Virginia Department of Health Radioactive Materials Program (804) 864-8150



TRAINING, EXPERIENCE AND PRECEPTOR ATTESTATION STATEMENT – C (Unsealed Radioactive Material Requiring Written Directive)

The Virginia Department of Health (VDH) is requesting disclosure of all information on this statement for the purpose of authorizing an individual to work with radioactive material. Failure to provide any information may result in denial or delay of authorizing an individual to work with radioactive material.

Instructions: Complete all applicable items. Refer to VAREG "Guidance for Medical Use of Radioactive Material." Use supplementary sheets where necessary. Retain one copy and submit original of the document to the Virginia Department of Health, Radioactive Materials Program, 109 Governor Street, Room 730, Richmond, VA 23219.

Name of Individual State Licensure						
	A copy of license to	practice medicine in Virginia is attached				
Requested Authorization(s) (check all that apply)						
	12VAC5-481-1950 Use of unsealed radioactive material for which a written directive is required					
	31-1950 Oral administration of sodium iodide I-131 requiring a written directive in quantities less than or equal to 1.22 Gigabecquerels (33 millicuries)					
12VAC5-481-1950 Oral administration of sodium (33 millicuries)	12VAC5-481-1950 Oral administration of sodium iodide I-131 requiring a written directive in quantities greater than 1.22 Gigabecquerels					
12VAC5-481-1950 Parenteral administration of any beta emitter, or a photon- emitting radionuclide with a photon energy less than 150 keV, for which a written directive is required 12VAC5-481-1950 Parenteral administration of any other radionuclide, for which a written directive is required						
PART I TRAINING AND EXPERIENCE						
Describe training and experience in sufficient detail to match the training and experience criteria in applicable regulations.						
1. Certification (attach copy of current certificate)						
Specialty Board Category Month and Year Certified						
Provide documentation on supervised clinical case experience. The table in section 4 may be used.						
Note: Items 2-3 do not need to be completed when using Board Certification to meet 12VAC5-481 , Part VII , training and experience requirements.						

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2. Classroom and Laboratory Training.					
Description of Training	Locat	ion	Clock Hours	Dates of Training	
Radiation Physics and Instrumentation	,	-			
Radiation Protection	,	-			
Mathematics Pertaining to Use and Measurement of Radioactivity	,	-			
Chemistry of Radioactive Material for Medical Use	,	-			
Radiation Biology	,	-			
3. Supervised Work Experience					
Description of Experience	Da	tes and Clock Hours o	of Experience		
Ordering, receiving and unpacking radioactive materials 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 preparing patient or human research subject dosages.					
Using administrative controls to prevent a medica					

a. Supervising Individual – Identification and Qualifications

Using procedures to contain spilled radioactive material and using

use of unsealed material.

proper decontamination procedures.

an purpos tioning amount and a committee of the committee	
The training and experience indicated above was obtained under the supervision of (if requirements In 12VAC5-481, Part VII, provide the following information for each)	
☐ 12VAC5-481-1980; ☐ 12VAC5-481-1990; ☐ 12VAC5-481-2000	12VAC5-481-2001
With experience administering dosages of:	
Oral NaI-131 requiring a written directive in quantities less than or equal to 1.	22 Gigabecquerels (33 millicuries)
Oral NaI-131 in quantities greater than 1.22 Gigabecquerels (33 millicuries) Parenteral administration of any beta emitter, or a photon- emitting radionuclic written directive is required	de with a photon energy less than 150 keV, for which a
Parenteral administration of any other radionuclide, for which a written directive	ve is required
Name of Supervising Individual	
Name of License on which Supervising Individual is Authorized	Materials License Number –(Indicate which State

or if NRC)

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4.	Supervised	Clinical	Case	Experience
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Description	Number of Cases Involving Personal Participation	Location	Date 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 a 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 a. Supervising Individual – Identification and Qualifications				
The training and experience indicated above was requirements In 12VAC5-481 , Part VII , provide 12VAC5-481-1980; 12VAC5-481-1	the following information for each			
With experience administering dosages of: Oral NaI-131 requiring a written directive	in quantities less than or equal to	1 22 Gigabecquerels (33 mill	licuries)	
Oral NaI-131 in quantities greater than 1.	•		incuries)	
Parenteral administration of any beta emitter, or a 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				
Name of Supervising Individual				
Name of License on which Supervising Ind	Materials License Num or if NRC)	nber –(Indicate which State		

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TRAINING,	EXP	ERIENCE	AND	PRE	CEPTO	R A	TTEST	ATION	STATI	EMEN'	T – (C
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(Authorized User – Unsealed Written Directive)

PART II – PRECPTOR ATTESTATION						
	Note: This part must be completed by the individual's preceptor. If more than one preceptor is necessary to document experience, obtain a separate preceptor statement from each.					
5. Precepto	or Approval and Attestation					
I attest that the	e individual named in Item 1:					
a. Has sat	isfactorily completed the training requirements in (che	eck all applicable):				
	12VAC5-481-1980 (Use of radioactive materia	ll authorized by 12VAC5-481-1950)				
	12VAC5-481-1990 (Limited to use of sodium i	odide I-131 in quantities ≤ 33 mCi)				
	12VAC5-481-2000 (Limited to use of sodium iodide I-131 in quantities ≥ 33 mCi)					
	12VAC5-481-2001(for the parental administration of unsealed byproduct material requiring a written directive)					
b. Has sa	atisfactorily completed the required clinical case	experience required in 12VAC5-481-1980 (as listed is section 4)				
c. Has re	ceived a level of competency sufficient to functi	on independently as an authorized user for the following:				
	☐ 12VAC5-481-1980 (Use of radioactive material authorized by 12VAC5-481-1950)					
	■ 12VAC5-481-1990 (Limited to use of sodium iodide I-131 in quantities ≤ 33 mCi)					
	12VAC5-481-2000 (Limited to use of sodium iodide I-131 in quantities ≥ 33 mCi)					
12VAC5-481-2001(for the parental administration of unsealed byproduct material requiring a written directive)						
I meet VDH's	requirements to be a preceptor authorized user for:					
	12VAC5-481-1980					
☐ 12VAC5-481-1990						
☐ 12VAC5-481-2000						
	12VAC5-481-2001					
Name of Lic	ense on which Preceptor is Authorized	Materials License Number –(Indicate which State or if NRC)				
Print Name of Preceptor						
SIGNATUR	GNATURE - Preceptor Date Signed					

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