NRC FORM 313A (AUT) U.S. NUCLEAR REGULATORY COMMISSION						
				APPROVED BY OMB: NO. 3150-0120 EXPIRES: MM/DD/YYYY		
Name of	Propose	d Authorized User		State or Territory Where Licen	sed	
Reques	ted Aut	norization(s) (check all that apply):	:			
3	35.300	Use of unsealed byproduct mate	erial for whic	ch a written directive is requi	red	
OR						
3	35.300	Oral administration of sodium ioc 1.22 gigabecquerels (33 millicuri		equiring a written directive in	quantities less than or equal to	
3	35.300	Oral administration of sodium ioc gigabecquerels (33 millicuries)	dide I-131 re	equiring a written directive in	quantities greater than 1.22	
3	35.300	Parenteral administration of any than 150 keV for which a written			clide with a photon energy less	
3	35.300	Parenteral administration of any	other radio	nuclide for which a written di	rective is required	
				G AND EXPERIENCE hree methods below)		
applicat experier	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.					
<u> </u>	Board (Certification				
a.	Provide	a copy of the board certification.				
		390, provide documentation on su I to document this experience.	upervised cl	linical case experience. The	table in section 3.c. may	
	c. 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.					
d.	Skip to	and complete Part II Preceptor At	ttestation.			
<u> </u>	Current	35.300, 35.400, or 35.600 Autho	orized Use	r Seeking Additional Autho	prization	
		zed User on Materials License			der the requirements below or	
	equival	ent Agreement State requirement	s (check all	that apply):		
	35.	390 35.392 3	35.394	35.490 35.6	90	
	require	ntly authorized for a subset of clin d supervised case experience. The nce. Also provide completed Part	he table in s	section 3.c. may be used to a		
	docum case ex	ntly authorized under 35.490 or 35 entation on classroom and laborat sperience. The tables in sections ovide completed Part II Preceptor	tory training 3.a., 3.b., a	l, supervised work experience and 3.c. may be used to docu	e, and supervised clinical	

FORM 313A (AUT) ⁽⁶⁾ AUTHORIZED USER TRAIN	U.S. NU	UCLEAR REGULA					
3. Training and Experience for			,				
a. Classroom and Laboratory Tr		.394	35.396				
Description of Training	Location of Training	Clock Hours	Dates of Training*				
Radiation physics and instrumentation							
Radiation protection							
Mathematics pertaining to the use and measurement of radioactivity							
Chemistry of byproduct material for medical use							
Dediction biology							
Radiation biology		Total Hours of Training:					
	Total Hours of Training:						
b. Supervised Work Experience	35.390 35.392 35. individual is necessary to document supervised train	ining, provide m					
b. Supervised Work Experience If more than one supervising	35.390 35.392 35.		Dates of				
of this page.	individual is necessary to document supervised train	ining, provide m	nultiple copies				
 b. Supervised Work Experience If more than one supervising of this page. Description of Experience Ordering, receiving, and unpacking radioactive materials safely and performing the related radiation surveys Performing quality control procedures on instruments used to determine the activity of dosages and performing checks for proper operation of 	individual is necessary to document supervised train	ining, provide m	Dates of				
 b. Supervised Work Experience If more than one supervising of this page. Description of Experience Ordering, receiving, and unpacking radioactive materials safely and performing the 	individual is necessary to document supervised train	ining, provide m	Dates of				
 b. Supervised Work Experience If more than one supervising of this page. Description of Experience Ordering, receiving, and unpacking radioactive materials safely and performing the related radiation surveys Performing quality control procedures on instruments used to determine the activity of dosages and performing checks for proper operation of survey meters Calculating, measuring, and safely preparing patient or human research subject 	individual is necessary to document supervised train	ining, provide m	Dates of				

C FORM 313A (AUT)	U.S. NUCLEAR REGULATORY COMMISSION			
AUTHORIZED USER TRAINING AND EXPERIENCE AND PRECEPTOR ATTESTATION (continued)				
. <u>Training and Experience for Proposed Authorized User</u> (continued)				
b. Supervised Work Experience (continued)				
Supervising Individual	License/Permit Number listing supervising individual as an authorized user			
Supervising individual meets the requirements below, or equivalent Agreement State requirements (check all that apply)**:				
gigabecquerels (33 millicu	written directive in quantities less than or equal to 1.22 uries)			
Parenteral administration of be	greater than 1.22 gigabecquerels (33 millicuries)			
	of beta-emitter, or photon-emitting radionuclide with a photon requiring a written directive is required			
Parenteral administration	of any other radionuclide requiring a written directive			
 ** Supervising Authorized User must have experience requesting authorized user status. 	e in administering dosages in the same dosage category or categories as the individual			

c. Supervised Clinical Case Experience If more than one supervising individual is necessary to document supervised work experience, provide multiple copies of this page.

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 iodide I-131 requiring a written directive in quantities less than or equal to 1.22 gigabecquerels (33 millicuries)			
Oral administration of sodium iodide I-131 requiring a written directive in quantities greater than 1.22 gigabecquerels (33 millicuries)			
Parenteral administration of any beta-emitter, or photon-emitting radionuclide with a photon energy less than 150 keV for which a written directive is required			
Parenteral adminstration of any other radionuclide for which a written directive is required			
(List radionuclides)			

NRC (10-20	FORM 313A (AUT)	U.S. NUCLEAR REGULATORY COMMISSION				
(10-20-	*	AINING AND EXPERIENCE AND PRECEPTOR ATTESTATION (continued)				
3.	3. Training and Experience for Proposed Authorized User (continued)					
	c. Supervised Clinical Case E	xperience (continued)				
	Supervising Individual	License/Permit Number listing supervising individual as an authorized user				
	Supervising individual meets th apply)**:	he requirements below, or equivalent Agreement State requirements (check all that				
	35.390 With experience	e administering dosages of:				
		1 requiring a written directive in quantities less than or equal to 1.22 rels (33 millicuries)				
		1 in quantities greater than 1.22 gigabecquerels (33 millicuries)				
	Parenteral a	administration of beta-emitter, or photon-emitting radionuclide with a photon than 150 keV requiring a written directive is required				
	Parenteral a	administration of any other radionuclide requiring a written directive				
	** Supervising Authorized User must requesting authorized user status	t have experience in administering dosages in the same dosage category or categories as the individual				
	d. Provide completed Part II F	Preceptor Attestation.				
		PART II – PRECEPTOR ATTESTATION				
Note	individual as long as the pro-	ed by the individual's preceptor. The preceptor does not have to be the supervising eceptor provides, directs, or verifies training and experience required. If more than to document experience, obtain a separate preceptor statement from each.				
	st Section eck one of the following for ea	ich requested authorization:				
	<u>For 35.390:</u>					
	Board Certification					
	I attest that	has satisfactorily completed the training and experience				
	requirements in 35.390	(a)(1).				
	OR					
	Training and Experience					
	I attest that	has satisfactorily completed the 700 hours of training				
		ne of Proposed Authorized User				
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(10-2006) AUTHORIZED U	ISER TRAINING AND EXPERIE	NCE AND PRECEPTOR ATTESTATION (continued)		
Preceptor Attestation	continued)			
First Section (continu	ued)			
For 35.392 (Identical	Attestation Statement Regard	ess of Training and Experience Pathway):		
I attest that		has satisfactorily completed the 80 hours of classroom		
_	Name of Proposed Authorized User			
-	training, as required by 10 CFR 3 uired in 35.392(c)(2).	5.392(c)(1), and the supervised work and clinical case		
For 35.394 (Identical	Attestation Statement Regard	ess of Training and Experience Pathway):		
I attest that		has satisfactorily completed the 80 hours of classroom		
	Name of Proposed Authorized User			
	raining, as required by 10 CFR 3 uired in 35.394(c)(2).	5.394 (c)(1), and the supervised work and clinical case		
Second Section				
I attest that		has satisfactorily completed the required clinical case		
	Name of Proposed Authorized User			
experience requ	uired in 35.390(b)(1)(ii)G listed be	low:		
	1 requiring a written directive in q rels (33 millicuries)	uantities less than or equal to 1.22		
Oral Nal-13	1 in quantities greater than 1.22 g	jigabecquerels (33 millicuries)		
	Parenteral administration of beta-emitter, or photon-emitting radionuclide with a photon energy less than 150 keV requiring a written directive is required			
Parenteral a	dministration of any other radion	uclide requiring a written directive		
Third Section				
I attest that		has satisfactorily achieved a level of competency to		
	Name of Proposed Authorized User			
function indepe	ndently as an authorized user for	:		
	1 requiring a written directive in q rels (33 millicuries)	uantities less than or equal to 1.22		
Oral Nal-13	1 in quantities greater than 1.22 g	jigabecquerels (33 millicuries)		
	dministration of beta-emitter, or p than 150 keV requiring a written	photon-emitting radionuclide with a photon directive is required		
Parenteral a	dministration of any other radion	uclide requiring a written directive		

NRC FORM 313A (AUT) (10-2006)			U.S. NUCLEAR REGULA	ATORY COMMISSION	
	INING AND EXPERI	ENCE AND PRECEPTO	OR ATTESTATION (co	ontinued)	
Fourth Section					
<u>For 35.396:</u>					
Current 35.490 or 35.690 at	<u>ithorized user:</u>				
I attest that	of Proposed Authorized User	is an authorized us	er under 10 CFR 35.4	90 or 35.690	
laboratory training, as rec experience required by 3	or equivalent Agreement State requirements, has satisfactorily completed the 80 hours of classroom and laboratory training, as required by 10 CFR 35.396 (d)(1), and the supervised work and clinical case experience required by 35.396(d)(2), and has achieved a level of competency sufficient to function independently as an authorized user for:				
	Parenteral administration of any beta-emitter, or photon-emitting radionuclide with a photon energy less than 150 keV for which a written directive is required				
Parenteral adminstrat	on of any other radio	nuclide for which a writte	en directive is required		
		OR			
Board Certification:					
I attest that		has satisfactorily c	ompleted the board ce	rtification	
Name of Proposed Authorized User requirements of 35.396(c), has satisfactorily completed the 80 hours of classroom and laboratory training required by 10 CFR 35.396 (d)(1) and the supervised work and clinical case experience required by 35.396(d)(2), and has achieved a level of competency sufficient to function independently as an authorized user for:					
	Parenteral administration of any beta-emitter, or photon-emitting radionuclide with a photon energy less than 150 keV for which a written directive is required				
Parenteral adminstrat	Parenteral adminstration of any other radionuclide for which a written directive is required				
Fifth Section Complete the following for prece	ptor attestation and	signature:			
I meet the requirements below, or equivalent Agreement State requirements, as an authorized user for:					
35.390 35.392	35.394	35.396			
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 (33 millicuries)					
Parenteral administration of beta-emitter, or photon-emitting radionuclide with a photon energy less than 150 keV requiring a written directive is required					
Parenteral administration of any other radionuclide requiring a written directive					
Name of Preceptor	Signature		Telephone Number	Date	
License/Permit Number/Facility Name	1		1	- 1	