

NRC	FORM 374A		U.S. NUCLEAR	REGULATORY COMMISSION		PAGE 2 OF 6 PAGES
	MATERIALS LICENSE SUPPLEMENTARY SHEET		License Number 13-32637-01MD	Docket or Refer 030-37428	Docket or Reference Number 030-37428	
- -			Amendment No. 7			
6.	Byproduct, source, and/or special nuclear material	7. Chemical and	/or physical form	may possess at any one time	e 9.	Authonized use 2
F.	Any byproduct material authorized under 10 CFR 35.65	F. Sealed Sour	SUCLEAN	F. 75 milliouries total	F.	For use in calibration and checking of the licensee's instruments. For redistribution of sealed sources initially distributed by a manufacturer licensed in accordance with 10 CFR 32.74 to authorized recipients for medical use and for non-medical use to authorized recipients.
G.	Uranium- depleted in Uranium-235	G. Metal		G. 500 kilograms total	G.	For shielding for molybdenum-99/technetium-99m generators.
н.	Any byproduct material identified in 10 CFR 35.400	H. Sealed Sou		H. I currie total	. <b>Н</b> .	For redistribution of sealed sources initially distributed by a manufacturer licensed in accordance with 10 CFR 32.74 that have been registered either with NRC under 10 CFR 32.210 or with an Agreement State and have been distributed in accordance with an NRC or Agreement State specific license authorizing distribution to persons specifically authorized by an NRC or Agreement State license to receive, possess, and use the authorized device.
1.	Any byproduct material permitted by 10 CFR 31.11	I. Prepackage	d Kits	I. 20 millicuries total	. <b>I.</b>	For redistribution to specific licensees or to general licensees in accordance with 10 CFR 31.11, provided the packaging and labeling remain unchanged.

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	CONDITIONS					
10. Licensed material may be used or stored at the licensee's facilities opated at 6538 Corporate Drive, Indianapolis, Indiana.						
11. The Radiation Safety Officer (RSO) for th	is license is Brian K. Hardesty, R.Ph.	A D	•			
12. Licensed material shall only be used by, or under the supervision of:						
A. A pharmacist working of designated	as an authorized nuclear pharmaciscine		)(2)(1) 01 (4).			
Authorized Nuclear Pharmacists	Material and Use	S.				
Brian K. Hardesty, R.Ph. Stephen L. Piepenbrink, R.Ph.		S				
Matthew O. Broshears, R.Ph. Jason J. Wilson, R.Ph.	All - Caller Main	N				
Yogesh P. Patel, Pharm.D.	All Mark A SS	6				
Kevin R. Boyd, R.Ph.						
Joseph M. Stamm, K.Ph.						
	7.40					

13. A. Sealed sources shall be tested for leakage and/or contamination at intervals not to exceed the intervals specified in the certificate of registration issued by the U.S. Nuclear Regulatory Commission under 10 CFR 32.210 or by an Agreement State. In the absence of a registration certificate, sealed sources shall be tested for leakage and/or contamination at intervals not to exceed 6 months, or at such other intervals as specified.

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- B. In the absence of a certificate from a transferor indicating that a leak test has been made within the intervals specified in the certificate of registration issued by the U.S. Nuclear Regulatory Commission under 10 CFR 32.210 or by an Agreement State, prior to the transfer, a sealed source received from another person shall not be part increase until tested and the test results received.
- C. Sealed sources need not be tested if they contain only hydrogen-3; or they contain only a radioactive gas; or the half-life of the isotope is 30 days or less; or they contain not more than 100 microcuries of beta- and/or gamma-emitting material or not more than 10 microcuries of alpha-emitting material.
- D. Sealed sources need not be tested if they are in storage and are not being used. However, when they are removed from storage for use or transferred to another person, and have not been tested within the required leak test interval, they shall be tested before use or transfer. No sealed source shall be stored for a period of more than 10 years without being tested for leakage and/or contamination.
- E. The leak test shall be capable of detecting the presence of 185 becquerels (0.005 microcuries) of radioactive material on the test sample. If the test reveals the presence of 185 becquerels (0.005 microcuries) or more of removable contamination, a report shall be filed with the U.S. Nuclear Regulatory Commission in accordance with 10 CFR 30.50(c)(2), and the source shall be removed immediately from service and decontaminated, repaired, or disposed of in accordance with Commission regulations.
- F. Tests for leakage and/or contamination shall be performed by the licensee or other persons specifically licensed by the U.S. Nuclear Regulatory Commission or an Agreement State to perform such services. In addition, the licensee is authorized to collect leak test samples for analysis by persons specifically licensed by the Commission or an Agreement State to perform such services.
- G. Records of leak test results shall be kept in units of becquerels (microcuries) and shall be maintained for 3 years.
- 14. Sealed sources containing licensed material shall not be opened or sources removed from source holders by the licensee, except as specifically authorized.

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15. The licensee shall conduct a physical inventory every 6 months, or at other intervals approved by the U.S. Nuclear Regulatory Commission,						

- 15. The licensee shall conduct a physical inventory every 6 months, or at other intervals approved by the U.S. Nuclear Regulatory Commission, to account for all sealed sources and/or devices received and possessed under the license. Records of inventories shall be maintained for 3 years from the date of each inventory, and shall include the radionucles, quantities, manufacturer's name and model numbers, and the date of the inventory.
- 16. Except for maintaining labeling as required by 40 CFR Part 20, or Part 71, the licensee shall obtain authorization from the U.S. Nuclear Regulatory Commission before making any changes in the sealed source, device, or source-device combination that would alter the description or specifications as indicated in the respective certificate of registration issued either by the U.S. Nuclear Regulatory Commission pursuant to 10 CFR 32.210 or by an Agreement State.
- 17. The licensee is authorized to hold radioactive material with a physical half-life of less than or equal to 120 days for decay-in-storage before disposal in ordinary trash provided:
  - A. Before disposal as ordinary trash, the waste shall be surveyed at the container surface with the appropriate survey instrument set on its most sensitive scale and with no interposed shielding to determine that its radioactivity cannot be distinguished from background. All radiation labels shall be removed or obliterated, except for radiation labels on materials that are within containers and that will be managed as biomedical waste after they have been released from the licensee.
  - B. A record of each such disposal permitted under this license condition shall be retained for 3 years. The record must include the date of disposal, the date on which the byproduct material was placed in storage, the radionuclides disposed, the survey instrument used, the background dose rate, the dose rate measured at the surface of each waste container, and the name of the individual who performed the disposal.
- 18. The licensee is authorized to retrieve, receive and dispose of radioactive waste from its customers, limited to radiopharmacy-supplied syringes and vials and their contents.
- 19. This license does not authorize distribution to persons exempt from licensing.

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20. Except as specifically provided otherv representations, and procedures cont	vise in this license, the licensee shall conc ained in the documents, including any end	uct its program in accordance wi	th the statements, se condition applies only to

those procedures that are required to be submitted in accordance with the regulations. The U.S. Nuclear Regulatory Commission's regulations shall govern unless the statements, representations, and procedures in the licensee's application and correspondence are more restrictive than the regulations.

A. Application dated December 29, 2016 (MM)004A159)

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B. Letter dated June 26, 2017 (ML17177A205)

FOR THE U.S. NUCLEAR RECULATORY COMMISSION By:

JUN 2 9 2017 Date:

Bryan A. Parker **Region 3** 

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