XV. Separate Document(s)

• Application for Federal Assistance SF–424 Form.

• Project Narrative Attachment Form (This form includes the Project Narrative, Budget, Tribal Resolution, and Critical Information page).

XVI. Paperwork Reduction Act

The information collection requirements contained in SF–424, Application for Federal Assistance have been reviewed and approved by the Office of Management and Budget (OMB) under the Paperwork Reduction Act, 44 U.S.C. 3504(h). The OMB control number is 4040–0004. The authorization expires on December 31, 2022. An agency may not conduct or sponsor, and you are not required to respond to, any information collection that does not display a currently valid OMB Control Number.

XVII. Authority

This is a discretionary grant program authorized under the Snyder Act (25 U.S.C. 13) and the Further Consolidated Appropriations Act, 2020 (Pub. L. 116–94). The Snyder Act authorizes the BIA to expend such moneys as Congress may appropriate for the benefit, care, and assistance of Indians for the purposes listed in the Act. LLGP grants facilitate one of the purposes listed in the Snyder Act: “General support and civilization, including education.” The Further Consolidated Appropriations Act, 2020, authorizes the BIA to “carry out the operation of Indian programs by direct expenditures, contracts, cooperative agreements, compacts, and grants, either directly or in cooperation with States and other organizations.” Further, the Conference Report specifies, the agreement continues $3,000,000 for grants to federally recognized Indian Tribes and Tribal organizations to provide native language instruction and immersion programs to Native students not enrolled at BIE schools, including those Tribes and organizations in states without Bureau-funded schools.

Bryan Newland,
Assistant Secretary—Indian Affairs.

[FR Doc. 2021–26401 Filed 12–3–21; 8:45 am]
BILLING CODE 4337–15–P

INTERNATIONAL TRADE COMMISSION

[Investigation Nos. 701–TA–534–537 and 731–TA–1274–1278 (Review)]

Certain Corrosion-Resistant Steel Products From China, India, Italy, Korea, and Taiwan; Notice of Commission Determination To Conduct Full Five-Year Reviews


ACTION: Notice.

SUMMARY: The Commission hereby gives notice that it will proceed with full reviews pursuant to the Tariff Act of 1930 to determine whether revocation of the countervailing duty orders on certain corrosion-resistant steel products from China, India, Italy, and Korea and the antidumping duty orders on certain corrosion-resistant steel products from China, India, Italy, Korea, and Taiwan would be likely to lead to continuation or recurrence of material injury within a reasonably foreseeable time. A schedule for the reviews will be established and announced at a later date.

DATES: September 7, 2021.


General information concerning the Commission may also be obtained by accessing its internet server (https://www.usitc.gov). The public record for these reviews may be viewed on the Commission’s electronic docket (EDIS) at https://edis.usitc.gov.

For further information concerning the conduct of these reviews and rules of general application, consult the Commission’s Rules of Practice and Procedure, part 201, subparts A through E (19 CFR part 201), and part 207, subparts A, D, E, and F (19 CFR part 207).

SUPPLEMENTARY INFORMATION: On September 7, 2021, the Commission determined that it should proceed to full reviews in the subject five-year reviews pursuant to section 751(c) of the Tariff Act of 1930 (19 U.S.C. 1675(c)). The Commission found that the domestic interested party group response and the respondent interested party group response from Taiwan to its notice of institution (86 FR 29283, June 1, 2021) were adequate, and determined to conduct a full review of the order on imports from Taiwan. The Commission also found that the respondent interested party group responses from China, India, Italy, and Korea were inadequate but determined to conduct full reviews of the orders on certain corrosion-resistant steel products from those countries in order to promote administrative efficiency in light of its determination to conduct a full review of the order with respect to Taiwan. A record of the Commissioners’ votes, the

Data Availability

• Applicability. The Department of the Interior is committed to basing its decisions on the best available science and providing the American people with enough information to thoughtfully and substantively evaluate the data, methodology, and analysis used by the Department to inform its decisions.

• Use of Data. The regulations at 2 CFR 200.315 apply to data produced under a Federal award, including the provision that the Federal Government has the right to obtain, reproduce, publish, or otherwise use the data produced under a Federal award as well as authorize others to receive, reproduce, publish, or otherwise use such data for Federal purposes.

• Availability of Data. The recipient shall make the data produced under this award and any subaward(s) available to the Government for public release, consistent with applicable law, to allow the Government for public release, and publication, or otherwise use the data produced under a Federal award as well as authorize others to receive, reproduce, publish, or otherwise use such data for Federal purposes.

• Methodology, including models, used to gather and analyze data.

XIV. Questions and Requests for OIED Assistance

OIED staff may provide technical assistance, upon written request by an applicant. The request must clearly identify the type of assistance sought. Technical assistance does not include funding to prepare a grant proposal, grant writing assistance, or pre-determinations as to the likelihood that a proposal will be awarded. The applicant is solely responsible for preparing its grant proposal. Technical assistance may include clarifying application requirements, and registration information for SAM or ASAP.

XV. Separate Document(s)

• Application for Federal Assistance SF–424 Form.

• Project Narrative Attachment Form (This form includes the Project Narrative, Budget, Tribal Resolution, and Critical Information page).
DEPARTMENT OF JUSTICE

Drug Enforcement Administration

[Docket No. DEA–935]

Importer of Controlled Substances Application: Johnson Matthey Inc.

AGENCY: Drug Enforcement Administration, Justice.

ACTION: Notice of application.

SUMMARY: Johnson Matthey Inc., has applied to be registered as an importer of basic class(es) of controlled substance(s). Refer to SUPPLEMENTARY INFORMATION listed below for further drug information.

DATES: Registered bulk manufacturers of the affected basic class(es), and applicants therefore, may file written comments on or objections to the issuance of the proposed registration on or before January 5, 2022. Such persons may also file a written request for a hearing on the application on or before January 5, 2022.

ADDRESSES: Written comments should be sent to: Drug Enforcement Administration, Attention: DEA Federal Register Representative/DPW, 8701 Morrissette Drive, Springfield, Virginia 22152. All requests for a hearing must be sent to: Drug Enforcement Administration, Attn: Hearing Clerk/OALJ, 8701 Morrissette Drive, Springfield, Virginia 22152. All request for a hearing also should be sent to: (1) Drug Enforcement Administration, Attn: Hearing Clerk/OALJ, 8701 Morrissette Drive, Springfield, Virginia 22152; and (2) Drug Enforcement Administration, Attn: DEA Federal Register Representative/DPW, 8701 Morrissette Drive, Springfield, Virginia 22152.

SUPPLEMENTARY INFORMATION: In accordance with 21 CFR 1301.34(a), this is notice that on November 2, 2021, Johnson Matthey Inc., 2003 Nolte Drive, West Deptford, New Jersey 08066–0727, applied to be registered as an importer of the following basic class(es) of controlled substance(s):

<table>
<thead>
<tr>
<th>Controlled substance</th>
<th>Drug code</th>
<th>Schedule</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coca Leaves</td>
<td>9040</td>
<td>II</td>
</tr>
<tr>
<td>Thebaine</td>
<td>9333</td>
<td>II</td>
</tr>
<tr>
<td>Opium, raw</td>
<td>9600</td>
<td>II</td>
</tr>
<tr>
<td>Noroxymorphine</td>
<td>9668</td>
<td>II</td>
</tr>
<tr>
<td>Poppy Straw Concentrate</td>
<td>9670</td>
<td>II</td>
</tr>
<tr>
<td>Fentanyl</td>
<td>9801</td>
<td>II</td>
</tr>
</tbody>
</table>

The company plans to import Coca Leaves (9040), Opium raw (9600) and Poppy Straw Concentrate (9670) in order to bulk manufacture Active Pharmaceutical Ingredients (API) for distribution to its customers. The company plans to also import Thebaine (9333), Noroxymorphine (9668) and Fentanyl (9801) to use as analytical reference standards, both internally and to be sold to their customers to support testing of Johnson Matthey Inc.’s API’s only.

Approval of permit applications will occur only when the registrant’s business activity is consistent with what is authorized under 21 U.S.C. 952(a)(2). Authorization will not extend to the import of the Food and Drug Administration-approved or non-approved finished dosage forms for commercial sale.

Brian S. Besser,
Acting Assistant Administrator.

BILLCODE P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

[Docket No. DEA–936]

Importer of Controlled Substances Application: Fisher Clinical Services, Inc.

AGENCY: Drug Enforcement Administration, Justice.

ACTION: Notice of application.

SUMMARY: Fisher Clinical Services, Inc., has applied to be registered as an importer of basic class(es) of controlled substance(s). Refer to SUPPLEMENTARY INFORMATION listed below for further drug information.

DATES: Registered bulk manufacturers of the affected basic class(es), and applicants therefore, may file written comments on or objections to the issuance of the proposed registration on or before January 5, 2022. Such persons may also file a written request for a hearing on the application on or before January 5, 2022.

ADDRESSES: Written comments should be sent to: Drug Enforcement Administration, Attention: DEA Federal Register Representative/DPW, 8701 Morrissette Drive, Springfield, Virginia 22152. All requests for a hearing must be sent to: Drug Enforcement Administration, Attn: Hearing Clerk/OALJ, 8701 Morrissette Drive, Springfield, Virginia 22152. All request for a hearing also should be sent to: (1) Drug Enforcement Administration, Attn: Hearing Clerk/OALJ, 8701 Morrissette Drive, Springfield, Virginia 22152; and (2) Drug Enforcement Administration, Attn: DEA Federal Register Representative/DPW, 8701 Morrissette Drive, Springfield, Virginia 22152.

SUPPLEMENTARY INFORMATION: In accordance with 21 CFR 1301.34(a), this is notice that on August 25, 2021, Fisher Clinical Services, Inc., 700A–C Nestle Way, Breinigsville, Pennsylvania 18031–1522, applied to be registered as an importer of the following basic class(es) of controlled substance(s):

<table>
<thead>
<tr>
<th>Controlled substance</th>
<th>Drug code</th>
<th>Schedule</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maruahana Extract</td>
<td>7350</td>
<td>I</td>
</tr>
<tr>
<td>Pelocybin</td>
<td>7437</td>
<td>I</td>
</tr>
<tr>
<td>Methylphenidate</td>
<td>1724</td>
<td>II</td>
</tr>
<tr>
<td>Levorphanol</td>
<td>9220</td>
<td>II</td>
</tr>
<tr>
<td>Noroxymorphine</td>
<td>9668</td>
<td>II</td>
</tr>
<tr>
<td>Tapentadol</td>
<td>9780</td>
<td>II</td>
</tr>
</tbody>
</table>

The company plans to import the listed controlled substances for use in clinical trials only. No other activity for these drug codes is authorized for this registration.

Approval of permit applications will occur only when the registrant’s business activity is consistent with what is authorized under 21 U.S.C. 952(a)(2). Authorization will not extend to the import of Food and Drug Administration-approved or non-approved finished dosage forms for commercial sale.

Brian S. Besser,
Acting Assistant Administrator.

BILLCODE P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

[Docket No. DEA–934]

Importer of Controlled Substances Application: Kinetochem LLC

AGENCY: Drug Enforcement Administration, Justice.

ACTION: Notice of application.