NRC FORM 313A (AUT) U.S. NUCLEAR REGULATORY COMMISSION							
(06-2016) AUTHORIZED USER TRAIN AND PRECEPTOR (for uses defined [10 CFR 35.390, 35.392]					TESTATION der 35.300)		APPROVED BY OMB: NO. 3150-0120 EXPIRES: 06/30/2019
Nam	ne of Propos	ed Authoriz	ed User		State or Territory	Where License	ed
Rec	uested Aut	horization(	(s) (check all that	t apply):			
[	35.300	Use of ur	nsealed byproduc	ct material for whi	ch a written direc	ctive is require	ed
0	DR						
[	35.300		ninistration of sod abecquerels (33 r		equiring a writter	n directive in	quantities less than or equal to
	35.300		ninistration of sod Juerels (33 millicu		equiring a writter	n directive in	quantities greater than 1.22
	35.300			of any beta-emitte written directive is		itting radionu	clide with a photon energy less
[	35.300	Parenter	al administration	of any other radio	onuclide for which	n a written dir	ective is required
					NING AND EXP		
*	* 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.						rience since the required
	1. <u>Board</u>	<u>Certificati</u>	<u>on</u>				
	a. Provide	a copy of	the board certific	ation.			
			de documentatior It this experience		inical case expe	rience. The t	able in section 3.c. may
	c. For 35.396, provide documentation on classroom and laboratory training, supervised work experie 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 compl	lete Part II Prece	ptor Attestation.			
	2. <u>Curren</u>	<u>t 35.300, 3</u>	35.400, or 35.600	) Authorized Use	er Seeking Addi	tional Autho	rization
	a. Authori	zed User c	on Materials Licer	nse		under	the requirements below or
	equiva	lent Agree	ment State requi	rements (check al	ll that apply):		
	35.	.390	35.392	35.394	35.490	35.69	0
	required se	upervised of	case experience.	of clinical uses un . The table in sec Part II Preceptor A	tion 3.c. may be		tation on additional ment this
	document case expe	ation on clarience. Th	assroom and lab	ons 3.a., 3.b., and	upervised work e	experience, a	.396, provide nd supervised clinical nt this experience.

I

<ol> <li>Training and Experience for a. Classroom and Laboratory Tr</li> </ol>	-	ed Authorizo		35.392		35.394		35.396
								Dates of
Description of Training		Locati	on of Trai	ning			lours	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 H	ours of Trai	ning:					
<ul> <li>b. Supervised Work Experience If more than one supervising of this page.</li> </ul>		35.390 35.390		35.392 ment sup		85.394 raining,		35.396 nultiple copies
If more than one supervising	g individua	al is necessar			ervised t	raining,	provide n	
of this page.	g individua ork Exper	is necessar ience	ry to docu	Total Ho	ervised t	raining, perience	provide n	nultiple copies
If more than one supervising of this page. Supervised Wo Description of Experience Must Include: Ordering, receiving, and unpacking radioactive materials safely and performing the	g individua	is necessar ience	y to docu	Total Ho	ervised t	contraining, contr	provide n	nultiple copies
If more than one supervising of this page. Supervised Wo Description of Experience Must Include: 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	g individua	is necessar ience	y to docu	Total Ho	ervised t	contraining, perience	provide n e: onfirm Yes	nultiple copies
If more than one supervising of this page. Supervised Wo Description of Experience Must Include: 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	g individua	is necessar ience	y to docu	Total Ho	ervised t	contraining, perience	provide n e: onfirm Yes No Yes	nultiple copies
If more than one supervising of this page. Supervised Wo Description of Experience	g individua	is necessar ience	y to docu	Total Ho	ervised t	contraining, perience	provide n e: onfirm Yes No Yes No Yes	nultiple copies

<b>FORM 313A (AUT)</b>		U.S. NUCLEAR REGULA	TORY COMMISSION
,	NING AND EXPERIE	NCE AND PRECEPTOR ATTESTATION (co	ntinued)
Training and Experience for P	roposed Authorized	User (continued)	
b. Supervised Work Experience	e (continued)		
Supervising Individual	License/Permit Number listing supervising individual as a a authorized user		
Supervising individual meets the apply)**:	requirements below,	, or equivalent Agreement State requirements	(check all that
35.390 With experience a	administering dosage	s of:	
	requiring a written dir ls (33 millicuries)	ective in quantities less than or equal to 1.22	
Oral Nal-131		han 1.22 gigabecquerels (33 millicuries)	
Parenteral ad		mitter, or photon-emitting radionuclide with a a written directive is required	photon
		her radionuclide requiring a written directive	
		tering dosages in the same dosage category or categorie	s as the individual
requesting authorized user status.		tering usages in the same usage category of categorie	
multiple copies of this page. Description of Experience	Number of Cases Involving Personal	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)	Participation		
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 administration of any other radionuclide for which a written directive is required			
(List radionuclides)			

NRC FORM 313A (AUT) (06-2016)	U.S. NUCLEAR REGULATORY COMMISSION						
AUTHORIZED USER TRAINING AND EXPER	RIENCE AND PRECEPTOR ATTESTATION (continued)						
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 meets the requirements belo apply)**:	ow, or equivalent Agreement State requirements (check all that						
35.390 With experience administering dosa	iges of:						
☐ 35.392 ☐ Oral Nal-131 requiring a written gigabecquerels (33 millicuries)	directive in quantities less than or equal to 1.22						
$\square$ 35.394 $\square$ Oral Nal-131 in quantities greate	er than 1.22 gigabecquerels (33 millicuries)						
Parenteral administration of beta	a-emitter, or photon-emitting radionuclide with a photon ing a written directive is required						
Parenteral administration of any	other radionuclide requiring a written directive						
** Supervising Authorized User must have experience in adm requesting authorized user status.	inistering dosages in the same dosage category or categories as the individual						
d. Provide completed Part II Preceptor Attestation							
PART II – PRE	CEPTOR ATTESTATION						
individual as long as the preceptor provides, dir	lote: 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 a the position sought and not attesting to the indiv	attesting that the individual has knowledge to fulfill the duties of vidual's "general clinical competency."						
First Section Check one of the following for each requested authorization:							
<u>For 35.390:</u>							
Board Certification							
I attest that Name of Proposed Authorized Us	has satisfactorily completed the training and experience						
requirements in 35.390(a)(1).							
OR							
Training and Experience							
I attest that Name of Proposed Authorized Us	has satisfactorily completed the 700 hours of training						
and experience, including a minimum of 200 10 CFR 35.390 (b)(1).	0 hours of classroom and laboratory training, as required by						

NRC FORM 313A (AUT)		U.S. NUCLEAR REGULATORY COMMISSION
AUTHORIZED U	USER TRAINING AND EXPERIEN	CE AND PRECEPTOR ATTESTATION (continued)
Preceptor Attestation (c	continued)	
First Section (contin	iued)	
For 35.392 (Identica	I Attestation Statement Regardle	ss 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 35. quired in 35.392(c)(2).	.392(c)(1), and the supervised work and clinical case
For 35.394 (Identica	I Attestation Statement Regardles	ss 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 35. juired in 35.394(c)(2).	.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 req	uired in 35.390(b)(1)(ii)G listed belo	)W:
	31 requiring a written directive in qua erels (33 millicuries)	antities less than or equal to 1.22
Oral Nal-13	31 in quantities greater than 1.22 gig	gabecquerels (33 millicuries)
	administration of beta-emitter, or ph than 150 keV requiring a written di	oton-emitting radionuclide with a photon rective is required
Parenteral a	administration of any other radionuc	lide requiring a written directive
Third Section		
I attest that		has satisfactorily achieved a level of competency to
	Name of Proposed Authorized User	
function indepe	endently as an authorized user for:	
	31 requiring a written directive in qua erels (33 millicuries)	antities less than or equal to 1.22
Oral Nal-13	31 in quantities greater than 1.22 gig	gabecquerels (33 millicuries)
	administration of beta-emitter, or ph s than 150 keV requiring a written di	oton-emitting radionuclide with a photon rective is required
Parenteral a	administration of any other radionuc	lide requiring a written directive

NRC FORM 313A (AUT) (06-2016) AUTHORIZED USER TRAIN	ING AND EXPERIENCE AND PRECEPT	U.S. NUCLEAR REGULAT						
Fourth Section								
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								
or equivalent Agreement Sta laboratory training, as require experience required by 35.39	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 administration	of any other radionuclide for which a writt	en directive is required						
	OR							
Board Certification:								
I attest that		completed the board cert	ification					
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 administration	of any other radionuclide for which a writt	en directive is required						
	Complete the following for preceptor 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 of	directive						
Name of Preceptor	Signature	Telephone Number	Date					
License/Permit Number/Facility Name								