Administration (DEA) as importer of schedule I controlled substances.

SUPPLEMENTARY INFORMATION: The company listed below applied to be registered as an importer of basic class of controlled substances. Information on previously published notice is listed in the table below. No comments or objections were submitted and no requests for a hearing were submitted for this notice.

Company	FR Docket	Published
Sharp (Bethlehem),	84 FR	March 18,
LLC.	9837.	2019

The DEA has considered the factors in 21 U.S.C. 823, 952(a) and 958(a) and determined that the registration of the listed registrant to import the applicable basic class of schedule I controlled substances is consistent with the public interest and with United States obligations under international treaties, conventions, or protocols in effect on May 1, 1971. The DEA investigated the company's maintenance of effective controls against diversion by inspecting and testing the company's physical security systems, verifying the company's compliance with state and local laws, and reviewing the company's background and history.

Therefore, pursuant to 21 U.S.C. 952(a) and 958(a), and in accordance with 21 CFR 1301.34, the DEA has granted a registration as an importer for schedule I controlled substances to the above listed company.

Dated: June 3, 2019.

John J. Martin,

Assistant Administrator.

[FR Doc. 2019–14023 Filed 7–1–19; 8:45 am]

BILLING CODE 4410-09-P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration [Docket No. DEA-392]

Importer of Controlled Substances

ACTION: Notice of application.

DATES: Registered bulk manufacturers of the affected basic classes, and applicants therefore, may file written comments on or objections to the issuance of the proposed registration on or before August 1, 2019. Such persons may also file a written request for a hearing on the application on or before August 1, 2019.

Application: Bellwyck Clinical Services

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: Administrator, 8701 Morrissette Drive, Springfield, Virginia 22152. All requests for a hearing should also 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 April 17, 2019, Bellwyck Clinical Services, 8946 Global Way, West Chester, Ohio 45069 applied to be registered as an importer of the following basic class of controlled substances:

Controlled substance	Drug code	Schedule
Amphetamine	1100	
Methylphenidate	1724	
Oxycodone	9143	

The company plans to import the listed controlled substances in dosage form to conduct clinical trials. Approval of permit applications will occur only when the registrant's activity is consistent with what is authorized under 21 U.S.C. 952(a) (2). Authorization will not extend to the import of FDA approved or non-approved finished dosage forms for commercial sale.

Dated: June 18, 2019.

John J. Martin,

Assistant Administrator.

[FR Doc. 2019–14027 Filed 7–1–19; 8:45 am]

BILLING CODE 4410-09-P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

[Docket No. DEA-392]

Bulk Manufacturer of Controlled Substances Application: Pisgah Laboratories, Inc.

ACTION: Notice of application.

DATES: Registered bulk manufacturers of the affected basic classes, and applicants therefore, may file written comments on or objections to the issuance of the proposed registration on or before September 3, 2019.

ADDRESSES: Written comments should be sent to: Drug Enforcement

Administration, Attention: DEA Federal Register Representative/DPW, 8701 Morrissette Drive, Springfield, Virginia 22152.

SUPPLEMENTARY INFORMATION: In

accordance with 21 CFR 1301.33(a), this is notice that on March 5, 2019, Pisgah Laboratories, Inc., 3222 Old Hendersonville Highway, Pisgah Forest, North Carolina 28768 applied to be registered as a bulk manufacturer of the following basic classes of controlled substances:

Controlled substance	Drug code	Schedule
Difenoxin Diphenoxylate Levorphanol Meperidine inter- mediate-B.	9168 9170 9220 9233	

The company plans to manufacture the listed controlled substances in bulk for distribution to its customers.

Dated: June 19, 2019.

John J. Martin,

Assistant Administrator.

[FR Doc. 2019-14028 Filed 7-1-19; 8:45 am]

BILLING CODE 4410-09-P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

[Docket No. DEA-392]

Importer of Controlled Substances Application: Lipomed

ACTION: Notice of application.

DATES: Registered bulk manufacturers of the affected basic classes, and applicants therefore, may file written comments on or objections to the issuance of the proposed registration on or before August 1, 2019. Such persons may also file a written request for a hearing on the application on or before August 1, 2019.

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 hearing must be sent to: Drug Enforcement Administration, Attn: Administrator, 8701 Morrissette Drive, Springfield, Virginia 22152. All requests for hearing should also 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 March 28, 2019, Lipomed, 150 Cambridge Park Drive,

Suite 705, Cambridge, Massachusetts 02140 applied to be registered as an importer of the following basic classes of controlled substances:

Controlled substance	Drug code	Schedule
3-Fluoro-N-methylcathinone (3–FMC)	1233	1
Cathinone	1235	1
Methcathinone	1237	1
4-Fluoro-N-methylcathinone (4–FMC)	1238	!
Pentedrone (α-methylaminovalerophenone)	1246	!
Mephedrone (4-Methyl-N-methylcathinone)	1248	!
4-Methyl-N-ethylcathinone (4–MEC)	1249	1
Naphyrone	1258	1
N,N-Dimethylamphetamine	1475 1480	ł
Fenethylline	1503	i
Aminorex	1585	i
4-Methylaminorex (cis isomer)	1590	i
Gamma Hydroxybutyric Acid	2010	i
Methaqualone	2565	i
Mecloqualone	2572	i
JWH-250 (1-Pentyl-3-(2-methoxyphenylacetyl) indole)	6250	1
SR-18 (Also known as RCS-8) (1-Cyclohexylethyl-3-(2-methoxyphenylacetyl) indole)	7008	I
ADB-FÙBINACA (N-(1-amino-3,3-dimethyl-1-oxobutan-2-yl)-1-(4-fluorobenzyl)-1H-indazole-3-carboxamide)	7010	1
5-Fluoro-UR-144 and XLR11 ([1-(5-Fluoro-pentyl)1H-indol-3-yl](2,2,3,3-tetramethylcyclopropyl)methanone)	7011	1
AB-FUBINACA (N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-(4-fluorobenzyl)-1H-indazole-3-carboxamide)	7012	1
JWH-019 (1-Hexyl-3-(1-naphthoyl)indole)	7019	1
MDMB-FUBINACA (Methyl 2-(1-(4-fluorobenzyl)-1H-indazole-3-carboxamido)-3,3-dimethylbutanoate)	7020	I
FUB-AMB, MMB-FUBINACA, AMB-FUBINACA (2-(1-(4-fluorobenzyl)-1Hindazole-3-carboxamido)-3-	7021	1
methylbutanoate).		
AB-PINACA (N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-pentyl-1H-indazole-3-carboxamide)	7023	I
THJ-2201 ([1-(5-fluoropentyl)-1H-indazol-3-yl](naphthalen-1-yl)methanone)	7024	I
5F-AB-PINACA (N-(1-amino-3methyl-1-oxobutan-2-yl)-1-(5-fluoropentyl)-1H-indazole-3-carboxamide)	7025	I
AB_CHMINACA (N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-(cyclohexylmethyl)-1H-indazole-3-carboxamide)	7031	Į.
MAB_CHMINACA (N-(1-amino-3,3dimethyl-1-oxobutan-2-yl)-1-(cyclohexylmethyl)-1H-indazole-3-carboxamide)	7032	!
5F-AMB (Methyl 2-(1-(5-fluoropentyl)-1H-indazole-3-carboxamido)-3-methylbutanoate)	7033	l
5F-ADB; 5F-MDMB-PINACA (Methyl 2-(1-(5-fluoropentyl)-1H-indazole-3-carboxamido)-3,3-dimethylbutanoate)	7034	!
ADB-PINACA (N-(1-amino-3,3-dimethyl-1-oxobutan-2-yl)-1-pentyl-1H-indazole-3-carboxamide)	7035 7042	1
dimethylbutanoate).		
MMB-CHMICA , AMB-CHMICA (methyl 2-(1-(cyclohexylmethyl)-1H-indole-3-carboxamido)-3-methylbutanoate)	7044	1
APINACA and AKB48 (N-(1-Adamantyl)-1-pentyl-1H-indazole-3-carboxamide)	7048	l
5F-APINACA, 5F-AKB48 (N-(adamantan-1-yl)-1-(5-fluoropentyl)-1H-indazole-3-carboxamide)	7049	ļ.
JWH–081 (1-Pentyl-3-(1-(4-methoxynaphthoyl) indole)	7081	!
5F-CUMYL-P7AICA (1-(5-fluoropentyl)-N-(2-phenylpropan-2-yl)-1H-pyrrolo[2,3-b]pyridine-3-carboxamide)	7085	!
4-CN-CUMYL-BUTINACA, 4-cyano-CUMYL-BUTINACA, 4-CN-CUMYL BINACA, CUMYL-4CN-BINACA, SGT-78 (1-(4-cyanobutyl)-N-(2-phenylpropan-2-yl)-1H-indazole-3-carboximide).	7089	
SR-19 (Also known as RCS-4) (1-Pentyl-3-[(4-methoxy)-benzoyl] indole)	7104	!
JWH-018 (also known as AM678) (1-Pentyl-3-(1-naphthoyl)indole)	7118	!
JWH–122 (1-Pentyl-3-(4-methyl-1-naphthoyl) indole)	7122	!
UR-144 (1-Pentyl-1H-indol-3-yl)(2,2,3,3-tetramethylcyclopropyl)methanone)	7144 7173	1
JWH–200 (1-[2-(4-Morpholinyl)ethyl]-3-(1-naphthoyl)indole)	7173	i
AM2201 (1-(5-Fluoropentyl)-3-(1-naphthoyl) indole)	7201	i
JWH–203 (1-Pentyl-3-(2-chlorophenylacetyl) indole)	7203	i
NM2201, CBL2201 (Naphthalen-1-yl 1-(5-fluoropentyl)-1H-indole-3-carboxylate)	7221	i
PB–22 (Quinolin-8-yl 1-pentyl-1H-indole-3-carboxylate)	7222	i
5F–PB–22 (Quinolin-8-yl 1-(5-fluoropentyl)-1H-indole-3-carboxylate)	7225	i
Alpha-ethyltryptamine	7249	i
Ibogaine	7260	i
CP–47,497 (5-(1,1-Dimethylheptyl)-2-[(1R,3S)-3-hydroxycyclohexyl-phenol)	7297	1
CP-47,497 C8 Homologue (5-(1,1-Dimethyloctyl)-2-[(1Ř,3S)3-hydroxycyclohexyl-phenol)	7298	1
Lysergic acid diethylamide	7315	1
2,5-Dimethoxy-4-(n)-propylthiophenethylamine (2C-T-7)	7348	I
Marihuana extract	7350	1
Marihuana	7360	1
Tetrahydrocannabinols	7370	1
Parahexyl	7374	1
Mescaline	7381	1
2C-T-2, (2-(4-Ethylthio-2,5-dimethoxyphenyl) ethanamine)	7385	1
3,4,5-Trimethoxyamphetamine	7390	1
4-Bromo-2,5-dimethoxyamphetamine	7391	1
4-Bromo-2,5-dimethoxyphenethylamine	7392	1
4-Methyl-2,5-dimethoxyamphetamine	7395	1

Controlled substance	Drug code	Schedule
2,5-Dimethoxyamphetamine	7396	1
JWH-398 (1-Pentyl-3-(4-chloro-1-naphthoyl) indole)	7398	!
2,5-Dimethoxy-4-ethylamphetamine 3,4-Methylenedioxyamphetamine	7399 7400	
5-Methoxy-3,4-methylenedioxyamphetamine	7400	i
N-Hydroxy-3,4-methylenedioxyamphetamine		i
3,4-Methylenedioxy-N-ethylamphetamine	7404	I
3,4-Methylenedioxymethamphetamine		I
4-Methoxyamphetamine	7411	
5-Methoxy-N-N-dimethyltryptamine	7431 7432	
Alpha-methyltryptamine		1
Diethyltryptamine	7434	i
Dimethyltryptamine	7435	i
Psilocybin	7437	1
Psilocyn	7438	1
5-Methoxy-N,N-diisopropyltryptamine	7439	
N-Ethyl-1-phenylcyclohexylamine	7455 7458	
1-[1-(2-Thienyl)cyclohexyl]piperidine	7470	i
1-[1-(2-Thienyl)cyclohexyl]pyrrolidine	7473	li
N-Ethyl-3-piperidyl benzilate	7482	1
N-Methyl-3-piperidyl benzilate	7484	1
N-Benzylpiperazine	7493	
4-MePPP (4-Methyl-alphapyrrolidinopropiophenone)	7498	
2C-D (2-(2,5-Dimethoxy-4-methylphenyl) ethanamine)	7508 7509	l I
2C-H (2-(2,5-Dimethoxyphenyl) ethanamine)	7517	i
2C-I (2-(4-iodo-2,5-dimethoxyphenyl) ethanamine)	7518	i
2C-C (2-(4-Chloro-2,5-dimethoxyphenyl) ethanamine)	7519	1
2C-N (2-(2,5-Dimethoxy-4-nitro-phenyl) ethanamine)		
2C-P (2-(2,5-Dimethoxy-4-(n)-propylphenyl) ethanamine)	7524	
2C-T-4 (2-(4-Isopropylthio)-2,5-dimethoxyphenyl) ethanamine)		1
25B–NBOMe (2-(4-bromo-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl) ethanamine)	7536	i
25C–NBOMe (2-(4-chloro-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl) ethanamine)	7537	i
25I-NBOMe (2-(4-iodo-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl) ethanamine)	7538	I
Methylone (3,4-Methylenedioxy-N-methylcathinone)	7540	1
Butylone		!
Pentylone	7542	
N-Ethylpentylone, ephylone (1-(1,3-benzodioxol-5-yl)-2-(ethylamino)-pentan-1-one)	7543 7545	l I
α -PBP (alpha-pyrrolidinobutiophenone)	7546	i
AM-694 (1-(5-Fluoropentyl)-3-(2-iodobenzoyl) indole)	7694	i
Acetyldihydrocodeine	9051	1
Benzylmorphine	9052	1
Codeine-N-oxide	9053	
Cyprenorphine	9054	1
Desomorphine	9055 9056	
Codeine methylbromide	9070	i
Dihydromorphine	9145	1
Difenoxin	9168	1
Heroin	9200	!
Hydromorphinol	9301	
Methyldesorphine	9302	
Methyldihydromorphine	9304 9305	i
Morphine methylsulfonate	9306	li
Morphine-N-oxide	9307	i
Myrophine	9308	1
Nicocodeine	9309	l i
Nicomorphine	9312	!
Normorphine	9313	
Pholcodine	9314 9315	
Acetorphine	9319	li
Drotebanol	9335	i
U-47700 (3,4-dichloro-N-[2-(dimethylamino)cyclohexyl]-N-methylbenzamide)	9547	1
AH-7921 (3,4-dichloro-N-[(1-dimethylamino)cyclohexylmethyl]benzamide))	9551	1
MT-45 (1-cyclohexyl-4-(1,2-diphenylethyl)piperazine))	9560	1
Acetylmethadol	9601	I
Allylprodine	9602	

Controlled substance	Drug code	Schedule
Alphameprodine	9604	I
Alphamethadol	9605	!
Benzethidine	9606	1
Betacetylmethadol	9607 9608	i
Betamethadol	9609	i
Betaprodine	9611	i
Clonitazene	9612	1
Dextromoramide	9613	!
Diampromide	9615	1
Diethylthiambutene	9616 9617	1
Dimenovador	9618	i
Dimethylthiambutene	9619	i
Dioxaphetyl butyrate	9621	1
Dipipanone	9622	1
Ethylmethylthiambutene	9623	!
Etonitazene	9624	!
Etoxeridine	9625	!
Furethidine	9626 9627	i
Ketobemidone	9628	i
Levomoramide	9629	1
Levophenacylmorphan	9631	1
Morpheridiné	9632	1
Noracymethadol	9633	!
Norlevorphanol	9634	!
Normethadone	9635 9636	1
NorpipanonePhenadoxone	9637	i
Phenampromide	9638	i
Phenoperidine	9641	i
Piritramide	9642	1
Proheptazine	9643	1
Properidine	9644	1
Racemoramide	9645	!
Trimeperidine	9646	1
PhenomorphanPropiram	9647 9649	i
1-Methyl-4-phenyl-4-propionoxypiperidine	9661	i
1-(2-Phenylethyl)-4-phenyl-4-acetoxypiperidine	9663	I
Tilidine	9750	1
Acryl fentanyl (N-(1-phenethylpiperidin-4-yl)-N-phenylacrylamide)	9811	!
Para-Fluorofentanyl	9812	!
3-Methylfentanyl	9813 9814	1
Acetyl-alpha-methylfentanyl	9815	i
N-(2-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)propionamide	9816	i
Acetyl Fentanyl (N-(1-phenethylpiperidin-4-yl)-N-phenylacetamide)	9821	i
Butyryl Fentanyl	9822	1
Para-fluorobutyryl fentanyl	9823	1
4-Fluoroisobutyryl fentanyl (N-(4-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)isobutyramide)	9824	1
2-methoxy-N-(1-phenethylpiperidin-4-yl)-N-phenylacetamide	9825	1
Para-chloroisobutyryl fentanyl	9826 9827	1
Beta-hydroxyfentanyl	9830	i
Beta-hydroxy-3-methylfentanyl	9831	i
Alpha-methylthiofentanyl	9832	1
3-Methylthiofentanyl	9833	1
Furanyl fentanyl (N-(1-phenethylpiperidin-4-yl)-N-phenylfuran-2-carboxamide)	9834	!
Thiofentanyl	9835	1
Beta-hydroxythiofentanyl	9836	I
Para-methoxybutyryl fentanyl	9837 9838	i
Valeryl fentanyl	9840	i
N-(1-phenethylpiperidin-4-yl)-N-phenyltetrahydrofuran-2-carboxamide)	9843	i
Cyclopropyl Fentanyl	9845	1
Cyclopentyl fentanyl	9847	1
Fentanyl related-compounds as defined in 21 CFR 1308.11(h)	9850	1
Amphetamine	1100	II
Methamphetamine	1105	
Lisdexamfetamine	1205	II II
Phenmetrazine	1631	

Controlled substance	Drug code	Schedu
mobarbital	2125	II
entobarbital	2270	II
ecobarbital	2315	II
lutethimide	2550	II
ronabinol in an oral solution in a drug product approved for marketing by the U.S. Food and Drug Administration	7365	II
abilone	7379	II
Phenylcyclohexylamine	7460	II
nencyclidine	7471	II
NPP (4-Anilino-N-phenethyl-4-piperidine)	8333	II
nenylàcetone	8501	II
Piperidinocyclohexanecarbonitrile	8603	ii
phaprodine	9010	ii
nileridine	9020	ii
ocaine	9041	ii
odeine	9050	ii
orphine HCI	9059	ii
hydrocodeine	9120	ii
xycodone	9143	ii
	9150	lii
ydromorphone		
phenoxylate	9170	II
gonine	9180	II
hylmorphine	9190	II
/drocodone	9193	III
vomethorphan	9210	II
evorphinol	9220	II
omethadone	9226	II
eperidine	9230	II
eperidine-intermediate-A	9232	II
eperidine intermediate-B	9233	II
eperidine intermediate-C	9234	II
etazocineetazocine	9240	II
ethadone	9250	II
ethadone intermediate	9254	II
etopon	9260	II
extropropoxyphene, bulk (non-dosage forms)	9273	II
orphine	9300	II
ripavine	9330	II
nebaine	9333	II
hydroetorphine	9334	II
evo-alphacetylmethadol	9648	II
xymorphone	9652	II
proxymorphone	9668	II
nenazocine	9715	ii
niafentanil	9729	ii
minodine	9730	ii
acemethorphan	9732	lii
acemorphan	9733	ii
fentanil	9737	ii
emifentanil	9739	II II
ufentanil	9740	III
arfentanil	9743	II
apentadol	9780	II
ezitramide	9800	II
entanyi	9801	II
oramide-intermediate	9802	H

The company plans to import analytical reference standards for distribution to its customers for research and analytical purposes. Placement of these drug codes onto the company's registration does not translate into automatic approval of subsequent permit applications to import controlled substances. Approval of permit applications will occur only when the registrant's business activity is consistent with what is authorized in 21 U.S.C. 952(a)(2). Authorization will not extend to the import of FDA approved or non-approved finished dosage forms for commercial sale.

Dated: June 18, 2019.

John J. Martin,

Assistant Administrator.

[FR Doc. 2019-14026 Filed 7-1-19: 8:45 am]

BILLING CODE 4410-09-P

DEPARTMENT OF JUSTICE

[OMB Number 1125-0001]

Agency Information Collection Activities: Proposed Collection: Comments Requested; Application for Cancellation of Removal (42A) for Certain Permanent Residents; and **Application for Cancellation of** Removal and Adjustment of Status (42B) for Certain Nonpermanent Residents

AGENCY: Executive Office for Immigration Review, Department of Justice.

ACTION: 30-Day notice.

SUMMARY: The Department of Justice (DOJ), Executive Office for Immigration Review, will be submitting the following information collection request to the Office of Management and Budget (OMB) for review and approval in accordance with the Paperwork Reduction Act of 1995.

DATES: Comments are encouraged and will be accepted for an additional 30 days until August 1, 2019.

FOR FURTHER INFORMATION CONTACT: If

you have additional comments especially on the estimated public burden or associated response time, suggestions, or need a copy of the proposed information collection instrument with instructions or additional information, please contact Lauren Alder Reid, Assistant Director, Office of Policy, Executive Office for Immigration Review, 5107 Leesburg Pike, Suite 2500, Falls Church, VA 22041, telephone: (703) 305-0289. Written comments and/or suggestions can also be directed to the Office of Management and Budget, Office of

Information and Regulatory Affairs, Attention Department of Justice Desk Officer, Washington, DC 20530 or sent to OIRA submissions@omb.eop.gov.

SUPPLEMENTARY INFORMATION: Written comments and suggestions from the public and affected agencies concerning the proposed collection of information are encouraged. Your comments should address one or more of the following four points:

- —Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility;
- -Evaluate the accuracy of the agency's estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used;
- Enhance the quality, utility, and clarity of the information to be collected; and/or
- -Minimize the burden of the collection of information on those who are to respond, including through the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology, e.g., permitting electronic submission of responses.

Overview of This Information Collection

- 1. Type of Information Collection: Extension with changes to a currently approved collection.
- 2. The Title of the Form/Collection: Application for Cancellation of Removal for Certain Permanent Residents; and Application for Cancellation of Removal and Adjustment of Status for Certain Nonpermanent Residents.
- 3. The agency form number, if any, and the applicable component of the Department sponsoring the collection: The form numbers are EOIR-42A and EOIR-42B, Executive Office for Immigration Review, United States Department of Justice.
- 4. Affected public who will be asked or required to respond, as well as a brief abstract: Primary: Individual aliens determined to be removable from the United States, Other: None, Abstract: This information collection is necessary to determine the statutory eligibility of individual aliens who have been determined to be removable from the United States for cancellation of their removal, as well as to provide information relevant to a favorable exercise of discretion.
- 5. An estimate of the total number of respondents and the amount of time

estimated for an average respondent to respond: It is estimated that 27,999 respondents will complete the form annually with an average of 5 hours and 50 minutes per response.

6. An estimate of the total public burden (in hours) associated with the collection: The estimated public burden associated with this collection is 162.394 hours.

If additional information is required contact: Melody Braswell, Department Clearance Officer, United States Department of Justice, Justice Management Division, Policy and Planning Staff, Two Constitution Square, 145 N Street NE, 3E.405B, Washington, DC 20530.

Dated: June 27, 2019.

Melody D. Braswell,

Department Clearance Officer for PRA, U.S. Department of Justice.

[FR Doc. 2019-14064 Filed 7-1-19; 8:45 am]

BILLING CODE 4410-30-P

NUCLEAR REGULATORY COMMISSION

[Docket Nos. 52-025 and 52-026; NRC-2008-02521

Southern Nuclear Operating Company, Inc.; Vogtle Electric Generating Plant, Units 3 and 4; Passive Residual Heat **Removal Instrumentation Minimum Inventory Displays**

AGENCY: Nuclear Regulatory Commission.

ACTION: Exemption and combined license amendment; issuance.

SUMMARY: The U.S. Nuclear Regulatory Commission (NRC) is granting an exemption to allow a departure from the certification information of Tier 1 of the generic design control document (DCD) and is issuing License Amendment Nos. 162 and 160 to Combined Licenses (COLs), NPF-91 and NPF-92. The COLs were issued to Southern Nuclear Operating Company, Inc., and Georgia Power Company, Oglethorpe Power Corporation, MEAG Power SPVM, LLC, MEAG Power SPVJ, LLC, MEAG Power SPVP, LLC, and the City of Dalton, Georgia (collectively SNC); for construction and operation of the Vogtle Electric Generating Plant (VEGP) Units 3 and 4, located in Burke County, Georgia.

The granting of the exemption allows the changes to Tier 1 information asked for in the amendment. Because the acceptability of the exemption was determined in part by the acceptability of the amendment, the exemption and