DIVISION OF MEDICAL QUALITY ASSURANCE BOARD OF PHARMACY 4052 BALD CYPRESS WAY, BIN #C-04 TALLAHASSEE, FLORIDA 32399-3254 (850) 245- 4474



NONRESIDENT STERILE COMPOUNDING PERMIT APPLICATION FORNONRESIDENT PHARMACIES

JULY 2016

Nonresident Sterile Compounding Permit for Nonresident Pharmacies Information

A Nonresident Sterile Compounding Permit as authorized by Section 465.0158, *Florida Statutes* is required in order to ship, mail, deliver, or dispense in any manner, a compounded sterile product into Florida.

Definition:

For purposes of this application, when the term "affiliated person" is used, the term shall mean any person who has an ownership interest of 5% or greater in the pharmacy and any person who directly or indirectly manages, oversees, or controls the operation of the pharmacy.

Application Processing

1. Please mail the application and the \$255.00 application fee (check or money order made payable to the FLORIDA DEPARTMENT OF HEALTH) to the following address:

Department of Health Board of Pharmacy P.O. Box 6330 Tallahassee, Florida 32314-6320

OR, use the following address if you are using express mail:

Department of Health Board of Pharmacy 4052 Bald Cypress Way, Bin C-04 Tallahassee, FL 32399-3254

- **2.** Along with the application, Nonresident Pharmacies must submit the following:
 - **a.** A letter of licensure verification for both the facility and the Prescription Department Manager or Pharmacist in Charge or equivalent from the state, territory or district regulatory or licensing agency. The letter must include the original licensure date, the expiration date, and current licensure status.
 - **b.** A copy of a current inspection report from an inspection conducted by the regulatory or licensing agency of the state, territory, or district in which the applicant is located. The inspection report is current if the inspection was conducted within six months before the date of submission of this application. The current inspection report must demonstrate that the applicant is fully compliant with chapters 797, 71, 85, and 731 of the United States Pharmacopeia that are adopted in Rule 64B16-27.797(1), Florida Administrative Code.

If you are unable to submit a current inspection report demonstrating compliance with the applicable chapters of the pharmacopeia, due to acceptable circumstances as established by Rule 64B16-28.905, F.A.C. or if no current inspection has been performed, the applicant may:

 Submit a current and satisfactory inspection report from an entity approved by the board. Approved entities can be found on the Board's website at www.floridaspharmacy.gov; or

- Request the Department to perform an onsite inspection in which all costs are borne by the applicant.
- **c.** A copy of the applicant's existing policies and procedures for sterile compounding. The policies and procedures must comply with pharmaceutical standards in chapters 797, 71, 85 and 731 of the United States Pharmacopoeia.
- d. Any and all other documentation requested or mandated within this application.
- **3.** Once an application is complete and approved, the Department will issue a permit which you will receive within 7 days.

All pharmacies must answer the following questions. The questions will assist in the Board's review of your application to determine your pharmacy's compliance the applicable chapters of the United States Pharmacopeia. Please answer the following questions as completely and legibly as possible. Attach additional pages if needed.

Th	ese questions relate to your primary engineering controls.
a.	How many primary engineering controls do you have?
b.	What kind are they? (select all that apply)
	 □ Laminar Airflow Workbench (LAFW) □ Compounding Aseptic Isolator (CAI) □ Biological Safety Cabinet (BSC) □ Compounding Aseptic Containment Isolator (CACI) □ Integrated vertical clean bench □ Other: please describe
C.	Where are your primary engineering controls located? (select all that apply)
	 □ Positive Pressure ISO Class 7 buffer room with walls/doors □ Negative Pressure ISO Class 7 buffer room with walls/doors □ Positive Pressure ISO Class 7 anteroom □ Positive Pressure ISO Class 8 anteroom □ Non-ISO classed segregated compounding area for non-hazardous compounding □ Non-ISO classed containment segregated compounding room with 12 ACPH/negative pressure □ Other: please describe
d.	What was the date of the last certification of your primary and secondary engineering controls?
e.	Did the certification of the primary and (if applicable) secondary engineering controls include testing of non-viable particle counts and airflow pattern smoke testing <i>under dynamic operating conditions</i> (while pharmacy staff are working or simulating work in the area being tested)? Yes No

1.

2.	What kind of gloves and alcohol are in use at your pharmacy for sterile compounding activities?			
	Describe briefly:			
2	If your pharmacy upon inclutors (Compounding Apontic Inclutors or Compounding Apontic			
3.	If your pharmacy uses isolators (Compounding Aseptic Isolators or Compounding Aseptic Containment Isolators), describe how gloves are donned before compounding in your isolator(s).			
	☐ Not applicable because we do not use isolators for sterile compounding.			
	Describe briefly:			
	Describe briefly:			
4.	Primary engineering controls must be disinfected at frequent intervals with sterile 70% IPA during use but they also must be part of the daily cleaning routine. Briefly describe how the inside of your primary engineering controls are cleaned and disinfected (as well as the agents used) during your pharmacy's daily cleaning routine.			
5.	Before pharmacy staff or outsourced cleaning staff are allowed to perform daily and monthly cleaning activities, they must receive (at a minimum) training and competency verification in which two areas?			
	1.			
	2.			
	Describe briefly:			
6.	USP Chapter 797 requires that each compounding staff member successfully complete some training and testing before they are allowed to make compounded sterile preparations for human use. Briefly describe this type of training and testing at your facility.			

7.	These questions relate to viable air sampling. Please provide a short answer to each.					
	a.	How often does your pharmacy perform viable air sampling?				
	b.	Where is viable air sampling performed?				
		How large are the samples of air you are sampling?				
		levels?				
8.	Но	rface sampling is an environmental metric that is required "periodically" by USP Chapter 797. w is it performed at your pharmacy? Briefly describe under what conditions it is performed, how en, with what and where it is performed.				
	De	scribe briefly:				
	De	scribe briefly:				
9.		SP Chapter 797 requires gloved fingertip sampling. Briefly describe how and when your armacy performs gloved fingertip sampling.				
10		nat activities would occur at your pharmacy if the results (number of colony forming units) of one your environmental sampling samples exceeded the preselected Action Levels for that area.				
	De	scribe briefly:				
11		ease explain how the concept of "first air" is critical to executing sterile compounding with proper eptic technique.				
	De	scribe briefly:				
l						

12	If a pharmacy uses a 0.22 micron filter for the purposes of sterilization, what test is required before that batch may be released?
13	. According to USP Chapter 797, is sterility testing required if a beyond-use date of 30 days refrigerated is assigned to a medium risk level batch?
	Answer Yes or No and then briefly explain your rationale:
14	During a compounding process, the pharmacy removes the vial stopper from a product purchased from an FDA registered manufacturer. Does this change the risk level that should be assigned to the final compounded sterile product (CSP) made from that product and what risk level would you assign it?
	Answer: Yes or No then indicate the risk level you would assign this CSP and your rationale:
15	. Please describe your use of lyophilization in your pharmacy.
16	If a pharmacy has performed sterility testing on a batch (or outsourced it to a vendor who performs sterility testing in compliance with USP Chapter 71 on their behalf) and the batch fails, is it acceptable practice to retest that batch?
	Answer Yes or No and then briefly describe your rationale:



FLORIDA BOARD OF PHARMACY

P.O. Box 6330 | Tallahassee, FL 32314 (850) 245-4474 | www.floridaspharmacy.gov

NONRESIDENT STERILE COMPOUNDING APPLICATION FOR NONRESIDENT PHARMACIES

Please submit the application fee and unlicensed activity fee totaling \$255 with your application.					
Existing Nonresident Pharmacy Permit Number (If you do not have this permit, you must also submit an application for a Nonresident Pharmacy Registration.)					
	ile Compounding Permit Number (if a	applicable)			
Federal Employer Identification	Number (FEIN)				
1. Corporate Name		Telephone Number			
2. Doing Business As (d/b/a)		E-Mail Address (Optional)			
3. Mailing Address					
City	State	Zip			
4. Physical Address					
City	State	Zip			
5. Prescription Department Mar	• ,	• , , .			
Name	License No.	Start Date			
6. Contact Person	Telephone Nun	nber			
7. DEA Registration Number (If applicable)					
8. Do you have 24-hour access to patient records?YesNo (If no, please provide an explanation on a separate sheet of paper)					

9. Date of last inspection: Day Month Year							
Inspecting Authority							
the United States Pharmacop and your policies and proced	10. Was this inspection structured to ensure compliance with Chapters 797, 71, 85, and 731 of the United States Pharmacopeia? (Attach a copy of the inspection report, the floor plan and your policies and procedures manual). YesNo						
11. Prescription Department Oper Monday – Friday: Open:	•	12. Toll-Free Telephone Number (Available 6 days a week for 40 hours)				
	_ Close:	()					
13. Ownership Information							
 a. Type of Ownership IndividualCorporationPartnershipOther: CORPORATIONS & LIMITED PARTNERSHIPS: INCLUDE A COPY OF THE ARTICLES OF INCORPORATION ON FILE WITH THE STATE WHERE THE FACILITY IS LOCATED. b. List each principal, officer, agent, managing employee or affiliated person of the applicant. 							
Attach a separate sheet if necessary.							
Name/Title Da	ate of Birth / / / / / / / /	Mailing Address, City State, Zip Code	% Ownership % %				
Questions 14 through 18 are required pursuant to Section 456.0635(2), <i>Florida Statutes</i> . Please explain any "yes" answered to the following questions on a separate sheet, providing as much detail as possible. Supporting documentation must include at a minimum the official charging document and the official judgment and sentence.							
14. Has the applicant or any principal, officer, agent, managing employee, or affiliated person of the applicant been convicted of, or entered a plea of guilty or nolo contendere to, regardless of adjudication, a felony under Chapter 409, Chapter 817, or Chapter 893, Florida Statutes or a similar felony offense committed in another state or jurisdiction? (If "no", skip to question 15.)							
Yes No							

	state or jur	isdiction), ha	irst or second degree (or the equivalent level of felony in as it been more than 15 years from the date of the plea, y subsequent probation?
Yes		No	
state or j and com of the th	jurisdiction pletion of a ird degree), has it beei any subsequ	third degree (or the equivalent level of felony in another in more than 10 years from the date of the plea, sentence lent probation? This question does not apply to felonies on 893.13(6)(a), Florida Statutes or a similar felony offense urisdiction.
Yes		No	
state or j	urisdiction) committed	under Section in another s	hird degree (or the equivalent level of felony in another on 893.13(6)(a), Florida Statutes or a similar felony tate or jurisdiction has it been more than 5 years from nd completion of any subsequent probation?
Yes		No	
If "voo"	1 41		
person c	of the applic	cant success	
person c	of the application the felony of	cant success	sfully completed a drug court program that resulted in the g withdrawn or the charges dismissed?
yesHas the apregardle	applicant of policing to pullicant of adjudating to pu	No r any principen convicted ication to a f	
yesHas the a of the ap regardle 1396 (rel	applicant of policing to put a ting to put a 16.)	No r any principen convicted ication to a fallowing to the second control of the	sfully completed a drug court program that resulted in the g withdrawn or the charges dismissed? ———————————————————————————————————
yesHas the appregardle question yes If "yes",	applicant of policing to pure attention to pure	no r any principen convicted ication to a fiblic health, who of application	sfully completed a drug court program that resulted in the g withdrawn or the charges dismissed? Dal, officer, agent, managing employee, or affiliated person of, or entered a plea of guilty or nolo contendere to, felony under 21 U.S.C. ss. 801-970 or 42 U.S.C. ss. 1395-welfare, Medicare and Medicaid issues)? (If "no", skip to
yesHas the appregardle question yes If "yes",	applicant of policant of policant been at the date ent period	no r any principen convicted ication to a fabric health, v	sfully completed a drug court program that resulted in the g withdrawn or the charges dismissed? Dal, officer, agent, managing employee, or affiliated person of, or entered a plea of guilty or nolo contendere to, felony under 21 U.S.C. ss. 801-970 or 42 U.S.C. ss. 1395-welfare, Medicare and Medicaid issues)? (If "no", skip to (If yes, explain on a separate sheet providing accurate details) On more than 15 years after the sentence and any nended?

16.	Has the applicant or any principal, officer, agent, managing employee, or affiliated person of the applicant ever been terminated for cause from the Florida Medicaid Program pursuant to Section 409.913, Florida Statutes? (If "no", skip to question 17.)				
	Yes	No			
	• • •	or any principal, officer, agent, neen reinstated and in good standi ent five years?	.		
	Yes	No			
17.	. Has the applicant or any principal, officer, agent, managing employee, or affiliated person of the applicant ever been terminated for cause, pursuant to the appeals procedures established by the state or from any other state Medicaid program? (If "no", skip to question 18)				
	Yes	No			
		or any principal, officer, agent, neen in good standing with a state			
	Yes	No			
	If "yes", did the termination	on occur at least 20 years prior to	the date of this application?		
	Yes	No			
18.	the applicant listed on the	ncipal, officer, agent, managing e United States Department of Hea f Excluded Individuals and Entitie	olth Human Services Office of		
	Yes No				
	If "yes", are you listed bed	ause you defaulted or are delinqu	uent on a student loan?		
	Yes No				
	If "yes", is the student loa	n default or delinquency the only	reason you are listed on the		
	Yes No				
	100				
19.	permit type, and permit nu	ed or permitted in any other state umber for each permit. Attach a s			
163_	 State	 Permit Type	Permit Number		
		- 7 -			

20.	Has the applicant or any principal, officer, agent, managing employee, or affiliated person ever owned a pharmacy? If yes, provide the name of the pharmacy, the state where the pharmacy is located and the status of the pharmacy.				
	Yes	No(If yes, please list them b	elow, you may provide additional sheet)		
	Pharmacy Name	State	Status		
21.	issued to the applicant or person of the applicant in Yes No	n ever been taken against any lice any principal, officer, agent, man this state or any other? (If yes, explain on a separate sheagency who took the disciplinary action)	aging employee, or affiliated		
22.	ever been convicted of a formation of the second se	agent, managing employee, affiliated and selony or misdemeanor, excluding a line of the control	minor traffic convictions? elonies, even if adjudication was		
		ense for the purposes of this question.)	ing under the influence or driving while		
23.	Is there any other permit is location address on this a	ssued by the Department of Healt pplication?	h located at the physical		
	Yes No	(If yes, explain on a separate she	et providing accurate details)		
24.	Does the applicant or any principal, officer, agent, managing employee, or affiliated person of the applicant have any outstanding fines, liens or overpayments assessed by a final order of the department?				
	Yes No	(If yes, explain on a separate she	et providing accurate details)		
	• • •	nt or any principal, officer, agent, plicant have a repayment plan ap			
	Yes No				
25. insp	Has the applicant rece ection conducted by the FD	eived an FDA Form 483 or OA within the last 3 years?	Warning Letter following an		
	action plan, and supporting docum	(If yes, please submit the Form a nentation demonstrating how the corrective clude but is not limited to pictures, facility o	e action plan was implemented.		

APPLICANT SIGNATURE PAGE

Florida law requires that applicants supplement their applications as needed to reflect any material change in any circumstances or conditions stated in the application that takes place between the initial filing of the application and the final grant or denial of the license, which might affect the decision of the department of board.

I, the undersigned, certify that the statements contained in this application are true, complete, and correct					
I, the undersigned, certify that the statements contained in this application are true, complete, and correct and I agree that said statements shall form the basis of my application. I do authorize the Florida Board of Pharmacy and the Department to make any investigations that they deem appropriate and to secure any additional information concerning the applicant or me. I further authorize them to furnish any information they may have or have in the future concerning me to any person, corporation, institution association, board, or any municipal, county, state, or federal governmental agencies or units. I understand according to the Florida Board of Pharmacy Statutes that a Pharmacy Permit may be denied, revoked or suspended for presenting any false, fraudulent, or forged statement, certificate, diploma, or other thing, in connection with an application for a license or permit.					
I, the undersigned, hereby acknowledge that providing false information in relation to this application, may result in denial of licensure, discipline, and/ or criminal penalties pursuant to sections 456.067, 465.015 (5), 775.082, 775.083, and 775.084, <i>Florida Statutes</i> .					
I, the undersigned, have completely reviewed and read the foregoing document and state that the facts stated in it are true.					
SIGNATURE					



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ATTESTATION

Section 465.0158(3)(c), F.S., requires that applicants submit a written attestation by an owner or officer of the applicant and by the applicant's Prescription Department Manager (PDM) or Pharmacist In Charge (PIC).

I hereby attest that I have read and understand the laws and rules governing sterile compounding in the State of Florida, and that any sterile compounded product shipped, mailed, delivered, or dispensed into the State of Florida from our facility meets or exceeds the standards for sterile compounding set by the State of Florida and has not been compounded in violation of the laws and rules of the state, territory, or district in which our facility is located.

I declare tha	at I have read the for	egoing Attestation and th	nat the facts stated in it a	are true
SIGNATURE	(Owner/Officer)	TITLE	DATE	
SIGNATURE	(PDM/PIC of Sterile Comp	TITLE	DATE	