## APPLICATION FOR INDIVIDUAL MANUFACTURING QUOTA FOR A BASIC CLASS OF CONTROLLED SUBSTANCE

Drug Enforcement Administration	tion		FUR A BA	SIC CLASS	OF CONTE	(ULL	ED SUBSTANCE
SEE INSTRUCTIONS ON REVERSE SIDE	No in	ndividual manufacturing qu application form has	ota may be issued unless been received, 21 CFR 1		OM No		OMB APPROVAL No. 1117 - 0006
NAME OF BASIC CLASS DESIRED (Only one basic class per D				2. SCHEDULE NUMBER 3. DEA DE		CODE	
4. NAME AND ADDRESS O	F REGISTRANT (Includ	de No., Street, City, State	; State and ZIP Code)		YEAR FOR WHICH QUOTA IS REQUESTED  DEA REGISTRATION NUMBER		
				6	6. DEA REGIS	TRAIIC	ON NUMBER
7. NAME OF CONTACT PER	RSON	3	8. TELEPHONE NO. (Include ext., if applicable)			9. FA)	( NUMBER
NOTE: A	All quantities are to b	e expressed in grams	of anhydrous acid, b	ase or alkaloid	(not as salts	i).	
10.		QUOTAS PREVIOUSLY ISSUED BY DEA			EA		QUOTA REQUESTED
QUOTA HISTORY		2nd PRECEDING YEA			CURRENT YEAR		
		Gran	ns	Grams	G	rams	Grams
11. PRODUCTION DATA		2nd PRECEDING YEAR	1st PRECEDI YEAR	NG F	ESTIMATE FOR CURRENT YEAR		ESTIMATE FOR YEAR FOR WHICH QUOTA IS REQUESTED
I. INVENTORY AS OF DEC							
<ul><li>a. Bulk controlled substance</li><li>b. In-process material</li></ul>	ce						
c. Contained in FINISHED							
TOTAL $(a+b+c)$							
II. DISPOSITION / UTILIZAT	ΓΙΟΝ						
b. Exports							
III. ACQUISITION / PRODU	CTION						
a. Domestic Sources							
b. Importation							
12. IF THE PURPOSE IS TO		THER SUBSTANCE (S), I	 FURNISH THE FOLLOV	 Ving informat	ΓΙΟΝ:		
NAME OF DEA		AMOUNT USED FOR THIS PURPOSE					% YIELD
NEW SUBSTANCE	DRUG CODE NUMBER	2nd PRECEDING YEA	AR 1st PRECEDING	G YEAR C	CURRENT YEA	ιR	(Historical)
13. REMARKS							
TO. HEIMAHING							
SIGNATURE OF APPLICAN	Т	PI	RINT or TYPE NAME a	nd TITLE of SIG	NER		DATE

## INSTRUCTIONS FOR COMPLETING THE DEA FORM 189: Application For Individual Manufacturing Quota

The DEA-189 must be filed on or before May 1 of the year preceding the calendar year for which the manufacturing quota is being applied. Regulations governing quotas are included in Title 21, Code of Federal Regulations, Part 1300 to end. Copies of these regulations may be ordered from: The Government Printing Office, Superintendent of Documents, Attn: New Orders, P.O. Box 371954, Pittsburgh, PA 15250-7954. Submit the completed form to:

Drug Enforcement Administration Office of Diversion Control Drug & Chemical Evaluation Section (ODE) Washington, D.C. 20537

The following instructions are for those items which are not completely self-explanatory.

- **Item 11(I).** This is to include all factory and branch stocks which have reached that point in manufacturing as to be identifiable, whether in bulk form, in the process of manufacture, in finished form, or otherwise (e.g., damaged, defective, or impure substances awaiting disposal, substances held in quarantine, or substances maintained for extemporaneous compounding), as a basic class of controlled substance manufactured or otherwise acquired by a registrant, whether in bulk, commercial containers, or contained in pharmaceutical preparations in the possession of the registrant (including stocks held by the registrant under separate registration as a manufacturer, importer, exporter or distributor).
- Item 13. Enter any additional factors which may be relevant to the establishment of a manufacturing quota including the trend of and recent changes in the individual companies and the national rates of net disposal, production cycles and current inventory position, the economic and physical availability of raw materials for use in manufacturing and for inventory purposes, yield and stability problems, potential disruptions to production (including possible labor strikes) and recent unforeseen emergencies such as floods or fires.

Under the Paperwork Reduction Act, a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Public reporting burden for this collection of information is estimated to average 30 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to the FOI and Records Management Section, Drug Enforcement Administration, Washington, D.C. 20537; and to the Office of Management and Budget, Paperwork Reduction Project No. 1117-0006, Washington, D.C. 20503