

PUTNAM GENERAL
HOSPITAL
We're doing great things.

November 7, 2006

J-3

VIA FACSIMILE (610)337-5393 and United States Mail
 United States Nuclear Regulatory Commission
 Division of Nuclear Materials Safety
 475 Allendale Road
 Region 1
 King of Prussia, PA 19406-1415
 Attn: LAT, DNMS

Re: Putnam General Hospital
 License Number 47-23070-01 03020237
Request to Expedite License Amendment

Ladies and Gentlemen:

This letter will serve as notification of the change of ownership of Putnam General Hospital, 1400 Hospital Drive, Hurricane, West Virginia, 25526 (the "Hospital"). Effective November 11, 2006, at 12:01 a.m. (the "Effective Time"), the assets of the Hospital will be sold by Teays Valley Health Services, LLC (the "Seller") to CAMC Teays Valley Hospital, Inc. ("CAMC Teays Valley Hospital"). At the Effective Time, CAMC Teays Valley Hospital will commence operation of the Hospital under the name "CAMC Teays Valley Hospital."

From and after the Effective Time, the Radiation Safety Officer for CAMC Teays Valley Hospital will be Steven A. Artz, M.D. He currently serves as Radiation Safety Officer under License Number 47-15473-01 (Charleston Area Medical Center, Inc.). **There will be no changes in services or radioactive materials at the Hospital.**

From and after the Effective Time, the President and Vice President of CAMC Teays Valley Hospital will be:

Albert H. Michaels
 President

1400 Hospital Drive
 Hurricane, WV 25526
 304-757-1792

Randall H. Hodges
 Vice President

1400 Hospital Drive
 Hurricane, WV 25526
 304-757-1796

From and after the Effective Time, the following individuals will be authorized users for the material and use specified:

<u>Authorized User</u>	<u>Material and Use</u>	<u>Currently Licensed Under</u>
Steven A. Artz, M.D.	35.100; 35.200; 35.300; 35.400; 35.500; In vitro studies; H-3; C-14; Gd-153 Sealed Sources, P-32 Guidant Sealed Source(s), and the Cs-137 J.L. Shepherd Sealed Source for storage only; Depleted Uranium	47-15473-01
John J. Anton, M.D.	35.100; 35.200	47-15473-01
Michael Eugene Anton, M.D.	35.100; 35.200	47-15473-01
Timothy Connor, M.D.	35.100; 35.200; 35.300; 35.400	47-15473-01
Ronald Cordell, M.D.	35.100; 35.200	47-15473-01
Jeffrey C. Dameron, M.D.	35.100; 35.200; 35.300	47-15473-01
Stephen M. Elksnis, M.D.	35.100; 35.200	47-15473-01
Mohammed Haffar, M.D.	35.100; 35.200	47-23070-01
Jennifer Marie Smith, M.D.	35.100; 35.200; 35.300	47-15473-01
Russell F. King, III, M.D.	35.100; 35.200; 35.300	47-15473-01
Johnsey Leef, M.D.	35.100; 35.200; 35.300; 35.400	47-15473-01
Johnsey Lee Leef, III, M.D.	35.100; 35.200; 35.300	47-15473-01
Mary H. McJunkin, M.D.	35.100; 35.200	47-15473-01

<u>Authorized User</u>	<u>Material and Use</u>	<u>Currently Licensed Under</u>
John F. Mega, M.D.	35.100; 35.200	47-15473-01
Bassam Moushmoush, M.D.,	35.100; 35.200	47-23070-01
Frank Muto, M.D.	35.100; 35.200	47-15473-01
John Reifsteck, M.D.	35.100; 35.200	47-15473-01
Christopher A. Schlarb, M.D.	35.100; 35.200	47-15473-01
James T. Smith, M.D.	35.100; 35.200	47-15473-01
John Willis, M.D.	35.100; 35.200	47-15473-01

Please advise us if you require any further information in order to update your records with respect to CAMC Teays Valley Hospital. Thank you for your cooperation and assistance.

Very truly yours,

Teays Valley Health Services, LLC
d/b/a Putnam General Hospital

By: Ronda J. Moore

Title: COO/CEO

CAMC Teays Valley Hospital, Inc.

By: Michael J. T... ..

Title: President

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Amendment No. 51

MATERIALS LICENSE

Pursuant to the Atomic Energy Act of 1954, as amended, the Energy Reorganization Act of 1974 (Public Law 93-438), and Title 10, Code of Federal Regulations, Chapter I, Parts 30, 31, 32, 33, 34, 35, 36, 39, 40, and 70, and in reliance on statements and representations heretofore made by the licensee, a license is hereby issued authorizing the licensee to receive, acquire, possess, and transfer byproduct, source, and special nuclear material designated below; to use such material for the purpose(s) and at the place(s) designated below; to deliver or transfer such material to persons authorized to receive it in accordance with the regulations of the applicable Part(s). This license shall be deemed to contain the conditions specified in Section 183 of the Atomic Energy Act of 1954, as amended, and is subject to all applicable rules, regulations, and orders of the Nuclear Regulatory Commission now or hereafter in effect and to any conditions specified below.

<p style="text-align: center;">Licensee</p> <p>1. Charleston Area Medical Center</p> <p>2. P.O. Box 1547 Charleston, West Virginia 25326</p>	<p>In accordance with the letter dated May 30, 2006,</p> <p>3. License number 47-15473-01 is amended in its entirety to read as follows:</p> <hr/> <p>4. Expiration date July 31, 2006 (extended)</p> <hr/> <p>5. Docket No. 03009164 Reference No.</p>	
<p>6. Byproduct, source, and/or special nuclear material</p> <p>A. Any byproduct material permitted by 10 CFR 35.100</p> <p>B. Any byproduct material permitted by 10 CFR 35.200</p> <p>C. Any byproduct material permitted by 10 CFR 35.300</p> <p>D. Any byproduct material permitted by 10 CFR 35.400</p> <p>E. Any byproduct material permitted by 10 CFR 35.500</p>	<p>7. Chemical and/or physical form</p> <p>A. Any</p> <p>B. Any</p> <p>C. Any</p> <p>D. Sealed Sources (3M Models 6501, 6502, and 6503; Best Medical International Inc. Model 81-01, 2301, and 81-02)</p> <p>E. Sealed Sources (Bristol-Myers Squibb Medical Imaging Model NES-8412; North American Scientific, Inc. Model MED3601)</p>	<p>8. Maximum amount that licensee may possess at any one time under this license</p> <p>A. As needed</p> <p>B. As needed</p> <p>C. 1 curie</p> <p>D. 3 curies</p> <p>E. 0.3 curies per source and 3.6 curies total</p>

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- | | | |
|---|---|---|
| 6. Byproduct, source, and/or special nuclear material | 7. Chemical and/or physical form | 8. Maximum amount that licensee may possess at any one time under this license |
| F. Any byproduct material permitted by 10 CFR 31.11 | F. Prepackaged Kits | F. 5 millicuries |
| G. Cesium 137 | G. Sealed Sources (Isomedia Model ISO-1000; MDS Nordion Model C-1000 or C-1001) | G. No single source to exceed the maximum activity specified in the certificate of registration issued by the U. S. Nuclear Regulatory Commission or an Agreement State |
| H. Gadolinium 153 | H. Sealed Sources (Lunar Corporation Model GD-1; Amersham Model GDC-CY1) | H. 4 millicurie |
| I. Cesium 137 | I. Sealed Source | I. 51 millicuries |
| J. Phosphorus 32 | J. Sealed Sources (Guidant Model GD7-R-32 Series) | J. 600 millicuries |
| K. Hydrogen 3 | K. Any | K. 5 millicuries |
| L. Carbon 14 | L. Any | L. 5 millicuries |
| M. Depleted Uranium | M. Metal | M. 360 kilograms |

9. Authorized use:

- A. Any uptake, dilution and excretion study permitted by 10 CFR 35.100.
- B. Any imaging and localization study permitted by 10 CFR 35.200.
- C. Any diagnostic study or therapy procedure permitted by 10 CFR 35.300.
- D. Any manual brachytherapy procedure permitted by 10 CFR 35.400.
- E. Diagnostic medical use of sealed sources permitted by 10 CFR 35.500 in compatible devices registered pursuant to 10 CFR 30.32(g).
- F. In vitro studies.

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- G. For irradiation of materials in self-shielded irradiator devices in accordance with the certificate of registration issued by the U.S. Nuclear Regulatory Commission under 10 CFR 32.210 or with an Agreement State and which have been distributed in accordance with a Commission or Agreement State specific license authorizing distribution to persons specifically authorized by a Commission or Agreement State license to receive, possess, and use the devices.
- H. For storage only incident to disposal.
- I. For storage only incident to disposal in a J.L. Shepherd 78 series calibrator.
- J. For storage only incident to disposal.
- K. through L. Possession incident to decontamination or disposal.
- M. Shielding for molybdenum-99/technetium-99m generators.

CONDITIONS

- 10. A. Licensed material may be used or stored at the licensee's facilities located at Memorial Division, 3200 MacCorkle Avenue, Charleston, West Virginia.
- B. Licensed material in items 6.A - 6.E may be used or stored at the licensee's facilities located at :
 - a. General Division, 501 Morris Street, Charleston, West Virginia.
 - b. Women & Children's Division, 800 Pennsylvania Avenue, Charleston, West Virginia.
 - c. Braxton County Memorial Hospital, 100 Hoylman Drive, Gasaway, West Virginia.
- 11. The Radiation Safety Officer for this license is Steven Artz, M.D.
- 12. Licensed material is only authorized for use by, or under the supervision of:
 - A. Individuals permitted to work as an authorized user, and/or authorized nuclear pharmacist in accordance with 10 CFR 35.13 and 35.14.
 - B. The following individuals are authorized users for medical use as indicated:

<u>Authorized Users</u>	<u>Material and Use</u>
John J. Anton, M.D.	35.100; 35.200
Michael Eugene Anton, M.D.	35.100; 35.200

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Authorized Users

Material and Use

Steven A. Artz, M.D.

35.100; 35.200; 35.300; 35.400; 35.500; In vitro studies; H-3, C-14, Gd-153 Sealed Sources, P-32 Guidant Sealed Source(s), and the Cs-137 J.L. Shepherd Sealed Source for storage only; Depleted Uranium

Dilip K. Basu, M.D.

35.100; 35.200

Nicholas Cassis, M.D.

Oral administration of sodium iodide iodine-131

Timothy Connor, M.D.

35.100; 35.200; 35.300; 35.400

Ronald Cordell, M.D.

35.100; 35.200

Glenn Crotty, M.D.

Oral administration of sodium iodide iodine-131

Jeffrey Dameron, M.D.

35.100; 35.200; 35.300

Joseph Devono, III, M.D.

35.100; 35.200

Stephen M. Elkanis, M.D.

35.100; 35.200

Mary Elizabeth Faw, M.D.

35.300

Jean-Pierre M. Geagea, M.D.

35.100; 35.200

John Goad, M.D.

35.100; 35.200

Steven Grubb, M.D.

Oral administration of sodium iodide iodine-131

Mohammed Haffar, M.D.

35.100; 35.200

Omar Hallak, M.D.

35.100; 35.200

Michael Blake Harmon, M.D.

35.400

Ramakrishnan Iyer, M.D.

35.100; 35.200

Kshama Jawalekar, M.D.

35.400

Jennifer Marie Smith, M.D.

35.100; 35.200; 35.300

Russell F. King, III, M.D.

35.100; 35.200; 35.300

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<u>Authorized Users</u>	<u>Material and Use</u>
Marciano B. Lee, M.D.	35.100; 35.200
Johnsey Leef, M.D.	35.100; 35.200; 35.300; 35.400
Johnsey Lee Leef, III, M.D.	35.100; 35.200; 35.300
Donald Lilly, M.D.	35.100; 35.200
Steven McCormick, M.D.	35.100; 35.200
Mary McJunkin, M.D.	35.100; 35.200
John F. Mega, M.D.	35.100; 35.200
Muhammand S. Mian, M.D.	35.100; 35.200
Scott Miller, M.D.	35.100; 35.200
Bassam Moushmouth, M.D.	35.100; 35.200
Frank Muto, M.D.	35.100; 35.200
Brian Allen Plants, M.D.	35.400
Premkumar Raja, M.D.	35.400
John Reifsteck, M.D.	35.100; 35.200
Gary Roberts, D.O.	35.100; 35.200
Ahmed Sakkal, M.D.	35.100; 35.200
Christopher Schlarb, M.D.	35.100; 35.200
James Smith, M.D.	35.100; 35.200
James Stanton, M.D.	35.100; 35.200
Jashvantial Thakkar, M.D.	35.100; 35.200
Lewis A. Whaley, D.O.	35.400
John Willis, M.D.	35.100; 35.200

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Authorized Users

M. Babar Yousef, M.D.

Material and Use

35.100; 35.200

- C. Authorized nuclear pharmacist: Kim David Lowe, Pharm.D.
- D. Licensed material in Item 6.G. shall be used by, or under the supervision of, individuals who have received the training described in letter dated January 12, 2006, and have been designated, in writing, by the Radiation Safety Officer. The licensee shall maintain records of individuals designated as users for 3 years following the last use of licensed material by the individual.
13. In addition to the possession limits in Item 8, the licensee shall further restrict the possession of licensed material to quantities below the minimum limit specified in 10 CFR 30.35(d) for establishing decommissioning financial assurance.
14. Sealed sources or detector cells containing licensed material shall not be opened or sources removed from source holders by the licensee.
15. 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.
16. For sealed sources not associated with 10 CFR Part 35 use, the following conditions apply:
- A. Sealed sources shall be tested for leakage and/or contamination at intervals not to exceed six months or at the intervals specified in the certificate of registration issued by the U.S. Nuclear Regulatory Commission under 10 CFR 32.210 or under equivalent regulations of an Agreement State.
- B. Notwithstanding Paragraph A of this Condition, sealed sources designed to primarily emit alpha particles shall be tested for leakage and/or contamination at intervals not to exceed 3 months.
- C. 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 under equivalent regulations of an Agreement State, prior to the transfer, a sealed source received from another person shall not be put into use until tested and the test results received.

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- D. 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.
 - E. 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.
 - F. The leak test shall be capable of detecting the presence of 0.005 microcurie (185 becquerels) of radioactive material on the test sample. If the test reveals the presence of 0.005 microcurie (185 becquerels) 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.
 - G. Tests for leakage and/or contamination, including leak test sample collection and analysis, shall be performed by the licensee or by other persons specifically licensed by the U.S. Nuclear Regulatory Commission or an Agreement State to perform such services.
 - H. Records of leak test results shall be kept in units of microcuries and shall be maintained for 5 years.
17. The licensee shall not repair, remove, replace, or alter any of the following: electrical and mechanical systems that control source or shielding movement, the irradiator's shielding or sealed source, safety interlocks, or any component that may affect safe operation of the irradiator. These activities shall be performed by a person specifically licensed by the U.S. Nuclear Regulatory Commission or an Agreement State to perform such services.
18. The licensee is authorized to transport licensed material in accordance with the provisions of 10 CFR Part 71, "Packaging and Transportation of Radioactive Material."

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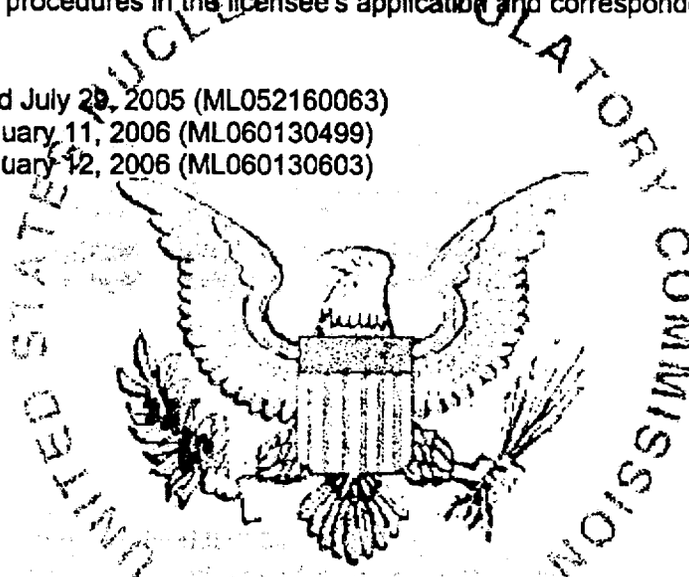
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19. Except as specifically provided otherwise in this license, the licensee shall conduct its program in accordance with the statements, representations, and procedures contained in the documents, including any enclosures, listed below. This license condition applies only to those procedures that are required to be submitted in accordance with the regulations. Additionally, this license condition does not limit the licensee's ability to make changes to the radiation protection program as provided for in 10 CFR 35.26. 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 July 29, 2005 (ML052160063)
- B. Letter dated January 11, 2006 (ML060130499)
- C. Letter dated January 12, 2006 (ML060130603)



For the U.S. Nuclear Regulatory Commission

Date July 13, 2006

By *Penny Lanzisera*

Penny Lanzisera
Medical Branch
Division of Nuclear Materials Safety
Region I
King of Prussia, Pennsylvania 19406

Thursday, July 13, 2006 12:33:13 PM

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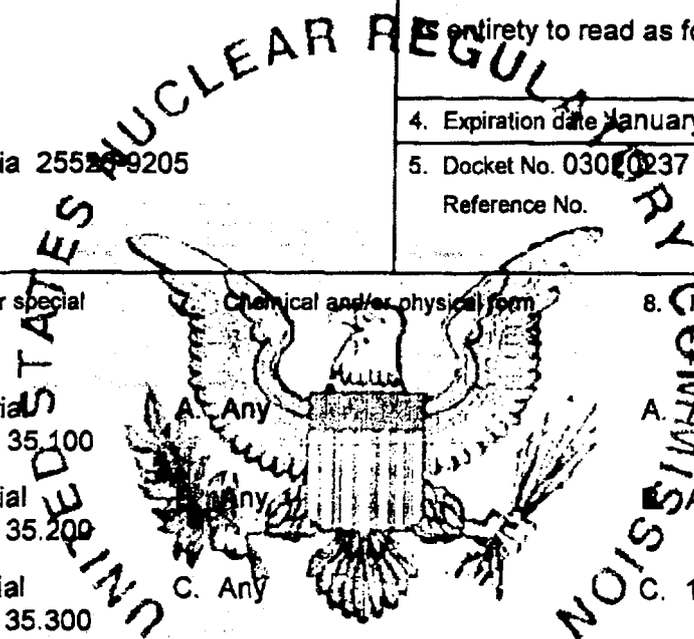
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Amendment No. 14

MATERIALS LICENSE

Pursuant to the Atomic Energy Act of 1954, as amended, the Energy Reorganization Act of 1974 (Public Law 93-438), and Title 10, Code of Federal Regulations, Chapter I, Parts 30, 31, 32, 33, 34, 35, 36, 39, 40, and 70, and in reliance on statements and representations heretofore made by the licensee, a license is hereby issued authorizing the licensee to receive, acquire, possess, and transfer byproduct, source, and special nuclear material designated below; to use such material for the purpose(s) and at the place(s) designated below; to deliver or transfer such material to persons authorized to receive it in accordance with the regulations of the applicable Part(s). This license shall be deemed to contain the conditions specified in Section 183 of the Atomic Energy Act of 1954, as amended, and is subject to all applicable rules, regulations, and orders of the Nuclear Regulatory Commission now or hereafter in effect and to any conditions specified below.

<p>Licensee</p> <p>1. Putnam General Hospital</p> <p>2. 1400 Hospital Drive Hurricane, West Virginia 25521-9205</p>	<p>In accordance with the application dated July 29, 2004,</p> <p>3. License number 47-23070-01 is amended in its entirety to read as follows:</p> <p>4. Expiration date January 31, 2015</p> <p>5. Docket No. 03010237 Reference No.</p>
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<p>6. Byproduct, source, and/or special nuclear material</p> <p>A. Any byproduct material permitted by 10 CFR 35.100</p> <p>B. Any byproduct material permitted by 10 CFR 35.200</p> <p>C. Any byproduct material permitted by 10 CFR 35.300</p>	<p>Chemical and/or physical form</p> <p>A. Any</p> <p>B. Any</p> <p>C. Any</p>	<p>8. Maximum amount that licensee may possess at any one time under this license</p> <p>A. As needed</p> <p>B. As needed</p> <p>C. 1000 millicuries</p>
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9. Authorized use:

A. Any uptake, dilution and excretion study permitted by 10 CFR 35.100.

B. Any imaging and localization study permitted by 10 CFR 35.200.

C. Any diagnostic study or therapy procedure permitted by 10 CFR 35.300.

CONDITIONS

- 10. Licensed material may be used or stored only at the licensee's facilities located at 1400 Hospital Drive, Hurricane, West Virginia.
- 11. The Radiation Safety Officer for this license is James Alan Cochrane, M.D.

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12. Licensed material is only authorized for use by, or under the supervision of:

- A. Individuals permitted to work as an authorized user, authorized nuclear pharmacist, and/or authorized medical physicist in accordance with 10 CFR 35.13 and 35.14.
- B. The following individuals are authorized users for medical use as indicated:

Authorized Users

Material and Use

James Alan Cochran, M.D.

35.100; 35.200; oral administration of sodium iodide I-131 for imaging and localization studies and for treatment of hyperthyroidism and cardiac dysfunction

Paul Dexter Akers, M.D.

35.100; 35.200; oral administration of sodium iodide I-131 for imaging and localization studies

Paul V. Akers, M.D.

35.100; 35.200; 35.300

Marsha S. Anderson, M.D.

35.100; 35.200; oral administration of sodium iodide I-131 for imaging and localization studies

Rodger A. Blake, M.D.

35.100; 35.200; 35.300

Paul H. Blom, M.D.

35.100; 35.200; 35.300

Peter A. Chirico, M.D.

35.100; 35.200; 35.300

Ricky J. Compton, M.D.

35.100; 35.200; oral administration of sodium iodide I-131 for imaging and localization studies

Hans G. Dransfeld, M.D.

35.100; 35.200; 35.300

Joseph W. Dransfeld, M.D.

35.100; 35.200; oral administration of sodium iodide I-131 for imaging and localization studies

Kellie K. Gooding, M.D.

35.100; 35.200; 35.300

Mohammed Y. Haffar, M.D.

35.200

Lee C. Haikal, M.D.

35.100; 35.200; oral administration of sodium iodide I-131 for imaging and localization studies

Michael V. Korona, M.D.

35.100; 35.200; oral administration of sodium iodide I-131 for imaging and localization studies

Donald R. Lewis, Jr., M.D.

35.100; 35.200; 35.300

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Authorized Users

Material and Use

Eric L. Leonard, M.D.

35.100; 35.200; oral administration of sodium iodide I-131 for imaging and localization studies

Richard E. McWhorter, M.D.

35.100; 35.200; 35.300

Bassam Moushmouth, M.D.

35.200

Maria Luna T. Navarro, M.D.

35.200

William S. Shiels, M.D.

35.100; 35.200; 35.300

Charles M. Siegler, M.D.

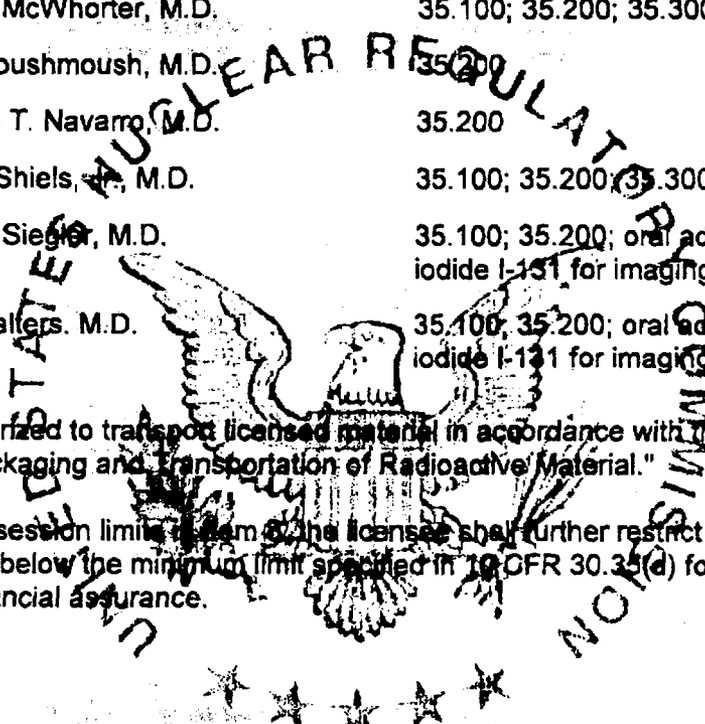
35.100; 35.200; oral administration of sodium iodide I-131 for imaging and localization studies

Torin P. Walters, M.D.

35.100; 35.200; oral administration of sodium iodide I-131 for imaging and localization studies

13. The licensee is authorized to transport licensed material in accordance with the provisions of 10 CFR Part 71, "Packaging and Transportation of Radioactive Material."

14. In addition to the possession limits of the licensee shall further restrict the possession of licensed material to quantities below the minimum limit specified in 10 CFR 30.35(d) for establishing decommissioning financial assurance.



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**MATERIALS LICENSE
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License Number
47-23070-01

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Amendment No. 14

15. Except as specifically provided otherwise in this license, the licensee shall conduct its program in accordance with the statements, representations, and procedures contained in the documents, including any enclosures, listed below. This license condition applies only to those procedures that are required to be submitted in accordance with the regulations. Additionally, this license condition does not limit the licensee's ability to make changes to the radiation protection program as provided for in 10 CFR 35.26. 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 January 12, 2005



For the U.S. Nuclear Regulatory Commission

Date January 19, 2005

By *Michelle Beardsley*
Michelle Beardsley

Medical Branch
Division of Nuclear Materials Safety
Region I
King of Prussia, Pennsylvania 19406

96458087