

Dated: September 8, 2004.

Michele M. Leonhart,

Deputy Administrator.

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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)(b) 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 21 CFR 1301.34(a), this is notice that on July 21, 2004, Tocris Cookson, Inc., 16144 Westwoods Business Park, Ellisville, Missouri 63021-4500, made application by renewal to the Drug Enforcement Administration (DEA) for registration as an importer of Tetrahydrocannabinols (7370), a basic class of controlled substance listed in Schedule I.

The company plans to import small quantities of the products for research purposes.

Any manufacturer who is presently, or is applying to be, registered with DEA to manufacture such basic class of controlled substance 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 or requests for hearing may be addressed, in quintuplicate, to the Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration, United States Department of Justice, Washington, DC 20537, Attention: DEA Federal Register Representative (CCD) and must be filed no later than (30 days from publication).

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 a basic class of any controlled substances 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.

Dated: September 16, 2004.

William J. Walker,

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

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DEPARTMENT OF JUSTICE

Drug Enforcement Administration

Value Wholesale Denial of Registration

On September 8, 2003, the Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration (DEA), issued an Order to Show Cause to Value Wholesale (Value) proposing to deny its November 6, 2001, application for DEA Certificate of Registration as a distributor of list I chemicals. The Order to Show Cause alleged that granting Value's application would be inconsistent with the public interest, as that term is used in 21 U.S.C. 823(h) and 824(a). The order also notified Value that should no request for a hearing be filed within 30 days, its hearing right would be deemed waived.

According to the DEA investigative file, the Order to Show Cause was sent by certified mail to Value at its proposed registered location at 15188 Eight Mile Road, Oak Park, Michigan 48237. It was received on September 16, 2003, and DEA has not received a request for a hearing or any other reply from Value or anyone purporting to represent the company in this matter.

Therefore, the Deputy Administrator of DEA, finding that (1) thirty days have passed since delivery of the Order to Show Cause, and (2) no request for a hearing having been received, concludes that Value has waived its hearing right. See *Aqui Enterprises*, 67 FR 12576 (2002). After considering relevant material from the investigative file, the Deputy Administrator now enters her final order without a hearing pursuant to 21 CFR 1309.53(c) and (d) and 1316.67 (2003). The Deputy Administrator finds as follows:

List I chemicals are those that may be used in the manufacture of a controlled substance in violation of the Controlled Substances Act. 21 U.S.C. 802(34); 21 CFR 1310.02(a). Pseudoephedrine and ephedrine are list I chemicals commonly used to illegally manufacture

methamphetamine, a Schedule II controlled substance. Phenylpropanolamine, also a list I chemical, is presently a legitimately manufactured and distributed product used to provide relief of the symptoms resulting from irritation of the sinus, nasal and upper respiratory tract tissues, and is also used for weight control. Phenylpropanolamine is also a precursor chemical used in the illicit manufacture of methamphetamine and amphetamine. Methamphetamine is an extremely potent central nervous system stimulant, and its abuse is an ongoing public health concern in the United States.

The Deputy Administrator's review of the investigative file reveals that an application dated November 6, 2001, was submitted on behalf of Value and signed by its President and only officer, Mr. John Loussia (Mr. Loussia). Value sought registration as a distributor of multiple list I chemicals, including pseudoephedrine (8112) and phenylpropanolamine (1225). There is no evidence in the investigative file that Value has sought to modify its pending application with regard to those two chemicals.

In January 1999, Value originally applied for DEA registration as a distributor of list I chemicals and during a pre-registration investigation, it was determined the company had been buying and selling list I chemical products for a number of years prior to filing this application for registration. However, on February 5, 1999, that application was approved and Value issued DEA Certificate of Registration 004000VHY.

On October 31, 2001, during the course of a regularly scheduled cyclic investigation, it was discovered Value's registration had expired, effective May 31, 2000, without any application for renewal having been filed. Nevertheless, investigators found that the firm had continued to order and sell list I chemical products after its registration had expired. Investigators also discovered Value had not been maintaining adequate or complete records of customer addresses as required by 21 CFR 1310.06. A DEA letter of admonition was issued the company and in reply, Mr. Loussia advised he would be submitting the instant application for registration and not be carrying list I chemical products until its approval.

In connection with the pending application, an on-site pre-registration investigation was conducted in March 2002. Mr. Loussia advised investigators that Value was a full-line wholesaler/distributor of groceries to local food