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 registration 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 4, 2010, 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:

<table>
<thead>
<tr>
<th>Drug</th>
<th>Schedule</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxydodeine (9143)</td>
<td>II</td>
</tr>
<tr>
<td>Hydromorphone (9150)</td>
<td>II</td>
</tr>
<tr>
<td>Fentanyl (9801)</td>
<td>II</td>
</tr>
</tbody>
</table>

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 26, 2010.

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.


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

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

Manufacturer of Controlled Substances; Notice of Application

Pursuant to § 1301.33(a) of Title 21 of the Code of Federal Regulations (CFR), this is notice that on March 5, 2010, Stepan Company, Natural Products Dept., 100 W. Hunter Avenue, Maywood, New Jersey 07607, made application by renewal to the Drug Enforcement Administration (DEA) as a bulk manufacturer of the basic classes of controlled substances listed in schedule II:

<table>
<thead>
<tr>
<th>Drug</th>
<th>Schedule</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cocaine (9041)</td>
<td>II</td>
</tr>
<tr>
<td>Benzoylecgonine (9180)</td>
<td>II</td>
</tr>
</tbody>
</table>

The company plans to manufacture the listed controlled substances in bulk for distribution to its 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 25, 2010.


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

DEPARTMENT OF JUSTICE

Federal Bureau of Investigation

Meeting of the CJIS Advisory Policy Board

AGENCY: Federal Bureau of Investigation (FBI), Department of Justice.

ACTION: Meeting notice.

SUMMARY: The purpose of this notice is to announce the meeting of the Criminal Justice Information Services (CJIS) Advisory Policy Board (APB). The CJIS APB is a federal advisory committee established pursuant to the Federal Advisory Committee Act (FACA). This meeting announcement is being published as required by Section 10 of the FACA.

The CJIS APB is responsible for reviewing policy issues and appropriate technical and operational issues related to the programs administered by the FBI’s CJIS Division, and thereafter, making appropriate recommendations to the FBI Director. The programs administered by the CJIS Division are the Integrated Automated Fingerprint Identification System, Interstate Identification Index, Law Enforcement Online, National Crime Information Center, National Instant Criminal Background Check System, National Incident-Based Reporting System, Law Enforcement National Data Exchange, and Uniform Crime Reporting.

The meeting will be open to the public on a first-come, first-seated basis. Any member of the public wishing to file a written statement concerning the CJIS Division’s programs or wishing to address this session should notify Senior CJIS Advisor Roy G. Weise at (304) 625–2730 at least 24 hours prior to the start of the session. The notification should contain the requestor’s name, corporate designation, and consumer affiliation or government designation along with a short statement describing the topic to be addressed and the time needed for the presentation. A requestor will ordinarily be allowed no more than 15 minutes to present a topic.

DATES AND TIMES: The APB will meet in open session from 8:30 a.m. until 5 p.m., on June 9–10, 2010.

ADDRESSES: The meeting will take place at The Renaissance, A Columbus Hotel, 50 North Third Street, Columbus, Ohio 43215, telephone (614) 233–7519.

FOR FURTHER INFORMATION CONTACT: Inquiries may be addressed to Mrs. Margery E. Broadwater; Management and Program Analyst; Advisory Groups Management Unit, Law Enforcement Support Section; FBI CJIS Division; Module C3, 1000 Custer Hollow Road,
Clarksburg, West Virginia 26306–0149; telephone (304) 625–2446, facsimile (304) 625–5090.

Dated: April 9, 2010.

Roy G. Weise,
Senior OIS Advisor, Criminal Justice Information, Services Division, Federal Bureau of Investigation.

[FR Doc. 2010–9444 Filed 4–23–10; 8:45 am]
BILLING CODE 4410–02–M

DEPARTMENT OF LABOR
Occupational Safety and Health Administration
[Docket No. OSHA–2009–0043]

Access to Employee Exposure and Medical Records; Extension of the Office of Management and Budget’s (OMB) Approval of Information Collection (Paperwork) Requirements

AGENCY: Occupational Safety and Health Administration (OSHA), Labor.

ACTION: Request for public comment.

SUMMARY: OSHA solicits public comments concerning its proposal to extend OMB approval of the information collection requirements contained in its Regulation on Access to Employee Exposure and Medical Records (29 CFR 1910.1020).

DATES: Comments must be submitted (postmarked or received) by June 25, 2010.

ADDRESSES:
Electronically: You may submit comments and attachments electronically at http://www.regulations.gov, which is the Federal eRulemaking Portal. Follow the instructions online for submitting comments.
Facsimile: If your comments, including attachments, are not longer than 10 pages, you may fax them to the OSHA Docket Office at (202) 693–1648.
Mail, hand delivery, express mail, messenger, or courier service: When using this method, you must submit three copies of your comments and attachments to the OSHA Docket Office, Docket No. OSHA–2009–0043, U.S. Department of Labor, Occupational Safety and Health Administration, Room N–2625, 200 Constitution Avenue, NW., Washington, DC 20210; telephone: (202) 693–2222.

SUPPLEMENTARY INFORMATION:

I. Background

The Department of Labor, as part of its continuing effort to reduce paperwork and respondent (i.e., employer) burden, conducts a preclearance consultation program to provide the public with an opportunity to comment on proposed and continuing information collection requirements in accordance with the Paperwork Reduction Act of 1995 (PRA–95) (44 U.S.C. 3506(c)(2)(A)). This program ensures that information is in the desired format, reporting burden (time and cost) is minimal, collection instruments are clearly understood, and OSHA’s estimate of the information collection burden is accurate. The Occupational Safety and Health Act of 1970 (the OSH Act) (29 U.S.C. 651 et seq.) authorizes information collection by employers as necessary or appropriate for enforcement of the OSH Act or for developing information regarding the causes and prevention of occupational injuries, illnesses, and accidents (29 U.S.C. 657). The Act also requires that OSHA obtain such information with minimum burden upon employers, especially those operating small businesses, and to reduce to the maximum extent feasible unnecessary duplication of efforts in obtaining information (29 U.S.C. 657).

Under the authority granted by the Act, OSHA published a health regulation governing access to worker exposure monitoring data and medical records. This regulation does not require employers to collect any information or to establish any new systems of records. Rather, it requires that employers provide workers, their designated representatives, and OSHA with access to worker exposure monitoring and medical records, and any analyses resulting from these records that employers must maintain under OSHA’s toxic chemical and harmful physical agent standards. In this regard, the regulation specifies requirements for record access, record retention, worker information, trade secret management, and record transfer. Accordingly, the Agency attributes the burden hours and costs associated with exposure monitoring and measurement, medical surveillance, and the other activities required to generate the data governed by the regulation to the health standards that specify these activities; therefore, OSHA did not include these burden hours and costs in the ICR.

Access to exposure and medical information enables workers and their designated representatives to become directly involved in identifying and controlling occupational health hazards, as well as managing and preventing occupationally-related health impairment and disease. Providing the Agency with access to the records permits it to ascertain whether or not employers are complying with the regulation, as well as the recordkeeping requirements of its other health standards; therefore, OSHA access provides additional assurance that workers and their designated representative are able to obtain the data they need to conduct their analyses.

II. Special Issues for Comment

OSHA has a particular interest in comments on the following issues:
• Whether the proposed information collection requirements are necessary for the proper performance of the Agency’s functions to protect workers, including whether the information is useful;
• The accuracy of OSHA’s estimate of the burden (time and costs) of the information collection requirement, including the validity of the methodology and assumptions used;
• The quality, utility, and clarity of the information collected; and
• Ways to minimize the burden on employers who must comply:

Instructions: All submissions must include the Agency name and OSHA docket number for the Information Collection Request (ICR) (OSHA–2009–0043). All comments, including any personal information you provide, are placed in the public docket without change, and may be made available online at http://www.regulations.gov. For further information on submitting comments see the “Public Participation” heading in the section of this notice titled “Supplementary Information.”

Docket: To read or download comments or other material in the docket, go to http://www.regulations.gov or the OSHA Docket Office at the address above. All documents in the docket (including this Federal Register notice) are listed in the http://www.regulations.gov index; however, some information (e.g., copyrighted material) is not publicly available to read or download through the Web site. All submissions, including copyrighted material, are available for inspection and copying at the OSHA Docket Office. You may contact Todd Owen at the address below to obtain a copy of the ICR.

FOR FURTHER INFORMATION CONTACT:
