6/30/2017

Materials Licensing Section
U.S. Nuclear Regulatory Commission
Region III
2443 Warrenville Road STE 210
Lisle, Illinois 60532-4352

Lic. No. 24-02490-03 Docket No. 030-02308

Dear Sir or Madam,

We are respectfully requesting an amendment to our material license number 24-02490-03 at SSM Health DePaul Hospital — St. Louis. We would like to add Radium Ra223 dichloride [Xofigo (an alpha emitting radiotherapeutic drug)] IV administration, to Rebecca J. Mueller, M.D. for authorized user.

She is already on the license for 10CFR35.10; 10CFR35.200; and 35.300 (limited to the oral administration of sodium iodine I-131). She was/has trained, assisted and done procedures with Andre Strzembosz, M.D on 7/26/16, 8/30/16, 9/27/16, 11/1/16 and 12/6/16.

Should you have any questions, please contact Joseph Pekala, Lead CNMT and RSO, for SSM Health DePaul Hospital – St. Louis at 314-447-5981(W) or 314-401-5374(C).

Sincerely,

Joseph Pekala BS, RSO, CNMT, ARRT(CT).

**Radiation Safety Officer** 

Andre Strzembosz, M.D.

NRC FORM 313A (AUT) (06-2016)

U.S. NUCLEAR REGULATORY COMMISSION

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## AUTHORIZED USER TRAINING AND EXPERIENCE AND PRECEPTOR ATTESTATION

(for uses defined under 35.300) [10 CFR 35.390, 35.392, 35.394, and 35.396] APPROVED BY OMB: NO. 3150-0120 EXPIRES: 06/30/2019

Name of Proposed Authorized User		ed Authorized User	State or Territory Where Licensed			
_/	Rebec	ca D. Mueller	Mo			
Red	uested Autl	horization(s) (check all that apply):				
[	35.300 Use of unsealed byproduct material for which a written directive is required					
(	OR					
[	35.300	Oral administration of sodium iodide I-131 requiring a written directive in quantities less than or equal to 1.22 gigabecquerels (33 millicuries)				
	35.300	Oral administration of sodium iodide I-131 requiring a written directive in quantities greater than 1.22 gigabecquerels (33 millicuries)				
[	35.300	35.300 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				
	35.300	Parenteral administration of any other radio	nuclide for which a written directive is required			
	PART I TRAINING AND EXPERIENCE (Select one of the three methods below)					
*	<ul> <li>* 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.</li> <li>1. Board Certification</li> <li>a. Provide a copy of the board certification.</li> </ul>					
		390, provide documentation on supervised cli document this experience.	nical 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 Attestation.					
	2. <u>Current</u>	35.300, 35.400, or 35.600 Authorized Use	r Seeking Additional Authorization			
	a. Authorized User on Materials License 24-02490-03 under the requirements below or equivalent Agreement State requirements (check all that apply):					
	35.3	390 35.392 35.394	35.490 35.690			
	b. If currently authorized for a subset of clinical uses under 35.300, provide documentation on additional required supervised case experience. The table in section 3.c. may be used to document this experience. Also provide completed Part II Preceptor Attestation.					
	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.					

CC FORM 313A (AUT) 2016) AUTHORIZED USER TRAI	NING AND EXPERIENCE AN			ATORY COMMISSION ontinued)
3. Training and Experience fo	r Proposed Authorized User			
a. Classroom and Laboratory Tr	aining 35.390	35.392 3	5.394	35.396
Description of Training	Location of Tr	aining	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				
Radiation biology				
	Total Hours of Training:			
b. Supervised Work Experience 35.390 35.392 35.394 If more than one supervising individual is necessary to document supervised training, provide roof this page.  Supervised Work Experience Total Hours of Experience:			35.396 multiple copies	
Description of Experience Must Include:	Location of Experience Permit Number of	ce/License or of Facility	Confirm	Dates of Experience*
Ordering, receiving, and unpacking radioactive materials safely and performing the related radiation surveys			☐ Yes	
Performing quality control procedures on instruments used to determine the activity of dosages and performing checks for proper operation of survey meters			☐ Yes	
Calculating, measuring, and safely preparing patient or human research subject dosages			☐ Yes	
Using administrative controls to prevent a medical event involving the use of unsealed byproduct material			☐ Yes	
Using procedures to contain spilled byproduct material safely and using proper decontamination procedures			☐ Yes	

## **AUTHORIZED USER TRAINING AND EXPERIENCE AND PRECEPTOR ATTESTATION (continued)**

Training and	d Experience for P	roposed Authoriz <u>ed</u>	<u>User</u> (continued)	
b. Supervise	ed Work Experience	(continued)		
Supervising Individual		License/Permit Number listing supervising individual as an authorized user		
Supervising apply)**:	individual meets the	requirements below,	; or equivalent Agreement State requirements	s (check all that
35.390 With experience administering dosages of:				
35.392	5.392 Oral Nal-131 requiring a written directive in quantities less than or equal to 1.22			
Oral Nal-131 in quantities greater than 1.22 gigabecquerels (33 millicuries)				
35.396 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				
c. Supervised Clinical Case Experience If more than one supervising individual is necessary to document supervised work experience, provide multiple copies of this page.				
Descriptio	on of Experience	Number of Cases Involving Personal Participation	Location of Experience/License or Permit Number of Facility	Dates of Experience*
iodide I-131 i directive in q	tration of sodium requiring a written uantities less than .22 gigabecquerels s)			
iodide I-131 r directive in q	tration of sodium requiring a written uantities greater pabecquerels (33			
any beta-emi photon-emitti with a photor	ing radionuclide n energy less than which a written			
other radionu written directi	dministration of any uclide for which a ive is required	5	SSM Health DePaul Hospital - St. Louis	7/20/16 8/30/16 9/27/16
(Xofi			V	11/1/16

AUTHORIZED USER TRAINING AND EXPERIENCE AND PRECEPTOR ATTESTATION (continued)					
3. Training and Experience for Proposed Authorized	<u>1 User</u> (continued)				
c. Supervised Clinical Case Experience (continued)	c. Supervised Clinical Case Experience (continued)				
Supervising Individual	License/Permit Number listing supervising individual as an authorized user				
Supervising individual meets the requirements below apply)**:	, or equivalent Agreement State requirements (check all that				
35.390 With experience administering dosage	es 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)				
25.396 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 of	ther radionuclide requiring a written directive				
** Supervising Authorized User must have experience in adminis requesting authorized user status.	** 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.					
PART II – PRECE	EPTOR 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 attesting that the individual has knowledge to fulfill the duties of the position sought and not attesting to the individual's "general clinical competency."					
First Section Check one of the following for each requested authorization:					
For 35.390:					
Board Certification					
attest that Rebecca Mueller has satisfactorily completed the training and experience					
requirements in 35.390(a)(1).					
OR					
Training and Experience	has action starily assembled the 700 house of table				
I attest that  Name of Proposed Authorized User	has satisfactorily completed the 700 hours of training				
and experience, including a minimum of 200 hours of classroom and laboratory training, as required by 10 CFR 35.390 (b)(1).					

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,	G AND EXPERIENCE AND PRECEPTOR ATTESTATION (continued)		
Preceptor Attestation (continued)			
First Section (continued)			
For 35.392 (Identical Attestation St	tatement Regardless of Training and Experience Pathway):		
I attest that	has satisfactorily completed the 80 hours of classroom		
and laboratory training, as requested in 35,392(	uired by 10 CFR 35.392(c)(1), and the supervised work and clinical case (c)(2).		
For 35.394 (Identical Attestation St	tatement Regardless of Training and Experience Pathway):		
I attest that	has satisfactorily completed the 80 hours of classroom		
and laboratory training, as requexperience required in 35.394	uired by 10 CFR 35.394 (c)(1), and the supervised work and clinical case (c)(2).		
Second Section			
Mattest that Rebecca L	has satisfactorily completed the required clinical case osed Authorized User		
experience required in 35.390(	b)(1)(ii)G listed below:		
Oral Nal-131 requiring a wr gigabecquerels (33 millicur	ritten directive in quantities less than or equal to 1.22 ies)		
Oral Nal-131 in quantities g	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	f any other radionuclide requiring a written directive		
Third Section			
Tattest that Rebecca	has satisfactorily achieved a level of competency to		
function independently as an a	uthorized user for:		
Oral Nal-131 requiring a wr gigabecquerels (33 millicuri	itten directive in quantities less than or equal to 1.22 ies)		
Oral Nal-131 in quantities g	reater than 1.22 gigabecquerels (33 millicuries)		
	f beta-emitter, or photon-emitting radionuclide with a photon equiring a written directive is required		
Parenteral administration of	f any other radionuclide requiring a written directive		

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NRC FORM 313A (AUT) (06-2016)		U.S. NUCLEAR REGULATORY COMMISSION			
AUTHORIZED USER TRAINII	NG AND EXPERIENCE AND PRECEPT	OR ATTESTATION (continued)			
Fourth Section					
For 35.396:					
Current 35.490 or 35.690 autho	rized user:				
I attest that	is an authorized u	ser under 10 CFR 35.490 or 35.690			
or equivalent Agreement Stat laboratory training, as require experience required by 35.39	Name of Proposed Authorized User 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 writ	ten directive is required			
	OR				
Board Certification:					
I attest that		completed the board certification			
requirements of 35.396(c), ha required by 10 CFR 35.396 (c	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 writ	ten directive is required			
Fifth Section Complete the following for preceptor	attestation and signature:				
I meet the requirements below, o	or equivalent Agreement State requireme	ents, as an authorized user for:			
35.390 35.392	35.394 35.396				
I have experience administering requesting authorization.	I have experience administering dosages in the following categories for which the proposed Authorized User is requesting authorization.				
Oral Nal-131 requiring a writted millicuries)	Oral Nal-131 requiring a written directive in quantities less than or equal to 1.22 gigabecquerels (33 millicuries)				
Oral Nal-131 in quantities gre	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 Andre Strzembosz	Signato	Telephone Number Date 314-344-6482			
License/Permit Number/Facility Name					
24-02490-03 030-0	12308 SSM Health De	Paul Hospital - St. Loms			

SSM Health DePaul Hospital Radiology Department-3<sup>rd</sup> Floor 12303 DePaul Drive Bridgeton, MO 63044





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Materials Licensing Section U.S. Nuclear Regulatory Commission Region III 2443 Warrenville Road STE 210 Lisle, Illinois 60532-4352

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