

Basic class	Established 2023 quotas (g)	Basic class	Established 2023 quotas (g)
Bezitramide .....	25	Phenmetrazine .....	25
Carfentanil .....	20	Phenylacetone .....	100
Cocaine .....	60,492	Piminodine .....	25
Codeine (for conversion) .....	1,085,024	Racemethorphan .....	5
Codeine (for sale) .....	21,003,397	Racemorphan .....	5
D-amphetamine (for sale) .....	21,200,000	Remifentanil .....	3,000
D,l-amphetamine .....	21,200,000	Secobarbital .....	172,100
d-amphetamine (for conversion) .....	20,000,000	Sufentanil .....	4,000
Dexmethylphenidate (for sale) .....	6,200,000	Tapentadol .....	11,941,416
Dexmethylphenidate (for conversion) .....	4,200,000	Thebaine .....	57,137,944
Dextropropoxyphene .....	35	<b>List I Chemicals</b>	
Dihydrocodeine .....	132,658	Ephedrine (for conversion) ...	41,100
Dihydroetorphine .....	25	Ephedrine (for sale) .....	4,136,000
Diphenoxylate (for conversion) .....	14,100	Phenylpropanolamine (for conversion) .....	14,878,320
Diphenoxylate (for sale) .....	770,800	Phenylpropanolamine (for sale) .....	7,990,000
Ecgonine .....	60,492	Pseudoephedrine (for conversion) .....	1,000
Ethylmorphine .....	30	Pseudoephedrine (for sale) ..	174,246,000
Etorphine hydrochloride .....	32		
Fentanyl .....	731,452		
Glutethimide .....	25		
Hydrocodone (for conversion) .....	1,250		
Hydrocodone (for sale) .....	27,239,822		
Hydromorphone .....	1,994,117		
Isomethadone .....	30		
L-amphetamine .....	30		
Levo-alphaacetylmethadol (LAAM) .....	25		
Levomethorphan .....	30		
Levorphanol .....	23,010		
Lisdexamfetamine .....	26,500,000		
Meperidine .....	681,289		
Meperidine Intermediate-A ...	30		
Meperidine Intermediate-B ...	30		
Meperidine Intermediate-C ...	30		
Metazocine .....	15		
Methadone (for sale) .....	25,619,700		
Methadone Intermediate .....	27,673,600		
Methamphetamine .....	150		
d-methamphetamine (for conversion) .....	485,020		
d-methamphetamine (for sale) .....	47,000		
l-methamphetamine .....	587,229		
Methylphenidate (for sale) ...	41,800,000		
Methylphenidate (for conversion) .....	15,300,000		
Metopon .....	25		
Moramide-intermediate .....	25		
Morphine (for conversion) ...	2,458,460		
Morphine (for sale) .....	21,747,625		
Nabilone .....	62,000		
Norfentanyl .....	25		
Noroxymorphone (for conversion) .....	22,044,741		
Noroxymorphone (for sale) ...	1,000		
Oliceridine .....	25,100		
Opium (powder) .....	250,000		
Opium (tincture) .....	530,837		
Oripavine .....	33,010,750		
Oxycodone (for conversion) .....	437,827		
Oxycodone (for sale) .....	53,840,608		
Oxymorphone (for conversion) .....	28,204,371		
Oxymorphone (for sale) .....	516,351		
Pentobarbital .....	33,843,337		
Phenazocine .....	25		
Phencyclidine .....	35		

**DEPARTMENT OF JUSTICE**

**Drug Enforcement Administration**

[Docket No. 1121]

**Bulk Manufacturer of Controlled Substances Application: Bulk Manufacturer of Marihuana: Alm Management**

**AGENCY:** Drug Enforcement Administration, Justice.

**ACTION:** Notice of application.

**SUMMARY:** The Drug Enforcement Administration (DEA) is providing notice of an application it has received from an entity applying to be registered to manufacture in bulk basic class(es) of controlled substances listed in schedule I. DEA intends to evaluate this and other pending applications according to its regulations governing the program of growing marihuana for scientific and medical research under DEA registration.

**DATES:** Registered bulk manufacturers of the affected basic class(es), and applicants therefore, may submit electronic comments on or objections to the issuance of the proposed registration on or before January 31, 2023.

**ADDRESSES:** DEA requires that all comments be submitted electronically through the Federal eRulemaking Portal, which provides the ability to type short comments directly into the comment field on the web page or attach a file for lengthier comments. Please go to <https://www.regulations.gov> and follow the online instructions at that site for submitting comments. Upon submission of your comment, you will receive a Comment Tracking Number. Please be aware that submitted comments are not instantaneously available for public view on <https://www.regulations.gov>. If you have received a Comment Tracking Number, your comment has been successfully submitted and there is no need to resubmit the same comment.

**SUPPLEMENTARY INFORMATION:** The Controlled Substances Act (CSA) prohibits the cultivation and distribution of marihuana except by persons who are registered under the CSA to do so for lawful purposes. In accordance with the purposes specified in 21 CFR 1301.33(a), DEA is providing notice that the entity identified below has applied for registration as a bulk manufacturer of schedule I controlled substances. In response, registered bulk manufacturers of the affected basic class(es), and applicants therefore, may submit electronic comments on or objections of the requested registration, as provided in this notice. This notice

The Administrator also establishes APQ for all other schedule I and II controlled substances included in 21 CFR 1308.11 and 1308.12 at zero. In accordance with 21 CFR 1303.13 and 21 CFR 1315.13, upon consideration of the relevant factors, the Administrator may adjust the 2023 APQ and AAN as needed.

**Signing Authority**

This document of the Drug Enforcement Administration was signed on November 29, 2022, by Administrator Anne Milgram. That document with the original signature and date is maintained by DEA. For administrative purposes only, and in compliance with requirements of the Office of the Federal Register, the undersigned DEA Federal Register Liaison Officer has been authorized to sign and submit the document in electronic format for publication, as an official document of DEA. This administrative process in no way alters the legal effect of this document upon publication in the **Federal Register**.

**Scott Brinks,**

*Federal Register Liaison Officer, Drug Enforcement Administration.*

[FR Doc. 2022-26351 Filed 11-30-22; 11:15 am]

**BILLING CODE P**

does not constitute any evaluation or determination of the merits of the application submitted.

The applicant plans to manufacture bulk active pharmaceutical ingredients (APIs) for product development and distribution to DEA registered researchers. If the application for registration is granted, the registrant would not be authorized to conduct other activity under this registration aside from those coincident activities specifically authorized by DEA regulations. DEA will evaluate the application for registration as a bulk manufacturer for compliance with all applicable laws, treaties, and regulations and to ensure adequate safeguards against diversion are in place.

As this applicant has applied to become registered as a bulk manufacturer of marihuana, the application will be evaluated under the criteria of 21 U.S.C. 823(a). DEA will conduct this evaluation in the manner described in the rule published at 85 FR 82333 on December 18, 2020, and reflected in DEA regulations at 21 CFR part 1318.

In accordance with 21 CFR 1301.33(a), DEA is providing notice that on October 25, 2022, Alm Management, 7460 Varna Avenue, North Hollywood, California 91605, applied to be registered as a bulk manufacturer of the following basic class(es) of controlled substances:

Controlled substance	Drug code	Schedule
Marihuana .....	7360	I

**Matthew Strait,**

*Deputy Assistant Administrator.*

[FR Doc. 2022-26208 Filed 12-1-22; 8:45 am]

**BILLING CODE 4410-09-P**

**DEPARTMENT OF JUSTICE**

**Drug Enforcement Administration**

[Docket No. 1122]

**Bulk Manufacturer of Controlled Substances Application: Bulk Manufacturer of Marihuana: Attitude Wellness**

**AGENCY:** Drug Enforcement Administration, Justice.

**ACTION:** Notice of application.

**SUMMARY:** The Drug Enforcement Administration (DEA) is providing notice of an application it has received from an entity applying to be registered to manufacture in bulk basic class(es) of controlled substances listed in schedule I. DEA intends to evaluate this and other pending applications according to its regulations governing the program of growing marihuana for scientific and medical research under DEA registration.

**DATES:** Registered bulk manufacturers of the affected basic class(es), and applicants therefore, may submit electronic comments on or objections to the issuance of the proposed registration on or before January 31, 2023.

**ADDRESSES:** DEA requires that all comments be submitted electronically through the Federal eRulemaking Portal, which provides the ability to type short comments directly into the comment field on the web page or attach a file for lengthier comments. Please go to <https://www.regulations.gov> and follow the online instructions at that site for submitting comments. Upon submission of your comment, you will receive a Comment Tracking Number. Please be aware that submitted comments are not instantaneously available for public view on <https://www.regulations.gov>. If you have received a Comment Tracking Number, your comment has been successfully submitted and there is no need to resubmit the same comment.

**SUPPLEMENTARY INFORMATION:** The Controlled Substances Act (CSA) prohibits the cultivation and distribution of marihuana except by persons who are registered under the CSA to do so for lawful purposes. In

accordance with the purposes specified in 21 CFR 1301.33(a), DEA is providing notice that the entity identified below has applied for registration as a bulk manufacturer of schedule I controlled substances. In response, registered bulk manufacturers of the affected basic class(es), and applicants therefor, may submit electronic comments on or objections of the requested registration, as provided in this notice. This notice does not constitute any evaluation or determination of the merits of the application submitted.

The applicant plans to manufacture bulk active pharmaceutical ingredients (APIs) for product development and distribution to DEA registered researchers. If the application for registration is granted, the registrant would not be authorized to conduct other activity under this registration aside from those coincident activities specifically authorized by DEA regulations. DEA will evaluate the application for registration as a bulk manufacturer for compliance with all applicable laws, treaties, and regulations and to ensure adequate safeguards against diversion are in place.

As this applicant has applied to become registered as a bulk manufacturer of marihuana, the application will be evaluated under the criteria of 21 U.S.C. 823(a). DEA will conduct this evaluation in the manner described in the rule published at 85 FR 82333 on December 18, 2020, and reflected in DEA regulations at 21 CFR part 1318.

In accordance with 21 CFR 1301.33(a), DEA is providing notice that on October 3, 2022, Attitude Wellness, 9741 South Industrial Drive, Ewart, Michigan 49631, applied to be registered as a bulk manufacturer of the following basic class(es) of controlled substances:

Controlled substance	Drug code	Schedule
Marihuana Extract .....	7350	I
Marihuana .....	7360	I
Tetrahydrocannabinols .....	7370	I