

UNITED STATES NUCLEAR REGULATORY COMMISSION REGION I 475 ALLENDALE ROAD KING OF PRUSSIA, PENNSYLVANIA 19406-1415

November 12, 2005

License No.

37-01893-01

 Docket No.
 03003013

 Control No.
 137927

Mary N. Mannix President and Chief Operating Officer Robert Packer Hospital Guthrie Healthcare System One Guthrie Square Sayre, PA 18840-1698

## SUBJECT: GUTHRIE HEALTHCARE SYSTEM AND GUTHRIE CLINIC, LICENSE AMENDMENT, CONTROL NO. 137927

Dear Ms. Mannix:

This refers to your license amendment request dated November 3, 2005. Enclosed with this letter is the amended license adding Charles F. Wild, Ph.D. as an Authorized Medical Physicist.

Please review the enclosed document carefully and be sure that you understand and fully implement all the conditions incorporated into the amended license. If there are any errors or questions, please notify the U.S. Nuclear Regulatory Commission, Region I Office, Licensing Assistance Team, (610) 337-5239, so that we can provide appropriate corrections and answers.

An environmental assessment for this action is not required, since this action is categorically excluded under 10 CFR 51.22(c)(14).

Current NRC regulations and guidance are included on the NRC's website at <u>www.nrc.gov</u>; select Nuclear Materials; Medical, industrial, and academic uses of nuclear material; then toolkit index page. Or you may obtain these documents by contacting the Government Printing Office (GPO) toll-free at 1-888-293-6498. The GPO is open from 7:00 a.m. to 9:00 p.m. EST, Monday through Friday (except Federal holidays).

Thank you for your cooperation.

Sincerely,

Original signed by Sandra Gabriel

Sandra Gabriel Senior Health Physicist Medical Branch Division of Nuclear Materials Safety

Enclosure: Amendmantial Ann711 in s record was deleted in accordance with the Freedom of Information Act, exemptions <u>2</u> FOIA <u>506-135</u>

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M. Mannix Guthrie Healthcare System and Guthrie Clinic 2

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cc: Asaf Durakovic, M.D., Radiation Safety Officer

M. Mannix 3 Guthrie Healthcare System and Guthrie Clinic

## DOCUMENT NAME: E:\Filenet\ML053200232.wpd

SISP Review Complete: <u>SGabriel</u> After declaring this document "An Official Agency Record" It<u>wlll not</u> be released to the Public.

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NAME	SGabriel /SLG2/						
DATE	11/12/05			_			

OFFICIAL RECORD COPY





NRC FOR	RM 374A U.S. NUCLEAR REGULATO	RY COMMISSION	PAGE 3 of 6 PAGES				
			License Number 37-01893-01				
	MATERIALS LICENSE SUPPLEMENTARY SHEET	Docket or Reference Number 030-03013					
ļ			Amendment No. 71				
11. The Radiation Safety Officer for this license is Asaf Durakovic, M.D.							
12. Lic	12. Licensed material is only authorized for use by, or under the supervision of:						
A.	A. Individuals permitted to work as an authorized user, authorized nuclear pharmacist, and/or authorized medical physicist in accordance with 10 CER 35 13 and 35 14						
B	The following individuals are authorized	D P P	dical use as indicated:				
0.							
·	Authorized Users	<u>Material ar</u>	nd Use				
	John M. Antos, M.D.	35.100; 35	5.200; 35.300; <u>In vitro</u> studies				
	Richard Foster, M.D.	35.100; 35	.200 <u>r In vitro</u> studies				
	Ralph D. Zehr, M.D.	35-100; 35	200: 35.300; 35.400; <u>In vitro</u> studies				
	Christopher Joy, M.D.	35,100; 35 	200 Oral administration of sodium iodide				
		: streatment	of hyperthyroidism and cardiac dysfunction				
	Thomas Gergel, M.D.	i iridium 192 A afterloadin	2 for use in a high dose rate remote g device; depleted uranium				
	Gary Proulx, M.D.	35,400; Irid afterloadin	dium 192 for use in a high dose rate remote g device depleted uranium				
	Asaf Durakovic, M.D.	35,100; 35	200; 35.300; <u>In vitro</u> studies				
	Duk K. Choi, M.D.	35.100; 35	5.200				
	Richard A. Kostick, D.O.	35.100; 35	.200; <u>In vitro</u> studies				
C.	The following individuals are authorized	d medical phys	sicists as indicated:				
	Authorized Medical Physicists	Material ar	nd Use				
	William F. Kendall, Ph.D.	Iridium-192 for calibrat	2 in a High Dose Rate Remote Afterloader Unit ions, spot-checks, and training				
	Charles F. Wild, Ph.D.	Iridium-192 for calibrat	2 in a High Dose Rate Remote Afterloader Unit ions, spot-checks, and training				

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		<u> </u>			
	D.	The following individual:	s are authorized ι	isers for nor	n-medical uses as indicated:
	ł	Authorized Users		<u>Materials</u>	
	ļ	Nan-Shang Chang, Ph.[	).	Hydrogen : Sulfur 35, 1	3, Carbon 14, Phosphorus 32, Phosphorus 33, Calcium 45, Chromium 51, Iodine 125
		John D. Noti, Ph.D.	GLEAF	Hydrogen Sulfur 35,	ຊື່ JCarbon 14, Phosphorus 32, Phosphorus 33, Calcium 45, Chromium 51, Iodine 125
	I	Carol L. Williams, Ph.D.		Hydrogen : Sulfur 35,	3, Carbon 14, Phosphorus 32, Phosphorus 33, Calcium 45, Chromium 51, Iodine 125
		Sydney Welt, M.D.		Chromium	51, lodine, 125, <u>Yt</u> trium 90
		Asaf Durakovic, MD		Americium	241 (storage), Strontium 90 (calibration)
13.	In ac mate esta	dition to the possession erial to quantities below blishing financial assura	1 limits in item 8, t the minimum limit ance for decommis	he licensee specified in ssioning t	shall further restrict the possession of licensed 10 GFR 30.35(d) 40.36(b), and 70.25(d) for
14.	The spea	licensee shall not use lice	censed material in nse.	i or on huma	an beings except as provided otherwise by
15.	The prov	licensee shall not use lic ided otherwise by speci	censed material in fic condition of thi	i field applic s license.	ations where activity is released except as
16.	The deca	licensee is authorized to ay-in-storage before disp	o hold byproduct rega	naterial with Ird to its radi	a physical half-life of less than 120 days for ioactivity if it:
	A.	Monitors byproduct mate cannot be distinguished survey meter set on its	erial at the surfact from the backgrc most sensitive sca	e before dis ound radiatio ale and with	posal and determines that its radioactivity on level with an appropriate radiation detection no interposed shielding; and
	В.	Removes or obliterates containers and that will licensee; and	all radiation labels be managed as b	s, except for iomedical w	r radiation labels on materials that are within aste after they have been released from the
	C.	Maintains records of the of disposal, the survey i at the surface of each v	<ul> <li>disposal of licent instrument used, t</li> <li>waste container, a</li> </ul>	sed material the backgrou nd the name	Is for 3 years. The record must include the date und radiation level, the radiation level measured e of the individual who performed the disposal.
17.	The 10 (	licensee is authorized to CFR Part 71, "Packaging	o transport license J and Transportati	ed material ir on of Radioa	n accordance with the provisions of active Material."

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U.S. NUCLEAR REGULATORY COMMISSION

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SUPPLEMENTARY SHEET

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18	For	sealed sources pot associated with 10 CER Part 35 up	the following conditions on hur
10.	101	sealed sources not associated with 10 Chill Part 35 da	se, the following conditions apply.
	Α.	Sealed sources shall be tested for leakage and/or con intervals specified in the certificate of registration issu under 10 CFR 32.210 or under equivalent regulations	ntamination at intervals not to exceed the red by the U.S. Nuclear Regulatory Commission s of an Agreement State.
	В.	Notwithstanding Paragraph A of this Condition seale particles shall be tested for leakage and/or contamina	d sources designed to primarily emit alpha ation at intervals not to exceed 3 months.
	C.	Each sealed source fabricated by the licensee shall b leakage, and contamination prior to any use or transf	e inspected and tested for construction defects, er as a sealed source.
	D.	In the absence of a certificate from a transferor indicate intervals specified in the certificate of registration issuunder 10 CFR 32.210 or under equivalent regulations sealed source received from another person shall not received.	ating that a leak test has been made within the ned by the U.S. Nuclear Regulatory Commission of an Agreement State, prior to the transfer, a be put into use until tested and the test results
	Ε.	Sealed sources need not be jested if they contain on gas; or the half-life of the isotope is 30 days of less of beta- and/or gamma-emitting material of not more that	y hydrogen-3; or they contain only a radioactive or they contain not more than 100 microcuries of an 10 microcuries of alpha-emitting material.
	F.	Sealed sources need not be tested if they are in stora are removed from storage for use of transferred to ar the required leak test interval, they shall be tested be stored for a period of more than 10 years without bein	ge and are not being used; however, when they other person and have not been tested within fore use of transfer. No sealed source shall be ng tested for leakage and/or contamination.
	G.	The leak test shall be capable of detecting the preser radioactive material on the test sample. If the test rev (185 becquerels) or more of removable contamination Regulatory Commission in accordance with 10 CFR 3 immediately from service and decontaminated, repair Commission regulations.	nce of 0.005 microcurie (185 becquerels) of veals the presence of 0.005 microcurie n, a report shall be filed with the U.S. Nuclear 00.50(c)(2), and the source shall be removed ed, or disposed of in accordance with
	H.	Tests for leakage and/or contamination, including leaperformed by the licensee or by other persons specific Commission or an Agreement State to perform such skept in units of microcuries and shall be maintained for	k test sample collection and analysis, shall be cally licensed by the U.S. Nuclear Regulatory services. Records of leak test results shall be or 5 years.
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License Number 27 04000 04

19. The licensee shall conduct a physical inventory every six months, or at other intervals approved by the U.S. Nuclear Regulatory Commission, to account for all sources and/or devices received and possessed under the license. Records of inventories shall be maintained for 5 years from the date of each inventory and shall include the radionuclides, quantities, manufacturer's name and model numbers, and the date of the inventory.

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		Amendment No. 71					
<ul> <li>20. Except as spaccordance of any enclosure be submitted licensee's at The U.S. Nurepresentation than the regression of the transformed of transformed</li></ul>	becifically provided otherwise in this license, the with the statements, representations, and pro- res, listed below. This license condition applied in accordance with the regulations. Addition bility to make changes to the radiation protect clear Regulatory Commission's regulations s ons, and procedures in the licensee's applicate ulations. ated December 18, 1990 ated August 17, 1995 ated November 30, 1995 ated Movember 30, 1995 ated March 16, 2001 except Quality Manage ated August 28, 2001 except QMP dated August 30, 2001 ated September 12, 2001 ated September 14, 2001 ated September 14, 2002 ated March 27, 2002 ated December 15, 2003 ated October 18, 2004	the licensee shall conduct its program in breedures contained in the documents, including les only to those procedures that are required to hally, this license condition does not limit the tion program as provided for in 10 CFR 35.26. hall govern unless the statements, tion and correspondence are more restrictive ment Program (QMP) [ML010880213] [ML012410090] [ML012420439] [ML012640354] [ML012640354] [ML020530307] [ML020950018] [ML043000249]					
	For the LL	S. Nuclear Regulatory Commission					
For the U.S. Nuclear Regulatory Commission							
Date Nov	ember 12, 2005 By	iginal signed by Sandra Gabriel					
	ndra Gabriel dical Branch						
	Div Re Kir	rision of Nuclear Materials Safety gion I Ig of Prussia, Pennsylvania 19406					

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