State of Florida Department of Business and Professional Regulation Division of Drugs, Devices, and Cosmetics

Application for Permit as a Nonresident Prescription Drug Manufacturer – Virtual Form No.: DBPR-DDC-236

APPLICATION CHECKLIST – IMPORTANT – Submit all items on the checklist below with your application to ensure faster processing.

APPLICATION	APPLICATION REQUIREMENTS
Application for Permit as a Nonresident Prescription Drug Manufacturer - Virtual	\$1,000 nonrefundable biennial application fee. If the applicant is applying for multiple manufacturing permits in the applicant's name and at applicant's address, you are only required to pay for the permit that has the highest fee. Make cashier's check, corporate check, or money order payable to the Florida Department of Business and Professional Regulation. If you answer "Yes" to any question in Section IV, be sure to provide a detailed explanation along with any relevant documentation. Submit photocopy of your license/permit(s) issued by your resident state that authorizes the distribution of prescription drugs from the applicant's establishment's address. Sign and date the Affidavit section of the application.
	Mail completed application to: Department of Business and Professional Regulation Division of Drugs, Devices and Cosmetics 2601 Blair Stone Road Tallahassee, FL 32399-1047

PLEASE NOTE:

- Telephone, email, and fax contact information is used to quickly resolve questions with applications. If such information is not provided, questions regarding applications will be mailed to the application contact's mailing address and may take longer to resolve.
- The disclosure of Social Security numbers is mandatory on all professional and occupational license applications, is solicited by the authority granted by 42 U.S.C. §§ 653 and 654, and will be used by the Department of Business and Professional Regulation pursuant to §§ 409.2577, 409.2598, 499.012(4)(a)f, 499.012(8)(o), 499.63(2), and 559.79(3), Florida Statutes, for the efficient screening of applicant and licensees by a Title IV-D child support agency to assure compliance with child support obligations. It is also required by § 559.79(1), Florida Statutes, for determining eligibility for licensure and mandated by the authority granted by 42 U.S.C. § 405(c)(2)(C)(i), to be used by the Department of Business and Professional Regulation to identify licensees for tax administration purposes.

State of Florida Department of Business and Professional Regulation Division of Drugs, Devices, and Cosmetics

Application for Permit as a Nonresident Prescription Drug Manufacturer - Virtual Form No.: DBPR-DDC-236

If you have any questions or need assistance in completing this application, please contact the Department of Business and Professional Regulation, Division of Drugs, Devices and Cosmetics, at **850.717.1800**. *For additional information* see the instructions at the beginning of this application.

CHECK ONE OF THE APPLICATION TYPES

Section I – Application Type

 New Application [3346/1020] New Application due to change in ownership. If checked, provide legal documentation for the change of ownership (i.e. Bill of Sale, stock transfer, merger). [3346/1020] Current Permit Number:
Section II – Applicant Information
APPLICANT INFORMATION
TAXPAYER IDENTIFICATION NUMBER OR FEDERAL EMPLOYER IDENTIFICATION NUMBER
This is a unique nine-digit number assigned by the Internal Revenue Service (IRS) to business entities operating in the United States for the purposes of identification. When the number is used for identification rather than employment tax reporting, it is usually referred to as a Taxpayer Identification Number (TIN), and when used for the purposes of reporting employment taxes, it is usually referred to as the Federal Employer Identification Number (FEIN).
Applicant's TIN/FEIN:
FULL LEGAL NAME
The "full legal name" is the complete name of the business entity that will be operating the establishment. This is generally the name that is on the documents that establish the existence or formation of the business entity. For example, a corporation's full legal name would normally be the name that is found in the corporation's articles of incorporation.
Applicant's Full Legal Name:
FICTITIOUS, TRADE, OR BUSINESS NAME
If the applicant intends to operate the permitted establishment under a name that is different from the Applicant's Full Legal Name listed above – e.g. fictitious, trade, or business name (also commonly referred to as a "dba", "D/B/A", or "doing business as" name – this name must be registered with the Florida Department of State, Division of Corporations). This is the name that will appear on the permit issued to the applicant by the department and must be the name that the applicant uses on operational documents for permitted activities.
☐ The applicant WILL NOT operate the permitted establishment under a name that is different from the Applicant's Full Legal Name listed above.
☐ The applicant WILL operate the permitted establishment under the following fictitious, trade, or business name:
The fictitious, trade, or business name listed directly above, is registered with the Florida Department of State, Division of Corporations and the applicant has been issued the following registration number:

·					
APPLICANT MAI	I ING A	DDRESS			
Street Address or P.O. Box:					
City:	State:		Zip Code (+4 optional):		
Country (if located outside the United States):	Telep	hone Number:	Fax Number:		
PHYSICAL ADDRESS OF ESTABLISHMENT TO BE PERMITTED (only if different from mailing address) Check if not applicable					
	aress) C	neck 🔝 if not ap	piicabie		
Street Address:					
City:		State:	Zip Code (+4 optional):		
Country (if located outside the United States):		Telephone Numb	er:		
Email Address:		Fax Number:			
APPLICATIO	N CON	ГАСТ			
The application contact is the person that the departr			e questions regarding the		
responses provided on, or the documentation submit					
also the person that will receive all official communica					
Last/Surname: First:		Middle:	Suffix:		
Address:					
City:		State:	Zip Code (+4 optional):		
Telephone Number:	Fax Nu	ımber:	1		
E-Mail Address:					
EMERGENCY CONT	ACT IN	FORMATION			
The emergency contact is the person that the dep During an emergency, the department will contact the hours listed below. The contact information provided reach and communicate with the person listed in the	his pers d should	on at times outsided be sufficient for t	e of the regular business		
Last/Surname: First:		Middle:	Suffix:		
			-		
Position/Title:					
Street Address:					
City:		State:	Zip Code (+4 optional):		
Telephone Number:	E-Mail	Address:	•		

	OPE	ERATING HO	URS		
List the establishment's daily	hours of operation			e. REMEMBER 1	to circle "a.m." or
"p.m." for each time indicated					
Mon:a.m./p.m. to _				n./p.m. to:_	
Tue:a.m./p.m. to _				m./p.m. to:_	
Wed:a.m./p.m. to _	:a.m./p.	.m. Sun _	: a.r	m./p.m. to:_	a.m./p.m.
Thu :a.m./p.m. to _	:a.m./p.	.m.			
Section III – Ownership Info	ormation				
		E OF OWNER	SHIP		
☐ Publicly Held Corporation	∐ C	Closely Held C	orporation	Limited Lia	bility Company
☐ Charitable Organization—	501(c)(3)	Sole Proprietor	ship	Governme	nt
☐ Partnership – General		Professional Cossociation	orporation	☐ Professional Pr	
Partnership – Other, Include Limited Liability Partnership a Limited Partnership		ər:			
List the state of incorpora Proprietorship). Business ent		nder non-U.S.	laws list the	country of orgai	
State:					
Proprietorship or Partnership Department of State, Division	List name and address of the applicant's registered agent for service of process in Florida (except Sole Proprietorship or Partnership – General) and provide documentation, such as a print out from the Florida Department of State, Division of Corporations' webpage, that the applicant's registered agent is registered with the Florida Department of State, Division of Corporations. N/A (Partnership – General or Sole Proprietorship) Name:				
Address:					
City:			State:		Code (+4
List the name, position/title, social security number, date of birth and address of each owner, partner, member, manager, officer, director, chief executive, or other person who directly or indirectly controls the operation of the business entity, as applicable. For example, corporations would list officers and directors, limited liability companies would list members and managers, etc.					
1. Name & Title:		Social Securi		Date of Birth:	% of Ownership:
Street Address:		City:		State:	Zip Code:
2. Name & Title:		Social Securi	ty #:	Date of Birth:	% of Ownership:

	Street Address:	City:	State:	Zip Code:
3.	Name & Title:	Social Security #:	Date of Birth:	% of Ownership:
	Street Address:	City:	State:	Zip Code:
4.	Name & Title:	Social Security #:	Date of Birth:	% of Ownership:
	Street Address:	City:	State:	Zip Code:
5.	Name & Title:	Social Security #:	Date of Birth:	% of Ownership:
	Street Address:	City:	State:	Zip Code:
6.	Name & Title:	Social Security #:	Date of Birth:	% of Ownership:
	Street Address:	City:	State:	Zip Code:
7.	Name & Title:	Social Security #:	Date of Birth:	% of Ownership:
	Street Address:	City:	State:	Zip Code:
8.	Name & Title:	Social Security #:	Date of Birth:	% of Ownership:
	Street Address:	City:	State:	Zip Code:
Lis	I t the name, social security number, date o	L birth and address of each	L ch person who c	owns 10 percent or
mo list	ore of the outstanding stock or equity intere the business entity name, FEID/FEIN and te of birth.	est in the business entity.	If such person is	s a business entity,
1.	Name:	SSN/FEID/FEIN#	Date of Birth: ☐ N/A	% of Ownership:
	Street Address:	City:	State:	Zip Code:
2.	Name:	SSN/FEID/FEIN#	Date of Birth:	% of Ownership:
	Street Address:	City:	State:	Zip Code:

3.	Name:	SSN/	FEID/FEIN#	Date of Birth:	% of Ownership:
	Street Address:	City:		State:	Zip Code:
4.	Name:	SSN/	FEID/FEIN#	Date of Birth: ☐ N/A	% of Ownership:
	Street Address:	City:		State:	Zip Code:
5.	Name:	SSN/	FEID/FEIN#	Date of Birth: ☐ N/A	% of Ownership:
	Street Address:	City:		State:	Zip Code:
6.	Name:	SSN/	FEID/FEIN#	Date of Birth: ☐ N/A	% of Ownership:
	Street Address:	City:		State:	Zip Code:
7.	Name:	SSN/	FEID/FEIN#	Date of Birth:	% of Ownership:
	Street Address:	City:		State:	Zip Code:
8.	Name:	SSN/	FEID/FEIN#	Date of Birth: ☐ N/A	% of Ownership:
	Street Address:	City:		State:	Zip Code:
	t all trade or business names used by the blicant does not use other trade or business				
cor <u>No</u> the	the applicant a subsidiary of another companies with percentages of ownership, te: A permit issued pursuant to this applicant's name and address. (If no, ple lines below).	using cation	additional sheet(s is only valid for the) if necessary). e applicant, and	
	rent Company Name		% of Ownership		

				or care, or chronic or rehab			
				ishment that is the subject			
	permit application? If so, please list the name of the company/companies providing such services below and provide the corresponding license or permit number(s) issued						
				ditional sheet(s) if necessar			
Nam		3 regulat	ory authority. (Ose aut	Permit/License No.:	Issuing Agency:		
- 10					iodag / igd.ioj.		
Sec	tion IV – Backg	round Qu					
				D QUESTIONS			
The	term "affiliated pa	rty" means	: (a) a director, officer,	trustee, partner, or committee	e member of a permittee or		
				mittee or applicant; (b) a pers			
partr	ner, shareholder, i	manager, n	nember, officer, director,	independent contractor, or e	employee of the permittee or		
				a personal information stateme			
				t or to renew a permit pursual t of the permittee or applicant.			
	_		·				
				you must provide detailed I documents. If needed, exp			
	g			,p	o oopa oo(o).		
1.	☐ Yes	□No	Has the applicant o	r any "affiliated party" (de	efined above) been found		
	If yes, explain			of adjudication), or pled n			
	in detail in			on of law that directly rela	ites to a drug, device, or		
	Section V		cosmetic?		and about bean food on		
2.	☐ Yes If yes, explain	☐ No		r any affiliated party (defir llatory agency in any state			
	in detail in			onstitute a violation of Chap			
	Section V		S. S. ISS C. IAC WOOLG CC	monato a violation of Onap	,		
3.	□Yes	☐ No	Has the applicant or	any affiliated party (define	ed above) been convicted		
	If yes, explain			cation) of any felony under	a federal, state (including		
	in detail in		Florida), or local law?	?			
	Section V		11 4	erry and a second			
4.	☐Yes	☐ No			ed above) been denied a		
	If yes, explain in detail in			any state (including Flori pters 456, 465, 499, or 893			
	Section V		regulated under Chap	picis 400, 400, 433, 01 030	, 1 .0.:		
5.	□Yes	□No	Has the applicant or	any affiliated party (define	ed above) had any current		
	If yes, explain			r license suspended or revo			
	in detail in			cal governmental agency r	elating to the manufacture		
	Section V		or distribution of drug	gs, devices, or cosmetics?			
6.	☐ Yes	□No	Has the applicant o	or any affiliated party (def	fined above) ever held a		
	If yes, explain	_			different name than the		
	in detail in		applicant's name? (If yes, provide the names			
	Section V		issued and at what a	ddress).			

Sec	Section V – Explanation(s) for "Yes" response(s) to background question(s)					
	EXPLANATION					
 						
 						
<u> </u>						
<u> </u>						
		_				
Sec	ction VI – Other Permits or Licenses					
	PERMITS OR LICENSES					
1.	Are there any permits or licenses issued by any agency of the State of Florida that authorize the purchase or possession of prescription drugs at the applicant's establishment or address? (If yes, please provide a list of all such permits including the issuing agency, the permit/license type, the permit/license number and the expiration date. If not, check the box indicating no other permits or licenses.)	☐ Yes	□No			
	licenses.). □ Permit/licensure list provided. □ No permits/licenses.					
2.	Is the applicant licensed or permitted to manufacture prescription drugs at the location of the establishment by the licensing or permitting authority in the state where the establishment is located? Yes – Resident license attached. No – Not permitted in resident state. No – Not permitted and not required to be permitted in resident state; written explanation attached with a copy of relevant regulation and/or laws showing that no permit is required.	☐ Yes	□ No			
3.	Is the applicant licensed in any other state as a manufacturer, repackager, distributor, or wholesale distributor of prescription drugs? (If yes, please provide a list all such permits including the state, the permit/license type, the permit/license number and the expiration date. If not, check the box indicating no other permits or licenses.). Permit/licensure list provided.	☐ Yes	□No			

No permits/licenses.

4.		prescription drugs into Florida? (If no, provide the name the drugs are sold into Florida in the spaces sheets if needed.)	☐ Yes ☐ No
4a.	Name	Physical Address	Florida Permit/License Number
5	Dravide information on the geta	till has antial that will ohio or physically transfer pres	ariation druge for
5.		blishment(s) that will ship or physically transfer presed the manufacturer, into or in Florida on the application	ant
5a.	Name	Address	Florida Permit/License Number
	tion VII – Prescription Drug Ma	unufacturing Activity	
Sec	tion vii – Frescription Drug Wa	indiacturing Activity	
Sec	· · ·	MANUFACTURING ACTIVITIES	
Gen	erally identify the applicant's in		t will purchase or
Gen rece	erally identify the applicant's in	MANUFACTURING ACTIVITIES Intended customers, the persons and entities that pplicant establishment after permit issuance. Wholesalers Pharmac	
Gen rece	lerally identify the applicant's in sive prescription drugs from the audanufacturers Hospitals /eterinarians Other (explain)	MANUFACTURING ACTIVITIES Intended customers, the persons and entities that pplicant establishment after permit issuance. Wholesalers	sies are Clinics
Gen rece	lerally identify the applicant's in sive prescription drugs from the audanufacturers Hospitals /eterinarians Other (explain)	MANUFACTURING ACTIVITIES Intended customers, the persons and entities that pplicant establishment after permit issuance. Wholesalers	sies are Clinics
Gen rece	nerally identify the applicant's in sive prescription drugs from the authorized Manufacturers Hospitals Veterinarians Other (explain)	MANUFACTURING ACTIVITIES Intended customers, the persons and entities that pplicant establishment after permit issuance. Wholesalers	sies are Clinics
Gen rece	derally identify the applicant's in sive prescription drugs from the applicant applica	MANUFACTURING ACTIVITIES Intended customers, the persons and entities that pplicant establishment after permit issuance. Wholesalers	cies are Clinics urer establishment ufacturer, not as a
Gen rece	lerally identify the applicant's in sive prescription drugs from the authorized Manufacturers Hospitals Veterinarians Other (explain) Intify the types of prescription drughich this establishment is considered by the constant of the const	MANUFACTURING ACTIVITIES Intended customers, the persons and entities that pplicant establishment after permit issuance. Wholesalers	cies are Clinics urer establishment ufacturer, not as a
Gen rece	Jerally identify the applicant's in sive prescription drugs from the authorized Manufacturers Hospitals Jeterinarians Other (explain) Intify the types of prescription dructify the types of prescription dructify this establishment is considered in the considered i	MANUFACTURING ACTIVITIES Intended customers, the persons and entities that pplicant establishment after permit issuance. Wholesalers	urer establishment ufacturer, not as a nufacturer, not as
Gen rece	derally identify the applicant's in sive prescription drugs from the authorized Manufacturers Hospitals Veterinarians Other (explain) Intify the types of prescription drugyhich this establishment is considered which this establishment is considered by the conside	MANUFACTURING ACTIVITIES Intended customers, the persons and entities that pplicant establishment after permit issuance. Wholesalers	urer establishment ufacturer, not as a nufacturer, not as PI or Otherwise) Otherwise) omers):
Gen rece	rerally identify the applicant's in sive prescription drugs from the authorized prescription dru	MANUFACTURING ACTIVITIES Intended customers, the persons and entities that pplicant establishment after permit issuance. Wholesalers	urer establishment ufacturer, not as a nufacturer, not as PI or Otherwise) Otherwise) omers):

Iden	Identify type of operation.					
(e.g.	FDA Drug Application Holder . NDA, ANDA, BLA, NADA, ADA holder)	Co-licensed partner FDA Drug Application H	Holder		el Manufa	cturer
Prov	Provide your Federal Food and Drug Administration (FDA) establishment registration number.					
	☐ FDA Establishment Registration	Number:				
	or					
L	No FDA Establishment Number	AND a written explanatio	n is attached	d ∐.		
1.	Are prescription drugs to be dis	stributed under this norm	eit intended '	for export?	Yes	☐ No
	(Note: A permit may be required handling prescription drugs in Flo	for Florida recipients that orida.)	t are freight	forwarders		
2.	Do you manufacture a prescriptio a separate sheet providing accur label.)				Yes	□No
3.	Does the applicant establishmen				☐ Yes	☐ No
	in the State of Florida throu contractors? (If yes, a Compl Please review sections 499.01 and	limentary Drug Distribute	or permit is			
4.	Will all required records be st			's physical	☐ Yes	☐ No
	address? (If no, provide the nam	me and address of the es	stablishments			_
4a.	required records will be stored an Name and physical address wher				L	
	Name:	10 10 4 4 11 2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	, 0.0.00.			
	Street Address:					
						
	City:		State:	Zip Cod	de (+4 opt	tional):
5.	Will the required records be comp	puterized, automated or s	tored electro	onically?	☐ Yes	☐ No
	If yes, will you have a back-up pro	ocedure to be able to pro-	vide required	d records?	☐ Yes	☐ No
	If electronically stored and not ma				☐ Yes	☐ No
	data maintained unchanged fro		, receipt, pu	urchase or		
	distribution, depending on the doc	cument type:				
6.	Section 499.0121(2), F.S., require					
	entry after hours and b) a securifacilitated or hidden by tamperin					
	description of the alarm and secu					
	the systems are monitored.	Alarm svs	etam dascrin	otion included	42 Yes	☐ No
		•	•			
	Sections 499.01(2)(a)1. and 499			otion included ers to establ		
7.	adhere to written policies and pro inventory, and distribution of pres	ocedures, which must be				
	Please provide the applicant's v	written policies and proc	edures on:	the receipt	security	storage
	inventory, distribution/disposition	n of prescription drugs;	distributing	g oldest app	proved st	ock first
4 '	(FIFO); identifying, recording a	nd reporting prescription	n drua loss	ses and the	fts: main	tenance.

	retrieval and retention of required records; prescription drug recalls and wirdisasters and other emergencies; and product tracing and other requirements Drug Supply Chain Security Act (DSCSA). Label each policy and procedure specifically identifying the subject matter in the covered by the policy or procedure. For example, the policy and procedure for labeled or identified as "Recall Policy and Procedure" or in another manner similar	under the e list above or recalls o	federal e that is could be
	Receipt, security, storage, inventory, distribution/disposition of prescription dru Distributing oldest approved stock first (FIF Identifying, recording and reporting prescription drug losses and the Maintenance, retrieval and retention of required recor Prescription drug recalls and withdraws Natural disasters and other emergence	FO) Yesefts Yeserds Yeserds Yeserds Yeserds	No No No No No
	Segregation and destruction of outdated prescription dru Temperature and humidity monitori Product Tracing and other DSCSA requirement	ıgs 🔲 Yes ing 🔲 Yes	No No
8.	Will the applicant establishment purchase and subsequently direct the distribution (e.g. drop shipment) of prescription drugs on its behalf to another company, including any prescription drug active pharmaceutical ingredient (API), for which the applicant establishment is not considered the manufacturer? (For assistance in determining the definition of "distribute" see Section 499.003, Florida Statutes.) If yes, you will need additional permit(s) depending on the activity. Refer to section 499.01(2), Florida Statutes.	Yes	□ No
9.	Will ANY prescription drugs, including active pharmaceutical ingredient, be stored, received, or warehoused – even temporarily – including, any customers' return(s) or recalled prescription drugs, at the location for which the applicant is seeking a permit? If yes, then the applicant is NOT eligible for this permit. Virtual manufacturer may not possess prescription drugs.	☐ Yes	□ No

Section VIII- Qualify as a Manufacturer

	QUALIFYING AS A MANUFACTURER (Check all that apply)	
	For the purpose of the questions below, the term "affiliate" means a business e relationship with another business entity in which, directly or indirectly: a. The business entity controls, or has the power to control, the other business entity; b. Third party controls, or has the power to control, both business entities.	-
	FDA approvals must be in the name of the applicant as listed on this application. If the is not in the same name as the applicant as listed on this application, you may manufacturer.	
1.	Does the applicant hold a FDA drug application (e.g., a New Drug Application (NDA), an Abbreviated New Drug Application (ANDA), a New Animal Drug Application (NADA), or an Abbreviated New Animal Drug Application (ANADA)) approved under the federal act? If yes, provide a list of all approved applications and licenses by number on a separate sheet with the drug's respective NDC number(s) listed with FDA, and provide copies of no more than 5 FDA approval letters. List of applications/licenses attached? Yes No Copies of approval letters attached? Yes No Does the applicant hold a Biologics License issued under s. 351 of the	☐ Yes ☐ No
۷.	Public Health Service Act, 42 U.S.C. s. 262 for a drug or biologic? If yes, provide a list of the approved licenses by number on a separate sheet, and provide a copy of no more than 5 FDA licenses for drugs or biologics. List of licenses attached? Yes No Copies of licenses attached? Yes No	
3.	Does the applicant "manufacture" drugs or biologics that are not the subject of an approved FDA application or license? If yes, please provide: a. All labeling associated with the drug or biologics manufactured and a listing of the drug's respective NDC number(s) listed with FDA by the applicant if not listed on the labeling; b. A written description of the applicant's intent with respect to the drug or biologic, i.e., clinical trial, distribution or commercial sale, etc.; c. Statement of reasoning for which the applicant claims the prescription drug can be marketed in the United States; and d. Documentation that the drug or biologic can be legally placed into interstate commerce as per FDA regulations, for example, a copy of section(s) of the Federal Register, Code of Federal Regulations (CFR) denoting the prescription drug Drug Efficacy Study Implementation (DESI) designation or a copy of section(s) of the CFR denoting the prescription drug remains pending final DESI review, or a copy and summary of material(s) and authoritative literature reviewed during the applicant's investigation supporting that the prescription drug has not yet been reviewed in the DESI process. Labeling attached? Yes No Statement of reasoning attached? Yes No Statement of reasoning attached? Yes No	☐ Yes ☐ No
4.	Is the applicant a co-licensed partner of a person described in 1, 2, or 3 above, who obtains drugs or biologics directly from a person described in 1, 2, 3 above, 5 below, or another co-licensed partner of such person? Please provide a complete, fully executed copy of no more than 5 co-licensing agreements between the applicant and the applicant's co-licensed partners. Complete agreements attached? Yes No Agreements are considered trade secret? Yes No	☐ Yes ☐ No

5.	Is the applicant an affiliate of a person describe another affiliate of such a person, that obtains from a person described in 1, 2, 3 or 4 above of person? If yes, please provide the following: a. If the applicant and the affiliate fall under the structure, i.e., one company is a parent, subsite of the other, provide written documentation destine companies, including, where applicable, the each company, an organizational chart with but but. The name and address of the manufacturer applicant obtains drugs or biologics.	drugs or biologics directly or another affiliate of such e same business / organizational diary, or sister / brother company scribing the relationships between the percentages of ownership in siness and d/b/a names; and or of the affiliate from whom the	☐ Yes ☐ No
	Relationship docun Documents are considere List of affi List of affiliates considere	ed trade secret? Yes No liates attached? Yes No	
Section IX – Affidavit			
AFFIDAVIT			
Pursuant to s. 559.79, F.S., each application for a license or renewal of a license issued by the Department of Business and Professional Regulation shall be signed under oath or affirmation by the applicant, or owner or chief executive of the applicant without the need for witnesses unless otherwise required by law.			
Pursuant to s. 559.791, F.S., any license issued by the Department of Business and Professional Regulation which is issued or renewed in response to an application upon which the person signing under oath or affirmation has falsely sworn to a material statement, including, but not limited to, the names and addresses of the owners or managers of the licensee or applicant, shall be subject to denial of the application or suspension or revocation of the license, and the person falsely swearing shall be subject to any other penalties provided by law.			
I UNDERSTAND THAT THE ISSUANCE OF A PERMIT BY THE DEPARTMENT ONLY AUTHORIZES THE APPLICANT TO CONDUCT REGULATED ACTIVITIES IN THE STATE OF FLORIDA UNDER THE NAME IN WHICH THE PERMIT IS ISSUED. IF THE PERMIT IS ISSUED IN THE NAME OF A DBA OR D/B/A THE APPLICANT MAY ONLY CONDUCT BUSINESS IN FLORIDA IN THE NAME OF THE DBA OR D/B/A.			
I FURTHER UNDERSTAND THAT PROVIDING ADDITIONAL DBA OR D/B/A NAMES TO THE DEPARTMENT AS PART OF THE APPLICATION PROCESS IS NOT, UPON LICENSURE, AN AUTHORIZATION TO CONDUCT BUSINESS IN FLORIDA UNDER THE NAME OF THOSE ADDITIONAL DBA'S OR D/B/A'S.			
I certify that I am empowered to execute this application as required by s. 559.79, F.S. I understand that my signature on this application has the same legal effect as if made under oath. To the best of my knowledge, all information contained on this application is true and correct. I understand the falsification of any information on this application may result in administrative action, including a fine, suspension, or revocation of the license.			
Sign	ature of Applicant, Owner or Chief Executive:	Date:	
		Title:	

Mail completed application to:
Department of Business and Professional Regulation
Division of Drugs, Devices and Cosmetics
2601 Blair Stone Road
Tallahassee, FL 32399-1047