

AUTHORIZED USER TRAINING, EXPERIENCE, AND PRECEPTOR ATTESTATION

(for uses defined under 35.300) [10 CFR 35.57, 35.390, 35.392, 35.394, and 35.396] APPROVED BY OMB: NO. 3150-0120 EXPIRES: (MM/DD/YYYY)

Name of Proposed Authorized User			State or Territory Where Licensed					
Reque	ested Aut	thorization(s) (check all that apply)	:					
	35.300 Use of unsealed byproduct material for which a written directive is required							
OF	₹							
35.300 Oral administration of sodium iodide 1.22 gigabecquerels (33 millicuries)			I-131 requiring a written directive in quantities less than or equal to					
	35.300 Oral administration of sodium iodide I-131 requiring a written directive in quantities greater than 1.22 gigabecquerels (33 millicuries)							
	35.300 Parenteral administration of any radioactive drug that contains a radionuclide that is primarily used for its electron emission, beta radiation characteristics, alpha radiation characteristics, or photon energy of less than 150 keV, for which a written directive is required.							
PART I TRAINING AND EXPERIENCE (Select one of the three methods below)								
da tra	* Training and Experience, including board certification, must have been obtained within the 7 years preceding the date of application or the individual must have related continuing education and experience since the required training and experience was completed. Provide dates, duration, and description of continuing education and experience related to the uses checked above.							
1.	Board	Certification						
а	. Provid	le a copy of the board certification.						
b	 For 35.390, provide documentation on supervised case experience. The table in section 3.c. may be used to document this experience. 							
С	superv	For 35. 396 , provide documentation on classroom and laboratory training, supervised work experience, and supervised clinical case experience. The tables in sections 3.a., 3.b., and 3.c. may be used to document this experience. Skip to and complete Part II Preceptor Attestation.						
d	. For a board certification issued on or before October 24, 2005 that is listed in 10 CFR 35.57(b)(2)(ii), provide the following:							
	(i) D	ocumentation that the individual p	erformed each use checked above on or before October 24, 2005.					
	(ii) Dates, duration, and description of continuing education and experience within the past seven years for each use checked above.							
е	. Stop h	iere.						
2 .	Curren	nt 35.300, 35.400, or 35.600 Auth	orized User Seeking Additional Authorization					
a.	Authori	ized User on Materials License	under the requirements below or					
	equiva	alent Agreement State requirement	s (check all that apply):					
	<u> </u>	35.390 35.392 3	35.394 35.490 35.690					
b.	supervi certified	ised case experience. The table in	ical uses under 35.300, provide documentation on additional required in section 3.c. may be used to document this experience. If board and stop here. If not board certified then provide completed Part II					

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c. If currently authorized under 35.490 or 35.690 and requesting authorization for 35.396, provide documentation on classroom and laboratory training, supervised work experience, and supervised clinical case experience. The tables in sections 3.a., 3.b., and 3.c. may be used to document this experience. Also provide completed Part II Preceptor Attestation. 3. Training and Experience for Proposed Authorized User 35.390 a. Classroom and Laboratory Training 35.392 35.394 35.396 Dates of Clock Description of Training Location of Training Hours Training* Radiation physics and instrumentation Radiation protection Mathematics pertaining to the use and measurement of radioactivity Chemistry of byproduct material for medical use Radiation biology **Total Hours of Training:** 35.390 35.392 35.394 b. Supervised Work Experience 35.396 (If more than one supervising individual is necessary to document supervised training, provide multiple copies of this page.) **Supervised Work Experience** Total Hours of Experience: Description of Experience Location of Experience/License or Dates of Confirm Permit Number of Facility Must Include: Experience* Ordering, receiving, and Yes unpacking radioactive materials safely and performing the No related radiation surveys Performing quality control procedures on instruments Yes used to determine the activity of dosages and performing No checks for proper operation of survey meters Calculating, measuring, and Yes safely preparing patient or human research subject No dosages Using administrative controls to Yes prevent a medical event involving the use of unsealed No byproduct material Using procedures to contain Yes spilled byproduct material safely and using proper No decontamination procedures

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. Supervised Work Experience	(continued)					
Supervising Individual		License/Permit Number listing supervising individual as an authorized user				
Supervising individual meets the check all that apply)**:	e requirements below,	or equivalent Agreement State requirements				
 ☐ 35.390 With experience administering dosages of: ☐ 35.392 ☐ Oral Nal-131 requiring a written directive in quantities less than or equal to 1.22 gigabecquerels (33 millicuries) ☐ 35.394 ☐ Oral Nal-131 in quantities greater than 1.22 gigabecquerels (33 millicuries) ☐ 35.396 ☐ Parenteral administration of any radioactive drug that contains a radionuclide that is primarily used for its electron emission, beta radiation characteristics, alpha radiation characteristics, or photon energy of less than 150 keV, for which a written directive is required. 						
 Supervising Authorized User must had individual requesting authorized user 	ave experience in administerstatus.	ering dosages in the same dosage category or categories	as the			
 Supervised Clinical Case Exp if more than one supervising individ this page. 		nment supervised work experience, provide multiple	copies of			
Description of Experience	Number of Cases Involving Personal Participation	Location of Experience/License or Permit Number of Facility	Dates of Experience*			
Oral administration of sodium odide I-131 requiring a written lirective in quantities less than or equal to 1.22 gigabecquerels 33 millicuries)						
Oral administration of sodium odide I-131 requiring a written directive in quantities greater han 1.22 gigabecquerels (33 millicuries)						
Parenteral administration of any radioactive drug that contains a radionuclide that is primarily used for its electron emission, beta radiation characteristics, alpha radiation characteristics, or photon energy of less than 150 keV, for which a written directive is required.						

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3. Training and Experience for Proposed Authorized User (continued)								
c. Supervised Clinical Case Experience (continued)								
Supervising Individual	License/Permit Number listing supervising individual as an authorized user							
Supervising individual mosts the requirements below or equiv	plant Agracoment State requirements (check all that apply)**							
Supervising individual meets the requirements below, or equiva								
☐ 35.390 With experience administering dosages of:								
Oral Nal-131 requiring a written directive in quantities less than or equal to 1.22 gigabecquerels (33 millicuries)								
35.394 Oral Nal-131 in quantities greater than	1.22 gigabecquerels (33 millicuries)							
	diation characteristics, alpha radiation characteristics, or or which a written directive is required.							
** Supervising Authorized User must have experience in administering dosages in the same dosage category or categories as the individual requesting authorized user status.								
d. Provide completed Part II Preceptor Attestation.								
DART II _ DRECE	PTOR ATTESTATION							
Note: This part must be completed by the individual's preceptor. The preceptor does not have to be the supervising individual as long as the preceptor provides, directs, or verifies training and experience required. If more than one preceptor is necessary to document experience, obtain a separate preceptor statement from each.								
By checking the boxes below, the preceptor is not	attesting to the individual's "general clinical competency."							
First Section Check one of the following for the requested authorization:								
For 35.390:								
I attest that	has satisfactorily completed the 700 hours of training							
Name of Proposed Authorized User and experience, including a minimum of 200 hours of classroom and laboratory training, as required by 10 CFR 35.390 (b)(1).								
For 35.392:								
I attest that Name of Proposed Authorized User	has satisfactorily completed the 80 hours of classroom							
and laboratory training, as required by 10 CFR 35.392(c)(1), and the supervised work and clinical case experience required in 35.392(c)(2).								
For 35.394:								
☐ I attest that	has satisfactorily completed the 80 hours of classroom							
Name of Proposed Authorized User								
and laboratory training, as required by 10 CFR 35.394 (c)(1), and the supervised work and clinical case experience required in 35.394(c)(2).								

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Second Section										
I attest that	has satisfactorily completed the required clinical case									
Name of Proposed Authorized User										
experience required in 35	experience required in 35.390(b)(1)(ii)G listed below:									
	Oral Nal-131 requiring a written directive in quantities less than or equal to 1.22 gigabecquerels (33 millicuries)									
Oral Nal-131 in quanti	Oral Nal-131 in quantities greater than 1.22 gigabecquerels (33 millicuries)									
Parenteral administration of any radioactive drug that contains a radionuclide that is primarily used for its electron emission, beta radiation characteristics, alpha radiation characteristics, or photon energy of less than 150 keV, for which a written directive is required.										
Third Section										
I attest that	is able to independently fulfill the radiation safety-related									
	f Proposed Authorized User									
duties as an authorized us	ser for the medical uses authorized under 10 CFR 35.300 for:									
Oral Nal-131 requiring gigabecquerels (33 mi	a written directive in quantities less than or equal to 1.22 llicuries)									
Oral Nal-131 in quanti	ties greater than 1.22 gigabecquerels (33 millicuries)									
Parenteral administration of any radioactive drug that contains a radionuclide that is primarily used for its electron emission, beta radiation characteristics, alpha radiation characteristics, or photon energy of less than 150 keV, for which a written directive is required.										
Fourth Section For 35.396:										
Current 35.490 or 35.690	authorized user:									
I attest that	is an authorized user under 10 CFR 35.490 or 35.690									
	f Proposed Authorized User									
laboratory training, as req experience required by 35	or equivalent Agreement State requirements, has satisfactorily completed the 80 hours of classroom and laboratory training, as required by 10 CFR 35.396 (b)(1), and the supervised work and clinical case experience required by 35.396(b)(2), and is able to independently fulfill the radiation safety-related duties as an authorized user under 10 CFR 35.300 for:									
Parenteral administration of any radioactive drug that contains a radionuclide that is primarily used for its electron emission, beta radiation characteristics, alpha radiation characteristics, or photon energy of less than 150 keV, for which a written directive is required.										
OR										
Board Certification:										
I attest that	has satisfactorily completed the board certification									
Name of Proposed Authorized User										
training required by 10	6(a)(3), has satisfactorily completed the 80 hours of classroom and laboratory CFR 35.396 (b)(1) and the supervised work and clinical case experience required by ble to independently fulfill the radiation safety-related duties as an authorized user									

under 10 CFR 35.300 for:

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Fifth Section									
Complete one of the following for the attestation and signature:									
Authorized User									
I meet the requirements below, or equivalent Agreement State requirements, as an authorized user for:									
	35	5.57 for 35.300 uses							
I have experience administering dosages in the following categories for which the proposed Authorized User is requesting authorization:									
Oral Nal-131 requiring a written directive in quantities less than or equal to 1.22 gigabecquerels (33 millicuries)									
Oral Nal-131 in quantities greater than 1.22 gigabecquerels	Oral Nal-131 in quantities greater than 1.22 gigabecquerels (33 millicuries)								
Parenteral administration of any radioactive drug that contains a radionuclide that is primarily used for its electron emission, beta radiation characteristics, alpha radiation characteristics, or photon energy of less than 150 keV, for which a written directive is required.									
OR									
Residency Program Director:									
I affirm that the attestation represents the consensus of the residency program faculty where at least one faculty member is an authorized user who meets the requirements below or equivalent Agreement State requirements:									
☐ 35.390 ☐ 35.392 ☐ 35.394 ☐ 35.39	96	35.57 for 35.300 uses							
I affirm that this facility member has experience in administering dosages in the same dosage category or categories for which the individual is requesting authorized user status and concurs with the attestation I am providing as program director.									
☐ I affirm that the residency training program is approved by the:									
Residency Review Committee of the Accreditation Council	for Gradua	te Medical Education							
Royal College of Physicians and Surgeons of Canada									
Council on Post-Graduate Training of the American Osteo	pathic Asso	ciation							
I affirm that the residency training program includes training and experience specified in:									
□ 35.390 □ 35.392 □ 35.394 □ 35.39	96								
Name of Facility:	License/Per	mit Number:							
Name of Preceptor or Residency Program Director (Typed or Printed)		Telephone Number	Date						
Signature		I							

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