NAC POPM 313M

(9-81)

10 CFR 35

U.S. NUCLEAR REGULATORY COMMISSION

APPLICATION FOR MATERIALS LICENSE - MEDICAL

Approved by OMB 3150-0041 Expires 9-30-83

INSTRUCTIONS - Complete I terms 1 through 26 if this R an initial application or an application for renewal of a license. Use supplemental sheets where necessary. I term 26 must be completed on all applications and signed. Retain one copy. Submit original and one copy of entire application to: Director, Office of Nuclear Materials Safety and Safeguards, U.S. Nuclear Regulatory Commission, Washington, D.C. 20555. Upon approval of this application, the applicant will receive a Materials License. An NRC Materials License is issued in accordance with the general requirements contained in Title 10, Code of Federal Regulations, Part 30, and the Licensee is subject to Title 10, Code of Federal Regulations, Part 170. The license fee category should be stated in Item 26 and the appropriate fee enclosed.

license fee category should				priate fee enclosed.	out of Federal Fregue	atrona, r o		****
1.a. NAME AND MAILING ADDRESS OF APPLICANT (institution, firm, clinic, physician, etc.) INCLUDE ZIP CODE				1.b. STREET ADDRESS(ES) AT WHICH RADIOACTIVE MATERIAL WILL BE USED (If different from 1.a.) INCLUDE ZIP CODE				
Research Medical Cente	er							
2316 East Meyer Boule								
Kansas City, MO 64132								
TELEPHONE NO.: AREA CODE)							
2. PERSON TO CONTACT REGARDING Morton H. Levitt, M.D Vice President, Clinic	cal S	erv	ices	3. THIS IS AN APPLI	SE IT TO LICENSE N			
TELEPHONE NO.: AREA CODE (816	6) 27	6_	4235	C. LJ RENEWAL C	OF LICENSE NO			
 INDIVIDUAL USERS (Name individual supervise use of radioactive material, Conformation of the formation of the supervise of the su	als who	will Suppi	use or directly lements A and B	5, RADIATION SAFE as radiation safety office me of training and expe	er. If other than indi	vidual us		
6.a. RADIOACTIVE MATERIAL FO	OR ME	DICA	AL USE	1			-	
RADIOACTIVE MATERIAL	DESIF		MAXIMUM POSSESSION LIMITS	ADDITION	AL ITEMS:	MAI ITEN DESIR	VIS	MAXIMUM POSSESSION LIMITS
LISTED IN:		"X"	(In millicuries)	8-4 117 2-1212			"X"	(In millicuries)
10 CFR 31.11 FOR IN VITRO STUDIES		X	5	OF HYPERTHYROID		ENT		
10 CFR 35, 100, SCHEDULE A, GROUP I			AS NEEDED	PHOSPHORUS 32 AS SOLUBLE PHOSPHATE FOR TREATMENT OF POLYCYTHEMIA				
10 CFR 35.100, SCHEDULE A, GROUP I	1		AS NEEDED	PHOSPHORUS-32 AS COLLOIDAL CHROMIC				
10 CFR 35.100, SCHEDULE A, GROUP III				PHOSPHATE FOR INTRACAVITARY TREAT- MENT OF MALIGNANT EFFUSIONS.				
10 CFR 35.100,SCHEDULE A, GROUP IV			AS NEEDED	GOLD-198 AS COLLOID FOR INTRA- CAVITARY TREATMENT OF MALIGNANT EFFUSIONS,				
10 CFR 35.100, SCHEDULE A, GROUP V			AS NEEDED	OF THYROID CARCINOMA				
10 CFR 35,100, SCHEDULE A, GROUP VI		X	1500	XENON-133 AS GAS OR GAS IN SALINE FOR BLOOD FLOW STUDIES AND PULMONARY FUNCTION STUDIES.				
6.b. RADIOACTIVE MATERIAL F calibration and reference standards a							0.1	
	ne autin	-	CHEMICAL	MAXIMUM NUMBER	T T T T T T T T T T T T T T T T T T T	- L131E	U.J	
ELEMENT AND MASS NUMBER		AND/OR PHYSICAL FORM		OF MILLICURIES OF EACH FORM	DESCRIBE PURPOSE OF USE			
Gadolinium-153	T	Gd		2000 8 F	BOTTEMINE	E Po	lensi	tometer
Aprieral Tell	20		ealed	E SE	BANDO	. 60		
Chack Mr. 2003	10	5	ource)	1	WW 5 8 18	35		
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Type of Fee IC Currel			MAN 28 1985			8 1985		
NAC FORM 313M	1/01	4.1				-		

(9-81)

Received By ...

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TROL NO. 78208

INH	FORMA	TION	RECUIRED	EDD	ITEMS 7	THROUGH 2	2
1170	UNIVER	VICTI	DECLUMED	FIJE	I LEIVIS F	I PIRELULIAN A	6.25

For Items 7 through 23, check the appropriate box(e	es) and submit a detailer	d description of all the requested info	ormation. Begin
each item on a separate sheet. Identify the item number	ber and the date of the	application in the lower right corner	of each page. If
you indicate that an appendix to the medical licensing	g guide will be followed	, do not submit the pages, but specif	y the revision
number and date of the referenced guide: Regulatory	y Guide 10.8 Rev	Date:	

MEDICAL ISOTOPES COMMITTEE	15.	GENERAL RULES FOR THE SAFE USE OF RADIOACTIVE MATERIAL (Check One)			
Names and Specialties Attached; and		Appendix G Rules Followed; or			
Duties as in Appendix B; or		Equivalent Rules Attached			
Equivalent Duties Attached		16. EMERGENCY PROCEDURES (Check One)			
8. TRAINING AND EXPERIENCE		Appendix H Procedures Followed; or			
Supplements A & B Attached for Each Individual User; and		Equivalent Procedures Attached			
Supplement A Attached for RSO.	17.	AREA SURVEY PROCEDURES (Check One)			
9. INSTRUMENTATION (Check One)		Appendix I Procedures Followed; or			
Appendix C Form Attached; or		Equivalent Procedures Attached			
List by Name and Model Number	18.	WASTE DISPOSAL (Check One)			
CALIBRATION OF INSTRUMENTS		Appendix J Form Attached; or			
Appendix D Procedures Followed for Survey Instruments; or		Equivalent Information Attached			
Equivalent Procedures Attached; and	19.	THERAPEUTIC USE OF RADIOPHARMACEUTICALS (Check One)			
Appendix D Procedures Followed for Dose Calibrator; or (Check One) Equivalent Procedures Attached		Appendix K Procedures Followed; or			
		Equivalent Procedures Attached			
FACILITIES AND EQUIPMENT	20.	THERAPEUTIC USE OF SEALED SOURCES			
Description and Diagram Attached		Detailed Information Attached; and			
PERSONNEL TRAINING PROGRAM		Appendix L Procedures Followed; or (Check One)			
Description of Training Attached		Equivalent Procedures Attached			
PROCEDURES FOR ORDERING AND RECEIVING RADIOACTIVE MATERIAL	21.	PROCEDURES AND PRECAUTIONS FOR USE OF RADIOACTIVE GASES (e.g., Xenon — 133)			
Detailed Information Attached		Detailed Information Attached			
PROCEDURES FOR SAFELY OPENING PACKAGES 14. CONTAINING RADIOACTIVE MATERIALS		PROCEDURES AND PRECAUTIONS FOR USE OF RADIOACTIVE MATERIAL IN ANIMALS			
(Check One)		Detailed Information Attached			
Appendix F Procedures Followed; or	23.	PROCEDURES AND PRECAUTIONS FOR USE OF RADIOACTIVE MATERIAL SPECIFIED IN ITEM 6.6			
Equivalent Procedures Attached	X	Detailed Information Attached			
	Equivalent Duties Attached TRAINING AND EXPERIENCE Supplements A & B Attached for Each Individual User; and Supplement A Attached for RSO. INSTRUMENTATION (Check One) Appendix C Form Attached; or List by Name and Model Number CALIBRATION OF INSTRUMENTS Appendix D Procedures Followed for Survey Instruments; or (Check One) Equivalent Procedures Attached; and Appendix D Procedures Followed for Dose Calibrator; or (Check One) Equivalent Procedures Attached FACILITIES AND EQUIPMENT Description and Diagram Attached PERSONNEL TRAINING PROGRAM Description of Training Attached PROCEDURES FOR ORDERING AND RECEIVING RADIOACTIVE MATERIAL Detailed Information Attached PROCEDURES FOR SAFELY OPENING PACKAGES CONTAINING RADIOACTIVE MATERIALS (Check One) Appendix F Procedures Followed; or	Names and Specialties Attached; and Duties as in Appendix B; or Equivalent Duties Attached TRAINING AND EXPERIENCE Supplements A & B Attached for Each Individual User; and Supplement A Attached for RSO. INSTRUMENTATION (Check One) Appendix C Form Attached; or List by Name and Model Number CALIBRATION OF INSTRUMENTS Appendix D Procedures Followed for Survey Instruments; or Equivalent Procedures Attached; and Appendix D Procedures Followed for Dose Calibrator; or Equivalent Procedures Attached FACILITIES AND EQUIPMENT Description and Diagram Attached PERSONNEL TRAINING PROGRAM Description of Training Attached PROCEDURES FOR ORDERING AND RECEIVING RADIOACTIVE MATERIAL Detailed Information Attached PROCEDURES FOR SAFELY OPENING PACKAGES CONTAINING RADIOACTIVE MATERIALS (Check One) Appendix F Procedures Followed; or 23.			

		24. PERSONNEL MONITOR	RING DEVICES		
(Check	TYPE appropriate box)	SUPPLIER	EXCHANGE FREQUENCY		
	FILM				
BODY	TLD				
	OTHER (Specify)				
	FILM				
. FINGER TLD					
	OTHER (Specify)				
	FILM				
. WRIST	TLD	Carrier Control			
	OTHER (Specify)				
. OTHER (Spe	ecify)				
HOSPITAL	25. FOR	PRIVATE PRACTICE APPLI	CANTS ONLY		
NAME OF H	OSPITAL	NTS CONTAINING RADIOACTI	b. ATTACH A COPY OF THE AGREEMENT LETTER		
MAILING A	DDRESS		SIGNED BY THE HOSPITAL ADMINISTRATOR.		
			c. WHEN REQUESTING THERAPY PROCEDURES, ATTACH A COPY OF RADIATION SAFETY PRECAU		
CITY STATE ZIP COD		STATE ZIP CODE	TIONS TO BE TAKEN AND LIST AVAILABLE RADIATION DETECTION INSTRUMENTS.		
	(7	26. CERTIFICATE This item must be completed by	y applicant)		
community w	t and any official executing this	certificate on behalf of the applic	ant named in Item 1a certify that this application is prepared in hat all information contained herein, including any supplements		
			b. APPLICANT OR CERTIFYING OFFICIAL (Signature)		
a. LICENSE FEE REQUIRED (See Section 170.31, 10 CFR 170)		Morton H. Levitt, MrQ.			
		Marke of the second			
1) LICENSE	FEE CATEGORY: 7C		Vice President, Clinical Services		
2) LICENSE I	FEE ENCLOSED & 120 00		c. DATE		
121 LICENSE FEE ENCLOSED: \$ 120.00			1-24-85		

RADIOACTIVE MATERIALS NOT LISTED IN ITEM 6A

Gadolinium-153

Manufacturer:

Model:

Gulf Nuclear, Inc. 202 Medical Center Blvd. Webster, Texas (713) 332-3581 Gd-1

See attached lunar data specification sheet for intended use.

The sealed source listed above will be used in a spine/femur bone mineral analyzer.

Source

Device

NRC Device Registration Number:

Gd-153

Lunar DP3

NR-430-D-101-S

We will also be using the lunar forearm scanner and have requested Group VI possession limits.

Source

Device

NRC Device Number

I-125

Lunar SP2

NR-430-D-102-S

Each lunar bone mineral analyzer is installed by a qualified expert who provides two days of installation and training. This training covers source installation, wipe testing, scan operations and data analysis and interpretation. The institution's radiation safety officer will be present for instruction on source replacement and wipe testing.

LUNAR RADIATION CORP.

916 Williamson Street Madison, Wisconsin 53703 (608) 258-8545

At Lunar Radiation our entire business is dedicated to the support of our customers' studies on bone. Lunar offers an unmatched line of systems for all research and clinical applications. Supporting these systems is a highly recognized staff dedicated to the advancement of this field. It is these systems and their support that have made Lunar the leader in bone assessment techniques. Let us help you choose the most effective system for your needs.

DUAL-PHOTON ABSORPTIOMETRY (153 Gd)

DP3 Spine/Femur Scanner. The most sensitive tool for diagnosis and monitoring of bone disease. Features fast-scan for screening or slow-scan for precision measurements. Enables critical evaluation of trabecular bone in order to monitor therapy in serial studies. (Spine measurements have a 2-3% precision in an area that changes 15% annually rather than \pm 1% annually as in the radius.)

DP4 Total Body Scanner. A rectilinear scanner for measuring the total skeleton as well as spine and femoral neck. Offers complete skeletal coverage for clinical research.

SINGLE-PHOTON ABSORPTIOMETRY (1251)

SP2 Forearm Scanner: A rectilinear scanner for high precision (1%) measurements on the limbs. For diagnosis and monitoring of all bone diseases, adult and infant studies, evaluation of the new ADFR treatment modality, and small animal studies. Capable of scanning irregular anatomical areas such as the distal radius and featuring a variety of scan speeds and step intervals.

BENEFITS OF LUNAR SYSTEMS

- -Automatic Bone Location by Intelligent Scanner
- -Automatic Edge and Baseline Detection
- -Data Base of Normals
- -Standards for System Calibration
- -Sophisticated Correction Factors
- -Extensive Application Support
- -24-Hour Replacement Service Program

WASTE DISPOSAL PROCEDURE

- Liquid waste will be disposed of in the sanitary sewer system in accordance with Section 20.303 of 10 CFR Part 20.
- 2. Mo-99/Tc-99m generators will be held for decay until radiation levels as measured with a low level survey meter and with all shielding removed, have reached background levels. All radiation labels will be removed or obliterated and the generators disposed of as normal trash.
- 3. Other solid waste will be held for decay until the radiation level as measured with a low level survey meter and with all shielding removed have reached background levels. All radiation labels will be removed or obliterated and the waste will be disposed of in normal trash. Those isotopes that are extremely long lived, particularly 3-H (tritium), will be disposed of by a commercial waste disposal service.
- 4. The commercial waste disposal service used will be:

Chem-Nuclear Systems, Inc. P.O. Box 1866 Bellevue, WA 98009

NRC/Agreement State License No. 46-19524-01

RADIATION SAFETY PROCEDURES FOR THERAPUTIC USE OF RADIOPHARMACEUTICALS

- 1. All patients treated with Iodine-131 or Gold-198 will be placed in a private room that has a toilet. Three end corridor rooms located on the oncology ward has been designated as the Hospital Radiopharmaceutical Therapy rooms. Each room has had the carpeting removed and has a megative air flow of at least 30 feet per minute into the room. The large surfaces in the rooms and toilet areas that are more likely to be contaminated will be covered with absorbant pads as appropriate to the amounts of contamination to be expected. Special attention will be given to objects likely to be touched by the patient, for example telephones, doorknobs and other items that would be difficult to decontaminate. Plastic bags or wrappings that are waterproof and easily disposable will be used on smaller items.
- 2. The patient's room will be properly posted or attended in accordance with Section 20.203 or 20.204 of 10 CFR part 20.
- 3. Surveys of the patient's room and surrounding areas will be conducted as soon as practical after administration of the treatment dose. Exposure rates will be measured at the patient's bedside and three feet from the patient after administration and at the entrance of the room. The Radiation Safety Officer or his designate will then determine how long a person may remain at these positions and will post these times on the patient's chart and on the door. The results of the daily surveys will be used to recalculate permitted times which will be posted on the patient's chart and on his door.
- 4. A form containing nursip instructions for patient's treated with radiopharmaceuticals will be completed immediately after administration of the treatment dose. A copy will be posted on the patient's chart.
- 5. Radiation levels in unrestricted areas will be maintained at less than the limits specified in paragraph 20.105 (B) of 10 CFR part 20.
- 6. All linens will be surveyed for contamination before being removed from the patient's room and if necessary, will be held for decay.
- 7. Disposable plates, cups, eating utensils, tissue, surgical dressings, and other similar waste items will be placed in a specially designated container. The material will be collected daily by the Radiation Safety Officer or his designee, checked for contamination, and disposed of as normal or radioactive waste, as appropriate.
- 8. Nondisposable items used for these patients will be held in plastic bags in the patient's room and will be checked for contamination by the Radiation Safety Officer or his designee. Items may be returned for normal use, held for decay, or decontaminated as appropriate.
- All excreta from Iodine-131 therapy patients will be released to the sanitary sewer system as permitted by Section 20.303 of 10 CFR part 20.
- 10. Before a therapy patient's room is reassigned to another patient, the room will be surveyed for contamination and decontaminated if necessary, and all radioactivity waste and waste containers will be removed.

- 11. Nursing Instructions: The guidelines outlined in Appendix K, Radiation Safety Procedures for Therapeutic use of Radiopharmaceutical of Regulatory Guide 10.8, A Guide of the Preparation of Applications for Medical Programs will be followed.
- 12. When contaminated wastes are transported to waste storage-disposable area, precautions will be taken to minimize external radiation of personnel. Stored wastes will be shielded to maintain exposure to personnel in restricted and unrestricted areas according to ALARA.

THERAPEUTIC USE OF I-131

- 1. In most cases, liquid iodine will be used for those therapy procedures requiring I-131. Liquid iodine, under most circumstances, will be delivered via a closed system. The system is a Paramedical Oral Radioisotope Administration Set (#32-27). Personnel will be instructed to wear gloves when administrating any liquid I-131 therapy. In those rare instances when an open delivery system must be used personnel will be instructed to open vials or remove liquid I-131 from the vial in a well ventillated area in order to minimize the possible intake of volatile iodine.
- 2. Liquid I-131 therapies requiring patient hospitalization will be administered in one of three designated therapy rooms located on the Oncology ward. Each room is designed as an isolation room and has a negative airflow of at least 30 feet per minute with an open door. The Oncology nursing staff will be instructed in proper radiation safety procedures on a yearly basis and will have their own personnel dosimetry monitor.
- 3. Immediately following the administration of liquid I-131 a whole body scan using a survey meter will be performed on the physician administering the therapy in an attempt to locate contamination. Should contamination be found, immediate decontamination procedures will be implemented to prevent the spread of radioactive materials.
- 4. Bioassays will be performed no earlier than six(6) hours but no later than seventy two(72) hours following administration of liquid I-131. Bioassays will be performed by a qualified Nuclear Medicine Technologist using a thyroid uptake probe and the rsults of the bioassay kept in a log book.

THERAPEUTIC USE OF I-131

- 4.(cont) The results of the bioassay will be interprelated to 24 hours which is assumed to be the time of maximal uptake(see enclosed graph which will be used to generate interprelative values).
- 5. Should a physician approach either the 40 hour MPC of 0.1003 uCi or the 520 hour MPC of 1.300 uCi, that physician will not be allowed to administer further therapies until the beginning of the next monitioring period. A running total of I-131 accumulated in the thyroid of all personnel involved in the administering of liquid I-131 will be maintained.
- 6. Bioassays will be performed and records maintained on any personnel involved in the cleanup of a spill of I-131.

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