U.S. DEPARTMENT OF JUSTICE - DRUG ENFORCEMENT ADMINISTRATION

## APPLICATION FOR PERMITTO EXPORT CONTROLLED SUBSTANCES PURSUANT TO SECTION 1003(a), (b), (c) & (d), TITLE III, PL 91-513 (See Reverse for Instructions and Privacy Act Information)

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Drug Enforcement Administration Office of Diversion Control International Drug Unit (ODOI) Washington, D.C. 20537		DATE		
		EXPORTER'S APPLICATION NUMBER		
Application is hereby made pursuar regulations prescribed thereunder f			nport and E	xport Act and the
1. NAME OF CONSIGNEE		2. ADDRESS OF CONSIGNEE		
3. BUSINESS OF CONSIGNEE		4. FOREIGN PORT OF ENTRY (City & Country)		
5.a PORT OF EXPORTATION (City & State of las U.S. Customs port)	5b. NAME OF EXPORTING (Air, Ship)	CARRIER OR VESSEL	5c. APPROX	C. DATE OF EXPORTATION
6. FOREIGN IMPORT LICENSE OR PERMIT FIELD	HEREWITH NO.	DA	ATED	
TO BE EXPORTED (Enter names as shown on labels; OR PREPA numbers and sizes of packages; strength of tablets, as acid, bas		ED SUBSTANCE CONTENT OF D RATION TO BE EXPORTED expr se or alkaloid (Enter name of con	essed trolled	7c. DATE EXPORTED AND ACTUAL QUANTITY (Completed by registrant at time of export)
capsules, etc., CSA Drug Code, and NDC Numb	preparation	stance contained in the drug; compound, or paration.)		DEA PERMIT NO.:
The mask ages to be averaged and labelled in conform	manaa mish 21 C.F.P. Post 2002	and to the heat of my knowledge	a and haliaf e	
The packages to be exported are labeled in conformand maintains a system for the control of these subcountry of import; the substances are to be applied substances for medical or scientific uses within successful substance will be processed within the country of i Convention on Narcotic Drugs, 1961.	stances; the drugs are consigned exclusively to medical or scien	d to a holder of such permits or tific uses within the country of in	licenses as ma mport; there is	by be required under the laws of the an actual need for the controlled
NAME OF EXPORTER		ADDRESS OF EXPORTER		
	ER'S DEA REGISTRATION NO.	SIGNATURE AND TITLE OF PERSON MAKING APPLICATION		
APPROVED EX	olled Substances may KPORT PERMIT NUMBER	not be exported by ma	<mark>ail or parc</mark> TPERMITNUN	el post. IBERISSUED
DEA USE ONLY				

## **INSTRUCTIONS AND INFORMATION, DEA-161**

This application must be completed in triplicate. Original is sent to DEA. See instruction (7) for copies two and three.

- (1) The name and address of the consignee as shown on this application and on the permit to export must correspond with that shown on the foreign import certificate.
- (2) To avoid delays in clearance at the port of export be sure to enter the correct port on this application. A copy of your export permit is sent directly to the District Director of Customs at the port indicated on the application for comparison with the permit presented for clearance of the shipment. The shipment will not clear at any other port without an amendment of the permit indicating a change to that effect.
- (3) The original or an authentic signed and/or notarized copy of the foreign import certificate must accompany this application. If this certificate is needed to accomplish entry of the drug into the country of destination, your request for its return to you should accompany the application.
- (4) Application should be made in the name of the registered legal entity, as shown on the DEA registration certificate, and signed by a responsible authorized official if a corporation, by a partner, or by the person registered as an individual. Only persons registered as exporters may be issued export permits. The registrations of manufacturers, distributors, practitioners, researchers, etc., do not entitle them to export controlled substances.
- (5) Permits will be mailed to the exporter at the address shown at the bottom of the application unless contrary instructions are attached to and made a part of this application.
- (6) Identification of drugs to be exported and the controlled substance content should be entered on the application in the following manner:

7a. NAME AND QUANITY OF DRUG OR PREPARATION TO BE EXPORTED	7B. CONTROLLED SUBSTANCE CONTENT OF DRUG OR PREPARATION TO BE EXPORTED (expressed as acid, base or alkaloid, not salt)		
3 bottles x 100 Secobarbital Sodium capsules (100 mg./capsule)	Secobarbital	27.47 Gm.	
2 boxes x 100 Medperidine HCL ampules (5%, 2ml. ampules)	Meperidine	17.43 G.m	
1 box x 100 Meperidine HCL vials (10%, 20 ml, vials)	Meperidine	174.30 Gm.	
2 x 1 Pt. Meperidine HCL Syrup (50 mg./5ml., pints)	Meperidine	8.24 Gm.	
1 box x 100 gm. Dextroamphetamine Sulfate powder	Dextroamphetamine	73.38 Gm.	
1 bottle x 500 Hydromorphone HCl tablets (4 mg./tablets)	Hydromorphone	1.77 Gm.	

(7) The following information must be entered in block 7c at the time of export: (1) DEA Export Permit Number and (2) actual quantity and date shipped. Copy 2 is sent to DEA, and Copy 3 is retained by the registrant.

## **PRIVACY ACT INFORMATION**

AUTHORITY: Section 1003 of the Controlled Substances Act of 1970 (PL 91-513).

PURPOSE: Control exportation of certain Controlled Substances into the United States.

**ROUTINE USES:** The Controlled Substances Act Registration Records produces special reports as required for statistical analytical purposes. Disclosures of information from this system are made to the following categories of users for the purposes stated:

- A. Other Federal law enforcement and regulatory agencies for law enforcement and regulatory purposes.
- B. State and local law enforcement and regulatory agencies for law enforcement and regulatorypurposes.
- C. Persons registered under the Controlled Substances Act (Public Law 91-513) for the purpose of verifying the registration of customers and practioners.

EFFECT: No permit will be issued.

According to the Paperwork Reduction Act of 1995, no persons are required to respond to a Collection of Information unless it displays a valid OMB control number. The valis OMB control number for this Information Collection is 1117-0004. The time required the complete this information collection is estimated to average 15 minutes per response, including the time to review instructions, search existing data resources, gather the data needed, and complete and review the information collection.