualized data retrieval systems, availability of requested data, and other variables, because this survey is intended to assess customer satisfaction rather than generate empirical data.

Methodology: The survey will be distributed and collected electronically, via electronic mail communication.

Contact: If additional information is required contact: Jerri Murray, Department Clearance Officer, Policy and Planning Staff, Justice Management Division, United States Department of Justice, Two Constitution Square, 145 N Street NE., Room 1407B, Washington, DC 20530.

Dated: June 25, 2013.

Jerri Murray,
Department Clearance Officer for PRA, United States Department of Justice.

[FR Doc. 2013–15566 Filed 6–28–13; 8:45 am]BILLING CODE 4410–02–P

NATIONAL AERONAUTICS AND SPACE ADMINISTRATION

[Notice: 13–070]

NASA Advisory Council; Science Committee; Planetary Science Subcommittee; Meeting.

AGENCY: National Aeronautics and Space Administration.

ACTION: Notice of meeting.

SUMMARY: In accordance with the Federal Advisory Committee Act, as amended, the National Aeronautics and Space Administration (NASA) announces a meeting of the Planetary Science Subcommittee of the NASA Advisory Council (NAC). This
Subcommittee reports to the Science Committee of the NAC. The Meeting will be held via Teleconference and WebEx for the purpose of soliciting, from the scientific community and other persons, scientific and technical information relevant to program planning.

DATES: Friday, July 19, 2013, 9:00 a.m. to 12:00 noon, local time.

ADDRESSES: This meeting will take place telephonically and by WebEx. Any interested person may call the USA toll free conference call number 800–857–7040, pass code PSS, to participate in this meeting by telephone. The WebEx link is https://nasa.webex.com/, meeting number 994 987 970, and password PSS@Jul19.

FOR FURTHER INFORMATION CONTACT: Ms. Marian Norris, Science Mission Directorate, NASA Headquarters, Washington, DC 20546, (202) 358–4452, fax (202) or mnorris@nasa.gov.

SUPPLEMENTARY INFORMATION: The agenda for the meeting includes the following topics:

—Briefing from the Mars Exploration Program Regarding the Recommendations of the Mars 2020 Science Definition Team

It is imperative that the meeting be held on this date to accommodate the scheduling priorities of the key participants.

Susan M. Burch, Acting Advisory Committee Management Officer, National Aeronautics and Space Administration.

[FR Doc. 2013–15677 Filed 6–28–13; 8:45 am]

NUCLEAR REGULATORY COMMISSION

[Docket No. 50–608; NRC–2013–0053]

SHINE Medical Technologies, Inc.

AGENCY: Nuclear Regulatory Commission.

ACTION: Notice; acceptance for docketing.

SUMMARY: The NRC staff has determined that the partial application for a construction permit, submitted by SHINE Medical Technologies, Inc., is acceptable for docketing.

ADDRESSES: Please refer to Docket ID NRC–2013–0053 when contacting the NRC about the availability of information regarding this document. You may access information related to this document, which the NRC possesses and is publicly available, using any of the following methods:

- Federal Rulemaking Web site: Go to http://www.regulations.gov and search for Docket ID NRC–2013–0053. Address questions about NRC dockets to Carol Gallagher; telephone: 301–492–3668; email: Carol.Gallagher@nrc.gov. For technical questions, contact the individual(s) listed in the FOR FURTHER INFORMATION CONTACT section of this document.
- NRC’s Agencywide Documents Access and Management System (ADAMS): You may access publicly available documents online in the NRC Library at http://www.nrc.gov/reading-rm/adams.html. To begin the search, select “ADAMS Public Documents” and then select “Begin Web-based ADAMS Search.” For problems with ADAMS, please contact the NRC’s Public Document Room (PDR) reference staff at 1–800–397–4209, 301–415–4737, or by email to pdr.resource@nrc.gov. The application is available in ADAMS under accession number ML130880226.
- NRC’s PDR: You may examine and purchase copies of public documents at the NRC’s PDR, Room O1–F21, One White Flint North, 11555 Rockville Pike, Rockville, Maryland 20852.


SUPPLEMENTARY INFORMATION: On March 26, 2013, SHINE Medical Technologies (SHINE) filed with the U.S. Nuclear Regulatory Commission (NRC), pursuant to Section 103 of the Atomic Energy Act and part 50 of Title 10 of the Code of Federal Regulations (10 CFR), a portion of an application for a construction permit for a medical radioisotope production facility in Janesville, Wisconsin (SMT–2013–012, NRC’s ADAMS Accession No. ML13088A192). A notice of receipt and availability of this application was previously published in the Federal Register (78 FR 29390) on May 20, 2013.

An exemption from certain requirements of 10 CFR 2.101(a)(5) granted by the Commission on March 20, 2013 (ADAMS Accession No. ML13072B195), in response to a letter from SHINE dated February 18, 2013 (ADAMS Accession No. ML13051A007), allowed for SHINE to submit its construction permit application in two parts. Specifically, the exemption allowed SHINE to submit a portion of its application for a construction permit up to six months prior to the remainder of the application regardless of whether or not an environmental impact statement or a supplement to an environmental impact statement is prepared during the review of its application. The first part of SHINE’s construction permit application consisted of the following information:

- The description and safety assessment of the site required by 10 CFR 50.34(a)(1)
- The environmental report required by 10 CFR 50.30(f)
- The filing fee information required by 10 CFR 50.30(e) and 10 CFR 170.21
- The general information required by 10 CFR 50.33
- The agreement limiting access to classified information required by 10 CFR 50.37

The NRC staff has determined that SHINE has submitted the information listed above in accordance with 10 CFR 2.101(a)(5), and that the partial application is acceptable for docketing. The docket number established for SHINE is 50–608.

The NRC staff will perform a detailed technical review of the partial construction permit application. Docketing of the partial construction permit application does not preclude the NRC from requesting additional information from the applicant as the review proceeds, nor does it predict whether the Commission will grant or deny the application. The NRC staff will also perform an acceptance review of the second and final part of the construction permit application when it is tendered. As stated in SHINE’s March 26, 2013, letter, the second and final part of SHINE’s application for a construction permit will contain the remainder of the preliminary safety analysis report required by 10 CFR 50.34(a) and will be submitted in accordance with the requirements of 10 CFR 2.101(a)(5). If, after completion of the acceptance review of the full construction permit application, the full construction permit application is found acceptable for docketing, the Commission will conduct a hearing in accordance with Subpart L, “Informal Hearing Procedures for NRC Adjudications.” of 10 CFR Part 2 and will receive a report on the construction permit application from the Advisory Committee on Reactor Safeguards consistent with 10 CFR 50.58, “Hearings and report of the Advisory Committee on Reactor Safeguards.” The Commission will announce in a future Federal Register notice, the opportunity to petition for leave to intervene in the hearing required for this application by 10 CFR 50.58, as well as the time and