



UNITED STATES  
NUCLEAR REGULATORY COMMISSION

REGION IV  
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ARLINGTON, TEXAS 76011-4125

October 27, 2008

Carol A. Couch, Ph.D., Director  
Georgia Department of Natural Resources  
Environmental Protection Division  
2 Martin Luther King, Jr. Drive, SE  
Suite 1152, East Tower  
Atlanta, Georgia 30334

Dear Dr. Couch:

The U.S. Nuclear Regulatory Commission (NRC) uses the Integrated Materials Performance Evaluation Program (IMPEP) in the evaluation of Agreement State programs. Enclosed for your review is the draft IMPEP report that documents the results of the Agreement State review held in Georgia on September 22-26, 2008. I was the team leader for the review. The review team's preliminary findings were discussed with you on the last day of the review. The review team's proposed recommendations are that the Georgia Agreement State Program be found adequate to protect public health and safety, but needs improvement, and compatible with NRC's program.

NRC conducts periodic reviews of Agreement State programs to ensure that public health and safety are adequately protected from the potential hazards associated with the use of radioactive materials and that Agreement State programs are compatible with NRC's program. The process, titled IMPEP, employs a team of NRC and Agreement State staff to assess Agreement State and NRC Regional radioactive materials programs. All reviews use common criteria in the assessment and place primary emphasis on performance. Three additional areas applicable to your program have been identified as non-common performance indicators and are also addressed in the assessment. A Management Review Board (MRB) composed of NRC managers and an Agreement State program manager who serves as a liaison to the MRB will make the final determination of adequacy and compatibility for each program, based on the review team's findings.

In accordance with procedures for implementation of IMPEP, we are providing you with a copy of the review team's draft report for your review and comment prior to submitting the report to the MRB. Comments are requested within 4 weeks from your receipt of this letter. This schedule will permit the issuance of the final report in a timely manner that will be responsive to your needs.

The team will review your response, make any necessary changes to the report, and issue it to the MRB as a proposed final report. Through previous coordination with Cynthia Sanders of your staff, we have scheduled the Georgia MRB meeting for December 4, 2008. The NRC will provide invitational travel for you or your designee to attend the MRB meeting at NRC Headquarters in Rockville, Maryland. NRC has video conferencing capability if it is more convenient for the State to participate through this medium. Please contact me if you desire to establish a video conference for the meeting.

C. Couch, Ph.D

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If you have any questions regarding the enclosed report, please contact me at (817) 860-8143.

Thank you for your cooperation.

Sincerely,

***/RA/***

Randy Erickson  
Regional State Agreements Officer

Enclosure:  
Draft Georgia IMPEP Report

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SUNSI Review Completed:  Yes      ADAMS:  Yes     No      Initials: RRE  
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INTEGRATED MATERIALS PERFORMANCE EVALUATION PROGRAM  
REVIEW OF THE GEORGIA AGREEMENT STATE PROGRAM

September 22-26, 2008

**DRAFT REPORT**

Enclosure

## 1.0 INTRODUCTION

This report presents the results of the review of the Georgia Agreement State Program. The review was conducted during the period of September 22-26, 2008, by a review team comprised of technical staff members from the U.S. Nuclear Regulatory Commission (NRC), the Commonwealth of Massachusetts, and the State of Ohio. Team members are identified in Appendix A. The review was conducted in accordance with the "Implementation of the Integrated Materials Performance Evaluation Program and Rescission of Final General Statement of Policy," published in the *Federal Register* on October 16, 1997, and NRC Management Directive 5.6, "Integrated Materials Performance Evaluation Program (IMPEP)," dated February 26, 2004. Preliminary results of the review, which covered the period of August 27, 2004, to September 26, 2008, were discussed with Georgia managers on the last day of the review.

[A paragraph on the results of the Management Review Board (MRB) meeting will be included in the final report.]

The Georgia Agreement State Program is administered by the Georgia Department of Natural Resources (the Department) and is located within the Program Coordination Branch (the Branch) of the Environmental Protection Division (the Division). The Branch is divided into six District Offices and five Atlanta-based programs. The Environmental Emergency Response and Radiation Control Program are Atlanta-based programs under the Branch and include the Radioactive Materials Program (the Program) and the Environmental Radiation Program that is responsible for environmental radiological surveillance and emergency response. Both are under the supervision of a single Program Manager. The Program administers the licensing and inspection duties, as well as the State's sealed source and device (SS&D) registration program. Organizational charts for the Department, the Branch, and Program are included in the report as Appendix B.

At the time of the review, the Georgia Agreement State Program regulated 521 specific licenses authorizing the possession and use of byproduct, source, and certain special nuclear materials. The review focused on the radioactive materials program as it is carried out under Section 274b (of the Atomic Energy Act, as amended) Agreement between NRC and the State of Georgia.

In preparation for the review, a questionnaire addressing the common and applicable non-common performance indicators was sent to the Office on June 13, 2008. The Office provided its initial response to the questionnaire on September 4, 2008, and provided supplemental information on September 9, 2008. A copy of the questionnaire response and supplemental information can be found in the NRC's Agencywide Documents Access and Management System (ADAMS) using the Accession Numbers ML082520665 and ML082540271, respectively.

The review team's general approach for conduct of this review consisted of: (1) examination of Georgia's response to the questionnaire; (2) review of applicable Georgia statutes and regulations; (3) analysis of quantitative information from the Program's database; (4) technical review of selected regulatory actions; (5) field accompaniments of five inspectors; and (6) interviews with managers and staff. The review team evaluated the information gathered against the established criteria for each common and applicable non-common performance

indicator and made a preliminary assessment of the Georgia Agreement State Program's performance.

Section 2.0 of this report covers the State's actions in response to recommendations made during the previous review. Results of the current review of the common performance indicators are presented in Section 3.0. Section 4.0 details the results of the review of the applicable non-common performance indicators, and Section 5.0 summarizes the review team's findings and recommendations. The review team's recommendations are comments that relate directly to program performance by the State. A response is requested from the State to all recommendations in the final report.

## 2.0 STATUS OF ITEMS IDENTIFIED IN PREVIOUS REVIEWS

During the previous review, which covered the period of April 8, 2000, to August 26, 2004, the review team made nine recommendations in regard to program performance. The current status of each of the previous recommendations is as follows:

1. The team recommends that the Program update its inspection procedures to eliminate extensions of license inspection due dates. (Section 3.2 of the 2004 IMPEP report)

Current Status: Following the 2004 review, the Program modified its inspection procedures and eliminated the provision that allowed the extension of inspection intervals based on licensee performance. This recommendation is closed.

2. The review team recommends that the Program ensure that Notices of Violation and licensee acknowledgment letters receive appropriate supervisory review and approval. (Section 3.3 of the 2004 IMPEP report)

Current Status: Following the 2004 review, the Program modified its procedures to require management review and approval of Notices of Violations and licensee acknowledgment letters. The review, approval, and disposition of these documents are tracked through an approval log. This recommendation is closed.

3. The review team recommends that the Program develop and implement a process for conducting annual accompaniments of all radiation compliance inspectors by a supervisor. (Section 3.3 of the 2004 IMPEP report)

Current Status: The Program did not develop and implement a process to ensure that all inspectors receive annual supervisory accompaniments. The Program performed staff accompaniments until late 2005 when the Program began to experience significant turnover. In an effort to expedite the training of new staff, management made the decision to forego accompanying the more experienced staff in lieu of accompanying the newer staff members. During this time, the more experienced staff did not receive supervisor accompaniments for up to 2 years. This recommendation remains open.

4. The review team recommends that the Program revise and implement procedures to address the handling of cases where inspections reveal a systemic breakdown in a licensee's radiation safety program and when a large number of health and safety violations are identified. (Section 3.3 of the 2004 IMPEP report)

Current Status: Following the 2004 review, the Program established enforcement guidance that addressed the importance of licensee compliance with Program requirements. The procedures describe the significance and severity of violations and provide a framework to address noncompliance with Program requirements. This recommendation is closed.

5. The review team recommends that the staff receive training on STP Procedure SA-300, identifying abnormal occurrences, and the schedule of reporting of significant events to the NRC Headquarters. (Section 3.5 of the 2004 IMPEP report)

Current Status: The review team verified that the Program had provided training to the staff on formerly Office of State and Tribal Programs (STP), now Office of Federal and State Materials and Environmental Management Programs (FSME) Procedure SA-300, "Reporting Material Events." This recommendation is closed.

6. The team recommends that the Program qualify one additional reviewer in SS&D evaluations to provide backup for the principal reviewer. (Section 4.2 of the 2004 IMPEP report)

Current Status: In January 2008, the Program hired an engineer as the backup to the primary SS&D reviewer; however, approximately 1 month later, the individual left the Program for another opportunity. Before the resulting vacancy could be filled, the Department imposed a hiring freeze. As a result, the Program Director must now perform all secondary SS&D reviews. This recommendation remains open.

7. The review team recommends that the Program develop written qualification requirements for SS&D reviewers. (Section 4.2 of the 2004 IMPEP report)

Current Status: In November 2006, the Program developed and implemented written procedures that identify the qualification and training requirements for SS&D reviewers. This recommendation is closed.

8. The team recommends that the Program establish an objective method to address defects and incidents involving SS&D evaluations that includes the identification of generic issues, trend analysis, and the communication of findings with other regulatory agencies. (Section 4.2 of the 2004 IMPEP report)

Current Status: Following the 2004 review, the Program established a written procedure for the investigation of SS&D defects and incidents. The Program also developed a license condition that requires manufacturers and distributors to report defects, deviations, or operations affecting the integrity of their products. This recommendation is closed.

9. The review team recommends that the staff with primary review and concurrence responsibilities for SS&D evaluations attend a training course on root cause analysis such as the NRC course "Root Cause/Incident Investigation Workshop" (G-205). (Section 4.2 of the 2004 IMPEP report)

Current Status: In November 2005, the primary SS&D reviewer completed NRC's Root Cause Analysis Course (G-205). This recommendation is closed.

### 3.0 COMMON PERFORMANCE INDICATORS

Five common performance indicators are used to review NRC Regional and Agreement State radioactive materials programs. The indicators are: (1) Technical Staffing and Training, (2) Status of Materials Inspection Program, (3) Technical Quality of Inspections, (4) Technical Quality of Licensing Actions, and (5) Technical Quality of Incident and Allegation Activities.

#### 3.1 Technical Staffing and Training

Issues central to the evaluation of this indicator include the Program's staffing level and staff turnover, as well as the technical qualifications and training histories of the staff. To evaluate this indicator, the review team examined the Program's response to the IMPEP questionnaire relative to this indicator, interviewed managers and staff, reviewed job descriptions and training records, and considered any possible workload backlogs.

When fully staffed, the Program is comprised of a Program Manager, nine Environmental Specialists (Specialist), one technical assistant, and one administrative staff member. One Specialist position is currently vacant due to a hiring freeze that was imposed earlier in the year. Specialists are assigned to one of six geographical regions within the State and are responsible for licensing, inspection, and incident response activities within that region. To be considered for a Specialist position, each candidate must possess, at a minimum, a Bachelor's degree in a science field.

The review team noted that seven Specialists left the Program during the review period. Five were fully qualified and experienced staff, and two were new hires who transferred to another environmental program shortly after starting with the Program. The Program successfully filled those vacancies and, for a time, was fully staffed. The two senior Specialists have an average of 10 years experience with the Program; five of the staff members average approximately 2 years experience; and the newest staff member was hired early in 2008.

The Program has a documented training program for Specialists that is similar to the training program established in NRC's Inspection Manual Chapter (IMC) 1246, "Formal Qualification Programs in the Nuclear Material Safety and Safeguards Program Area." The Program Manager indicated that she is committed to ensuring that newly hired Specialists receive the training required to qualify them for their positions. In addition to NRC training courses, the Program also sends Specialists to the five week health physics course held in Oak Ridge, Tennessee.

While the Program has a documented training program, they do not have a documented qualification program. Specialists are qualified through a combination of education and

experience, formal classroom training, in-house and on-the-job training, completion of specific tasks, and mentoring by more experienced staff. The Program has not developed a procedure designed to determine an individual's competency in each program area prior to authorizing them to work independently. Specialists are notified verbally by the Program Manager when she determines they are qualified. At the time of the review, five of the eight Specialists were considered fully qualified by the Program. As discussed further in Sections 3.3 and 3.4, the review team identified key areas where Specialists that were considered fully qualified could have benefitted from additional experience or on-the-job training prior to being approved to work independently. The review team recommends that the State develop, document, and implement a formal qualification program for licensing and inspection activities that includes written documentation and supervisor endorsement of competency in each program area.

Based on the IMPEP evaluation criteria, the review team recommends that Georgia's performance with respect to the indicator, Technical Staffing and Training, be found satisfactory, but needs improvement.

### 3.2 Status of Materials Inspection Program

The review team focused on five factors while reviewing this indicator: inspection frequency, overdue inspections, initial inspections of new licenses, timely dispatch of inspection findings to licensees, and performance of reciprocity inspections. The review team based its evaluation on the Program's questionnaire response relative to this indicator, data gathered from the Program's database, examination of completed inspection casework, and interviews with managers and staff.

The review team verified that the Program's inspection priorities, with one exception, are consistent with the inspection priorities prescribed by NRC's IMC 2800, "Materials Inspection Program." The review team noted that inspection reports for all gamma stereotactic radiosurgery units (gamma knife) indicated the next inspection was due in three years from the previous inspection, which is inconsistent with the Priority 2 designation identified in IMC 2800. Staff stated that the Program categorized the gamma knife as a teletherapy device, which is a specific license type identified in IMC 2800 having a three year inspection frequency. The Program committed to changing the priority code for gamma knife devices to match the two year inspection frequency found in IMC 2800.

The review team determined that, during the review period, the Program performed 229 Priority 1, 2, and 3 inspections. The Program completed 27 of these inspections overdue with four being overdue at the time of the review. The Program also completed 76 initial inspections, of which 13 were conducted overdue (greater than 12 months after license issuance). One initial inspection was overdue at the time of the review. The review team determined that the Program performed approximately 15 percent of the total Priority 1, 2, and 3, and initial inspections overdue during the review period. Discussions with Program staff revealed that most of the overdue inspections were due to staff turnover during the review period. The Program is now better staffed and the Program has a plan in place to catch up on overdue inspections.

The review team determined that the Program adequately planned for the initial set of Increased Controls inspections. The review team evaluated the Program's prioritization methodology and

found it acceptable. The Program identified 25 licensees that were subject to the Increased Controls and performed all of the initial inspections in a timely manner. At the time of the review, the Program had not performed any subsequent Increased Controls inspections of affected licensees. This issue is further discussed in Section 3.3.

The review team evaluated the Program's timeliness of issuing inspection reports. The Program has an effective and efficient process that helps ensure that inspection findings are communicated to licensees in a timely manner. Inspection findings are normally communicated to the licensee using a form similar to NRC Form 591M, "Safety Inspection Report and Compliance Inspection." These forms are generally used for minor violations or other deficiencies. A completed form is typically issued onsite at the completion of an inspection. For significant violations, the Specialist may elect to issue a Notice of Violation. Based on the 32 inspection files reviewed, the review team determined that the appropriate inspection correspondence was generally issued within 30 days of the inspection, with most being issued on site.

During the review period the Program granted approximately 180 reciprocity permits. Most of the licensees were candidates for inspection based upon the criteria in IMC 1220, "Processing of NRC Form 241 and Inspection of Agreement State Licensees Operating under 10 CFR 150.20." The review team determined that the Program inspected 20 percent of candidate licensees in only one of the four years covered by the review period; and, in each of the remaining years the Program inspected between 11 and 17 percent of candidate licensees. The review team noted staff turnover and the focus on completing routine inspections as factors why the Program did not inspect 20 percent of candidate reciprocity licensees each year.

Based on the IMPEP evaluation criteria, the review team recommends that Georgia's performance with respect to the indicator, Status of Materials Inspection Program, be found satisfactory, but needs improvement.

### 3.3 Technical Quality of Inspections

The review team evaluated inspection reports, enforcement documentation, inspection field notes, and interviewed the responsible inspectors for 32 radioactive materials inspections conducted during the review period. The casework examined included a cross-section of inspections conducted by four former and seven current inspectors and covered a wide variety of inspection types that included medical, academic, and research and development broad-scope licensees; industrial radiography; self-shielded irradiator; medical; nuclear pharmacy; and reciprocity licensees. Appendix C lists the inspection casework files reviewed and includes case-specific comments.

The review team found that the Program's inspection procedures are generally consistent with the inspection guidance found in IMC 2800. Specialists are responsible for conducting inspections of all the various types of licensees in their assigned region. Specialists use a specific inspection checklist for each license type to help ensure that all relevant aspects of a particular program are reviewed. Inspection documentation parallels the inspection checklist, limiting the amount of narrative documentation contained in routine reports; however, reports involving violations generally have additional documentation to support the enforcement action. The review team noted that inspection reports were generally consistent between Specialists

and addressed unresolved safety issues from previous inspections. The reports also noted discussions held with licensees during exit interviews.

All routine inspection documentation is entered into the Program's electronic filing system. Upon completion of the inspection, Specialists complete a "Change Form" identifying the inspection date, next due date, and inspection results. This form is then submitted to an administrative staff member who updates the Program's database with the new information.

The Program does not require management review of inspection documentation where no findings were identified, including both routine and Increased Controls inspections. Inspections identifying violations are sent to the licensees from the office following management review and approval. Based on the evaluation of casework, the review team determined that the Specialists reviewed the applicable components of the licensees' radiation safety programs.

As noted in Section 3.2, the Program successfully completed the initial round of Increased Controls inspections. However, the team found that subsequent health and safety inspections of those affected licensees did not include any followup to the Increased Controls as identified in RCPD 07-006 "Continuing Inspections of Increased Controls Licensees. In one instance the review team identified an Increased Controls inspection where violations were identified and a Notice of Violation was issued to the licensee, but the Program did not follow up with the licensee to ensure that the violations had been corrected. The review team further found that the Program's inspection procedures and enforcement guidance did not include requirements for Increased Controls inspection followup. The review team recommends that the State update their inspection procedures and enforcement guidance to include the requirements for timely followup of Increased Controls violations.

The review team determined that documents involving Increased Controls inspections were protected, segregated from the electronic file storage system, separated from license files, and maintained in a locked file cabinet with limited access. The documents observed were sufficiently marked as sensitive information to be withheld from public disclosure.

The Program has adequate numbers and types of radiation survey instruments to support their inspection efforts. Calibrated survey instruments, such as Geiger-Mueller detectors, scintillation detectors, micro-R meters and ion chambers, were available on site for the staff's use. Survey instruments and dosimeters are calibrated by licensed service providers and through a contract with the South Carolina Department of Health and Environmental Control. Laboratory sample analysis is provided through the mobile laboratory maintained by the Environmental Radiation Program. If needed, additional laboratory capabilities are available from licensed academic and research facilities within the State.

The review team noted that, over the review period, annual management accompaniments were not always performed on a routine basis. All Specialists were accompanied in 2005 and again in 2006. A significant number of staff left the Program in subsequent years and management made the decision to concentrate efforts on training new employees rather than accompanying the two senior staff. These two individuals went up to 2 years without being accompanied by management. As noted in Section 2.0, the recommendation from the 2004 review regarding supervisory accompaniments remains open.

A member of the review team accompanied five of the Program's Specialists during the week of August 4, 2008. The Specialists conducted inspections at two hospital nuclear medicine departments, one using therapeutic radioisotopes and one using only diagnostic radioisotopes; a nuclear pharmacy; a cancer center using a high dose-rate remote afterloading device (HDR); and a custom source manufacturer. Appendix C lists the inspector accompaniments. The Specialists were generally prepared for the inspections, having reviewed the previous inspection reports, and were thorough in their administrative audits of the licensees' radiation safety programs. They conducted interviews with licensee personnel and performed confirmatory measurements. The Specialists held entrance and exit meetings with the appropriate level of licensee management.

The review team determined that, while the inspections observed during the accompaniments were generally adequate to assess basic radiological health, safety, and security at the licensed facilities, the Program's newer inspectors would have benefitted from additional training prior to being authorized to work independently. The review team noted that the newer inspectors rarely observed licensed activities in progress and, instead, focused almost exclusively on documentation. One Specialist commented during his inspection that he did not understand the meaning of some of the documentation he was reviewing. Another Specialist inspecting an HDR unit stated he believed he was authorized to inspect HDR units independently before he was comfortably ready to do so.

The review team interviewed the Specialists during accompaniments and noted that some expressed concerns that they may have been released to work independently before they were fully ready. Specialists added that there were times when they performed inspections without the added benefit of having attended a training class for the type of inspection being performed, primarily because they were unable to get into the classes. Additionally, some of the staff stated that because they did not have backgrounds in health physics, they did not always fully comprehend all the complexities and nuances associated with some of the program areas they were responsible for licensing and inspecting. They indicated that while they have access to more experienced staff while in the office, they don't always have that luxury when in the field.

When notified of these observations, the Program Manager indicated that as the primary trainer for the Program with a large number of staff to train, there are certain restrictions on her time. She acknowledged that with her own administrative workload, little help to train the staff, and a pressing need to keep up with the work, she is often forced to shorten the training period so that work can be completed timely. The Program Manager stated that in an attempt to alleviate this problem, she had attempted to reorganize the current staff and hire an individual to assist with training, but was unsuccessful. She added that she is always available to help with questions and concerns, and that any Specialist can receive additional training, if requested.

Based on the IMPEP evaluation criteria, the review team recommends that Georgia's performance with respect to the indicator, Technical Quality of Inspections, be found satisfactory, but needs improvement.

### 3.4 Technical Quality of Licensing Actions

The review team interviewed the Specialists and examined completed licensing casework for 31 specific licenses. Licensing actions were reviewed for completeness, consistency, proper possession authorizations, qualifications of authorized users, adequacy of facilities and equipment, adherence to good health physics practices, financial assurance, operating and emergency procedures, appropriateness of license conditions, and overall technical quality. The casework was also reviewed for timeliness, use of appropriate deficiency letters and cover letters, reference to appropriate regulations, supporting documentation, consideration of enforcement history, pre-licensing visits, supervisory review, and proper signatures.

The casework was selected to provide a representative sample of licensing actions completed during the review period. Licensing actions selected for evaluation included one new license, six renewals, 18 amendments, and six license terminations. Files reviewed included a cross-section of license types, including: academic, medical diagnostic and therapy, brachytherapy, industrial radiography, and fixed and portable gauges. The casework sample represented work from each of the Specialists. A listing of the licensing casework reviewed, with case-specific comments, can be found in Appendix D.

The administrative staff member assigns licensing actions directly to the Specialist who is responsible for the region from which the licensing request originated. Tracking numbers are assigned and logged into a computer tracking system. Due to an unusual level of staff turnover since the 2004 review, several new Specialists are independently responsible for a geographical region. Periodically these newer Specialists receive licensing requests in program areas where they have little or no experience in reviewing, with little or no formalized on-the-job training. Some Specialists expressed concerns that they felt unqualified to conduct these reviews.

Currently, on-the-job training in the licensing process is informal and lacks an established set of qualification criteria. Utilizing State of Georgia licensing guidance that parallels NRC's NUREG-1556 series, Specialists independently review all casework assigned to them, including casework in program areas where they have little or no experience. They then submit the casework to the Program Manager for review and signature. The Program Manager provides a verbal clearance when a reviewer can independently sign licensing actions.

Overall, the review team concluded that licensing actions were adequate to protect the public health and safety; however, based on the evaluation of casework, the review team noted repeated examples of issues with thoroughness, completeness, consistency, clarity, technical quality, and adherence to existing licensing guidance. Specific examples included license terminations that were completed without receiving required documentation from the licensee verifying the appropriate disposal of radioactive sources or verifying that sealed sources were not leaking prior to transfer to another licensee; license amendments that were issued releasing locations of use where both long- and short-lived radionuclides were used without receiving close-out surveys that demonstrated the absence of residual contamination; and an industrial radiography license that did not authorize temporary job sites as requested in the licensee's application. The review team noted issues involving possession authorizations, and the application of the Increased Controls and fingerprinting requirements. Followup interviews with Specialists also revealed inconsistencies among reviewers in their application of licensing guidance that is used to perform licensing reviews.

The review team evaluated the Program's financial assurance program. The review team noted that all of the instruments were originally executed documents with original signatures and were stored in a locked safe. The review team determined that the Program's financial assurance program was adequate and that the instruments were properly secured from unauthorized access.

The review team evaluated the Program's industrial radiographer certification process. The Program offers a proctored examination to applicants several times each year in a variety of locations throughout the State. Exams are contracted through the State of Texas who grades the examinations and provides results to the Program. The review team determined the Program's overall implementation of this process to be adequate.

The team reviewed the State's program for the implementation of pre-licensing guidance. Currently, the State does not have a formal program to evaluate new applicants intended use of licensed material; however, the State does use the screening tool to evaluate requests to add risk significant quantities of radioactive material to a license.

Based on the IMPEP evaluation criteria, the review team recommends that Georgia's performance with respect the indicator, Technical Quality of Licensing Actions, be found satisfactory, but needs improvement.

### 3.5 Technical Quality of Incident and Allegation Activities

In evaluating the effectiveness of the Program's actions in responding to incidents and allegations, the review team examined the response to the questionnaire relative to this indicator, evaluated selected incidents reported for Georgia in the Nuclear Material Events Database (NMED) against those contained in the Program's files, and evaluated the casework for 17 radioactive materials incidents. A listing of the casework examined can be found in Appendix E. The review team also evaluated the Program's response to three allegations involving radioactive materials reported directly to the State during the review period.

When notified of an incident or an allegation, management and staff discuss the event and determine the level of initial response based on the health and safety risk associated with the incident. The Program maintains a database for tracking the status of all incidents and allegations. The actions taken in response to an incident are documented and filed, and if the incident meets the reporting thresholds, as established in FSME Procedure SA-300, the Program notifies NRC. If an investigation is complex and extends over a period of time, NMED is appropriately updated. The Program does not use the NMED data entry software program to place events into NMED, but instead uses an internal standardized form to document the response to incidents and allegations, and e-mails this form to the NMED contractor. Since the 2004 review, the review team identified a total of 67 incidents for Georgia in NMED, of which 28 required reporting to NRC. The review team evaluated the Program's timeliness in reporting incidents to the NRC Headquarters Operations Center, and determined that, following notification from the licensee, the Program was generally timely in reporting incidents.

The incidents selected for review included medical events; lost, stolen, or abandoned radioactive material; contamination events; damaged equipment, and transportation incidents.

The review team determined that the Program's responses to incidents were thorough, complete, and comprehensive. Initial responses were prompt and well coordinated, and the level of effort was commensurate with the health and safety significance. The Program immediately dispatched inspectors to a site when the possibility of an immediate threat to public health and safety existed. When no immediate threat was present and the Program determined that the licensee had qualified, competent individuals investigating the incident, the Program generally responded telephonically with an on-site followup at a later date. The review team noted that at the conclusion of investigations, Specialists documented findings and made appropriate notifications.

In evaluating the effectiveness of the Program's response to allegations, the review team evaluated the casework for three allegations. The review team concluded that the Program consistently took prompt and appropriate action in response to concerns raised. The review team noted that the Program thoroughly documented the investigations and retained all necessary documentation to appropriately close the allegations. The Program notified the allegers of the outcomes of the investigations when the allegers' identities were known. The review team determined that the Program adequately protected the identity of allegers.

Based on the IMPEP evaluation criteria, the review team recommends that Georgia's performance with respect to the indicator, Technical Quality of Incident and Allegation Activities, be found satisfactory.

#### 4.0 NON-COMMON PERFORMANCE INDICATORS

Four non-common performance indicators are used to review Agreement State programs: (1) Compatibility Requirements, (2) Sealed Source and Device Evaluation Program, (3) Low-Level Radioactive Waste Disposal Program, and (4) Uranium Recovery Program. Georgia's Agreement does not relinquish the authority for regulation of uranium recovery activities, so only the first three non-common performance indicators were applicable to this review.

##### 4.1 Compatibility Requirements

###### 4.1.1 Legislation

Georgia became an Agreement State in December 1969. Legislative authority to create an agency and enter into an Agreement with NRC was established in the Georgia Radiation Control Act (O.C.G.A. Title 31 Chapter 13, et seq., as amended). The Department is designated as the State's radiation control agency. The review team was informed that no new legislation was passed since the last review that would affect the Program or its authority.

###### 4.1.2 Program Elements Required for Compatibility

The Georgia Regulations for Control of Radiation, found in Chapter 391-3-17, Rules and Regulations for Radioactive Materials, apply to all ionizing radiation, whether emitted from radionuclides or devices. Georgia requires a license for possession and use of all radioactive material.

The review team examined the procedures used in the State's regulatory process and found that the process takes 9-12 months for rule promulgation. The public and other interested parties are offered an opportunity to comment on proposed rules during a 30-day comment period and during a public meeting. NRC is provided with drafts of the proposed rules for review and comment before the public comment period. Proposed rules are submitted to the Board of Natural Resources (the Board) for review and approval. The Board's calendar for rule adoption is tentatively set in January for that calendar year. All programs in the Department wishing to promulgate rules must get on the Board's calendar. After the proposed rules are approved by the Board, they are filed with the Secretary of State. The rules become final 20 days after filing with the Secretary of State. The Department's Rules and Regulations are not subject to "sunset" laws.

The review team evaluated the Program's response to the questionnaire, reviewed the status of regulations required to be adopted by the State under the Commission's adequacy and compatibility policy, and verified the adoption of regulations with data obtained from the State Regulations Status sheet that FSME maintains.

Since the previous review, the Program adopted 9 amendments in one rule package that became effective on September 24, 2008. Additionally, the Increased Controls requirements were adopted on November 10, 2005, and the fingerprinting requirements were adopted on June 2, 2008, both through the issuance of license conditions.

The review team recognized the amount of progress since the last review to address the number of overdue regulations. With the adoption of the previously mentioned rulemaking package, the State is up to date with regulation development. The Program is committed to ensuring that the State maintains compatibility with the NRC by addressing upcoming regulation changes.

Current NRC policy requires that Agreement States adopt certain equivalent regulations or legally binding requirements no later than 3 years after they become effective. The review team determined that the Program was up to date on all required NRC amendments at the time of the review.

The Program will need to address the following regulation in upcoming rulemakings or by adopting alternative legally binding requirements:

- "Requirements for Expanded Definition of Byproduct Material," 10 CFR Parts 20, 30, 31, 32, 33, 35, 61, and 150 amendment (72 FR 55864), that is due for Agreement State adoption by November 30, 2010.
- "Exemptions from Licensing, General Licenses, and Distribution of Byproduct Material: Licensing and Reporting Requirements," 10 CFR Parts 30, 31, 32 and 150 amendment (72 FR 58473), that is due for Agreement State adoption by December 17, 2010.
- "Occupational Dose Records, Labeling Containers, and Total Effective Dose Equivalent," 10 CFR Parts 19 and 20 amendment (72 FR 68043), that is due for Agreement State adoption by February 15, 2011.

Based on the IMPEP evaluation criteria, the review team recommends that Georgia's performance with respect to the indicator, Compatibility Requirements, be found satisfactory.

#### 4.2 Sealed Source and Device Evaluation Program

In reviewing this indicator, the review team used three subelements to evaluate the Program's SS&D Evaluation Program. These subelements were: (1) Technical Staffing and Training; (2) Technical Quality of the Product Evaluation Program; and (3) Evaluation of Defects and Incidents Regarding SS&Ds.

In assessing the Program's SS&D evaluation activities, the review team examined information contained in the Program's response to the IMPEP questionnaire for this indicator. The review team evaluated 8 of 27 SS&D actions processed during the review period. The Program completed seven new SS&D evaluations and issued 20 amendments and corrections to existing registrations since the last review. The review team noted the staff's use of guidance documents and procedures, interviewed staff members involved in SS&D evaluations, and verified the use of regulations and inspections to enforce commitments made in the applications.

##### 4.2.1 Technical Staffing and Training

The Program currently has two individuals who are qualified to perform safety evaluations of SS&D applications, one staff Specialist and the Program Manager. The Program previously had more trained Specialists capable of performing SS&D reviews; however, due to the significant staff turnover since the last review, the Program lost those other Specialists. In response to those losses, the Program successfully hired a Specialist with an engineering degree in January 2008; however, approximately one month later, the Specialist left the Program for a position in another environmental program. As noted in Section 2.0, the recommendation from the 2004 regarding training a backup SS&D reviewer remains open.

The review team interviewed staff members involved in the reviews and determined that they were familiar with the procedures used to evaluate sources and devices. They also had access to applicable reference documents.

##### 4.2.2 Technical Quality of the Product Evaluation Program

The review team examined eight certificates and supporting documentation (five new registrations and three amendments) for five vendors. Each of the actions was initially reviewed by a fully qualified Specialist and received secondary reviews by one of three other qualified Specialists. A listing of the SS&D certificates evaluated, with case-specific comments, can be found in Appendix F.

Analysis of the casework and interviews with staff members confirmed that the Program follows the recommended guidance from NRC's SS&D Workshop and NUREG-1556, Volume 3, Revision 1, "Consolidated Guidance About Materials Licenses – Applications for Sealed Source and Device Evaluation and Registration," with one exception as noted below. The review team determined that the depth and scope of the SS&D evaluations during the review period were adequate and addressed both the physical integrity of the product and the health and safety of

the users, the public, and the environment. The registrations clearly summarized the product evaluation and provided Specialists with adequate information to license the possession and use of the product.

The review team identified one amendment adding a new sealed source to a device certificate that was incorporated as a corrected page. This practice is not consistent with the guidance identified in NUREG 1556, Volume 3. The addition of a new sealed source must be evaluated for engineering design and safety prior to being added to the certificate. The primary Specialist noted that the new source had the same dimensions as the other registered source and that radiation profiles had been provided and approved; however, each affected page must be amended to add a new registered source.

#### 4.2.3 Evaluation of Defects and Incidents Regarding SS&Ds

The review team noted that following the 2004 review, the Program added a license condition to the licenses of manufacturers requiring them to report defects affecting the integrity of their products. The Program also developed written procedures for the investigation of defects and incidents involving SS&D devices. The procedure requires the primary SS&D reviewer to query NMED on a monthly basis to identify generic issues, analyze trends, and to communicate findings with other regulatory agencies. The review team noted that these queries were conducted twice in April 2007 and once in September 2008, which is not in accordance with the new procedures. The review team discussed with the Program the possibility of amending the NMED query procedure so that reviews are performed at a less frequent interval. The review team noted that the Program is aware of issues affecting the SS&D program via routine inspections and the Part 21 equivalent license condition.

The review team noted the Program's responsiveness to customer complaints involving devices manufactured under Georgia licenses by performing quality assurance inspections on two manufacturers since the 2004 review. These inspections were directly related to problems with devices registered by Georgia and demonstrated the Program's sensitivity to complaints.

Based on the IMPEP evaluation criteria, the review team recommends that Georgia's performance with respect to the indicator, Sealed Source and Device Evaluation Program, be found satisfactory.

#### 4.3 Low-Level Radioactive Waste Disposal Program

In 1981, NRC amended its Policy Statement, "Criteria for Guidance of States and NRC in Discontinuance of NRC Authority and Assumption Thereof by States Through Agreement," to allow a State to seek an amendment for the regulation of low-level radioactive waste (LLRW) as a separate category. Those States with Agreements with NRC prior to 1981 were determined to have continued LLRW disposal authority without the need of an amendment. Although Georgia has such disposal authority, NRC has not required States to have a program for licensing a disposal facility until such time as the State has been designated as a host State for a LLRW disposal facility. When an Agreement State has been notified or becomes aware of the need to regulate a LLRW disposal facility, it is expected to put in place a regulatory program that will meet the criteria for an adequate and compatible LLRW disposal program. There are no plans

for a commercial LLRW disposal facility in Georgia. Accordingly, the review team did not evaluate this indicator.

## 5.0 SUMMARY

As noted in Sections 3.0 and 4.0, the review team found Georgia's performance to be satisfactory, but needs improvement, for four performance indicators: Technical Staffing and Training, Status of Materials Inspection Program, Technical Quality of Inspections, and Technical Quality of Licensing Actions. The review team found Georgia's performance to be satisfactory for the three other performance indicators reviewed. The review team made two recommendations regarding the performance of the Georgia Agreement State Program and left two recommendations from the 2004 review open. Accordingly, the review team recommends that the Georgia Agreement State Program be found adequate to protect public health and safety, but needs improvement, and compatible with NRC's program. Based on the results of the current review, the review team recommends that NRC initiate a period of Monitoring of the Georgia Agreement State Program. The review team recommends that a periodic meeting take place in approximately 1 year and that next full review of the Georgia Agreement State Program take place in approximately 4 years.

Below are the recommendations, as mentioned earlier in the report, for evaluation and implementation, as appropriate, by the State.

1. The review team recommends that the State develop, document, and implement a formal qualification program for licensing and inspection activities that includes written documentation and supervisor endorsement of competency in each program area. (Section 3.1)
2. The review team recommends that the State update their inspection procedures and enforcement guidance to include the requirements for timely followup of Increased Controls violations. (Section 3.3)
3. The review team recommends that the Program develop and implement a process for conducting annual accompaniments of all radiation compliance inspectors by a supervisor. (Section 3.3 of the 2004 IMPEP report)
4. The team recommends that the Program qualify one additional reviewer in SS&D evaluations to provide backup for the principal reviewer. (Section 4.2 of the 2004 IMPEP report)

## LIST OF APPENDIXES

Appendix A	IMPEP Review Team Members
Appendix B	Georgia Organization Charts
Appendix C	Inspection Casework Reviews
Appendix D	License Casework Reviews
Appendix E	Incident Casework Reviews
Appendix F	Sealed Source & Device Casework Reviews

## APPENDIX A

### IMPEP REVIEW TEAM MEMBERS

<b>Name</b>	<b>Area of Responsibility</b>
Randy Erickson, Region IV	Team Leader Compatibility Requirements Inspector Accompaniments
James Kottan, Region I	Technical Staffing and Training Technical Quality of Incident and Allegation Activities
Bryan Parker, Region I	Status of the Materials Inspection Program
Kevin Null, Region III	Technical Quality of Licensing Actions
Stephen James, Ohio	Technical Quality of Inspections
Kenath Traegde, Massachusetts	Sealed Source and Device Evaluation Program

APPENDIX B

GEORGIA ORGANIZATION CHARTS

ADAMS ACCESSION NO.: ML082520656

## APPENDIX C

### INSPECTION CASEWORK REVIEWS

NOTE: CASEWORK LISTED WITHOUT COMMENT IS INCLUDED FOR COMPLETENESS ONLY.

File No.: 1

Licensee: Hurst Boiler and Welding  
Inspection Type: Routine, Unannounced  
Inspection Date: 6/23/08

License No.: GA 918-1  
Priority: 1  
Inspector: KR

Comment:

No follow-up of initial Special inspection on 6/23/08 inspection.

File No.: 2

Licensee: Hurst Boiler and Welding  
Inspection Type: Special, Announced  
Inspection Date: 4/4/07

License No.: GA 918-1  
Priority: 1  
Inspector: ED

File No.: 3

Licensee: JANX Integrity Group  
Inspection Type: Routine, Announced  
Inspection Date: 5/23/08

License No.: GA 1369-1  
Priority: 1  
Inspectors: EJ, JM

Comment:

No follow-up of initial Special inspection on 1/31/08 or 5/23/08 inspections.

File No.: 4

Licensee: JANX Integrity Group  
Inspection Type: Routine, Announced  
Inspection Date: 1/31/08

License No.: GA 1369-1  
Priority: 1  
Inspectors: EJ, JM

File No.: 5

Licensee: JANX Integrity Group  
Inspection Type: Special, Announced  
Inspection Date: 9/7/06

License No.: GA 1369-1  
Priority: 1  
Inspector: LP

File No.: 6

Licensee: Acuren Inspection  
Inspection Type: Routine, Unannounced  
Inspection Date: 3/19/08

License No.: GA 1115-1  
Priority: 1  
Inspector: JF

File No.: 7

Licensee: Acuren Inspection  
Inspection Type: Routine, Announced  
Inspection Date: 11/15/06

License No.: GA 1115-1  
Priority: 1  
Inspector: ED

File No.: 8

Licensee: Acuren Inspection  
Inspection Type: Special, Announced  
Inspection Date: 11/15/06

License No.: GA 1115-1  
Priority: 1  
Inspector: ED

File No.: 9

Licensee: Sowega Testing Services  
Inspection Type: Routine, Announced  
Inspection Date: 8/15/08

License No.: GA 923-1  
Priority: 1  
Inspector: KR

Comment:

No follow-up of initial Special inspection on 8/15/08 inspection.

File No.: 10

Licensee: Sowega Testing Services  
Inspection Type: Special, Announced  
Inspection Date: 4/12/07

License No.: GA 923-1  
Priority: 1  
Inspector: LP

Comment:

Initial Special inspection identified violations, NOV issued, no further follow-up.

File No.: 11

Licensee: Sowega Testing Services  
Inspection Type: Routine, Announced  
Inspection Date: 2/14/07

License No.: GA 923-1  
Priority: 1  
Inspector: LP

File No.: 12

Licensee: Theragenics  
Inspection Type: Routine, Announced  
Inspection Date: 8/27/08

License No.: GA 881-5  
Priority: 2  
Inspectors: EJ, KR, CS

File No.: 13

Licensee: Theragenics  
Inspection Type: Special, Announced  
Inspection Date: 4/23/08

License No.: GA 881-5  
Priority: 2  
Inspector: EJ

Comment:

Special Inspection report sent four months after inspection date.

File No.: 14

Licensee: Theragenics  
Inspection Type: Initial, Announced  
Inspection Date: 11/15/05

License No.: GA 881-5  
Priority: 2  
Inspector: EJ

File No.: 15

Licensee: Hopewell Design  
Inspection Type: Routine, Announced  
Inspection Date: 2/15/06

License No.: GA 1434-1  
Priority: 3  
Inspector: EJ

Comment:

Inspection report sent 4 months after inspection date.

File No.: 16

Licensee: Atlanta Oncology Associates  
Inspection Type: Routine, Unannounced  
Inspection Date: 4/14/08

License No.: GA 1178-1  
Priority: 2  
Inspector: TC

File No.: 17

Licensee: Cardinal Health  
Inspection Type: Routine, Unannounced  
Inspection Date: 3/26/08

License No.: GA 463-3MD  
Priority: 2  
Inspectors: JM, IB

File No.: 18

Licensee: The Medical Center  
Inspection Type: Routine, Announced  
Inspection Date: 8/7/08

License No.: GA 392-2  
Priority: 2  
Inspector: TC

File No.: 19

Licensee: Newton Health System  
Inspection Type: Routine, Unannounced  
Inspection Date: 8/4/08

License No.: GA 632-1  
Priority: 3  
Inspector: KS

File No.: 20

Licensee: St. Joseph's Hospital  
Inspection Type: Routine, Unannounced  
Inspection Date: 7/20/06

License No.: GA 296-4  
Priority: 3  
Inspectors: LS, LP, KS

File No.: 21

Licensee: Analytics, Inc.  
Inspection Type: Routine, Unannounced  
Inspection Date: 8/8/08

License No.: GA 742-1  
Priority: 3  
Inspector: EJ

File No.: 22

Licensee: Analytics, Inc.  
Inspection Type: Routine, Unannounced  
Inspection Date: 6/20/05

License No.: GA 742-1  
Priority: 3  
Inspector: EJ

File No.: 23

Licensee: GE Healthcare  
Inspection Type: Routine, Announced  
Inspection Date: 8/6/08

License No.: GA 1166-1MD  
Priority: 2  
Inspector: IB

File No.: 24

Licensee: GE Healthcare  
Inspection Type: Routine, Announced  
Inspection Date: 11/27/06

License No.: GA 1166-1MD  
Priority: 2  
Inspector: IB

File No.: 25

Licensee: Columbus Radiation Oncology  
Inspection Type: Routine, Unannounced  
Inspection Date: 1/22/08

License No.: GA 1256-1  
Priority: 2  
Inspector: TC

File No.: 26

Licensee: Medical College of Georgia  
Inspection Type: Special, Announced  
Inspection Dates: 6/12-16/06

License No.: GA 1110-1  
Priority: 2  
Inspector: ED

File No.: 27

Licensee: Landis International  
Inspection Type: Routine, Unannounced  
Inspection Date: 1/14/08

License No.: GA 941-1  
Priority: 3  
Inspector: KR

File No.: 28

Licensee: Emory University  
Inspection Type: Special, Announced  
Inspection Date: 2/22/07

License No.: GA 153-1  
Priority: 3  
Inspector: LS

File No.: 29

Licensee: Emory University  
Inspection Type: Routine, Announced  
Inspection Dates: 7/10-13/06

License No.: GA 153-1  
Priority: 3  
Inspector: LS

File No.: 30

Licensee: St. Joseph Hospital  
Inspection Type: Routine, Unannounced  
Inspection Date: 8/5/08

License No.: GA 359-1  
Priority: 5  
Inspector: JF

File No.: 31

Licensee: Rome Imaging Center  
Inspection Type: Routine, Announced  
Inspection Date: 8/28/08

License No.: GA 1413-1  
Priority: 3  
Inspector: JM

File No.: 32

Licensee: Nucletron  
Inspection Type: Reciprocity, Unannounced  
Inspection Date: 3/19/08

License No.: Reciprocity  
Priority: 1  
Inspector: IB

### INSPECTOR ACCOMPANIMENTS

The following inspector accompaniments were performed prior to the on-site IMPEP review:

Accompaniment No.: 1

Licensee: Newton Health System  
Inspection Type: Routine, Unannounced  
Inspection Date: 8/4/08

License No.: GA-632-1  
Priority: 3  
Inspector: KS

Accompaniment No.: 2

Licensee: St. Joseph Hospital  
Inspection Type: Routine, Unannounced  
Inspection Date: 8/5/08

License No.: GA-359-1  
Priority: 3  
Inspector: JF

Accompaniment No.: 3

Licensee: GE Healthcare  
Inspection Type: Routine, Announced  
Inspection Date: 8/6/08

License No.: GA-1166359-1MD  
Priority: 2  
Inspector: IB

Accompaniment No.: 4

Licensee: The Medical Center  
Inspection Type: Routine, Announced  
Inspection Date: 8/7/08

License No.: GA-239-2  
Priority: 2  
Inspector: TC

Accompaniment No.: 5

Licensee: Analytics, Inc.  
Inspection Type: Routine, Unannounced  
Inspection Date: 8/8/08

License No.: GA-742-1  
Priority: 3  
Inspector: EJ

## APPENDIX D

### LICENSE CASEWORK REVIEWS

NOTE: CASEWORK LISTED WITHOUT COMMENT IS INCLUDED FOR COMPLETENESS ONLY.

File No.: 1

Licensee: Allmond Lab Services

Type of Action: Termination

Date Issued: 6/2/06

License No.: GA 1340-1

Amendment No.: 03

License Reviewer: KR

Comment:

License was terminated without following up on licensee's statement that two licensed devices had been stolen.

File No.: 2

Licensee: Ram K. Puri, M.D

Type of Action: Termination

Date Issued: 10/18/06

License No.: GA 1533-1

Amendment No.: 01

License Reviewer: LP

File No.: 3

Licensee: Savannah State University

Type of Action: Amendment

Date Issued: 4/1/08

License No.: GA 1102-1

Amendment No.: 03

License Reviewer: JF

Comment:

Location of use was removed without a close-out survey.

File No.: 4

Licensee: PET Imaging Center of Savannah

Type of Action: Amendment

Date Issued: 7/16/08

License No.: GA 1122-1

Amendment No.: 02

License Reviewer: JF

Comment:

Location of use was removed without a close-out survey.

File No.: 5

Licensee: Evans Memorial Hospital

Type of Action: Renewal

Date Issued: 9/20/07

License No.: GA 999-1

Amendment No.: 04

License Reviewer: JF

File No.: 6

Licensee: Piedmont Hospital.

Type of Action: Amendment

Date Issued: 9/2/08

License No.: GA 292-1

Amendment No.: 54

License Reviewer: TC

File No.: 7

Licensee: The Medical Center

Type of Action: Amendment

Date Issued: 4/9/08

License No.: GA 239-2

Amendment No.: 26

License Reviewer: TC

Comment:

License amendment was issued without a required license condition.

File No.: 8

Licensee: St. Joseph's Hospital

Type of Action: Amendment

Date Issued: 7/8/08

License No.: GA 296-4

Amendment No.: 65

License Reviewer: KS

File No.: 9

Licensee: Hurst Boiler and Welding, Inc

Type of Action: Amendment

Date Issued: 6/2/08

License No.: GA 918-1

Amendment No.: 17

License Reviewer: KR

Comment:

License did not authorize temporary job sites as the licensee had requested in its application for license renewal.

File No.: 10

Licensee: Northside Hospital

Type of Action: Amendment

Date Issued: 6/2/08

License No.: GA 39-1

Amendment No.: 46

License Reviewer: CS

File No.: 11

Licensee: Northeast Georgia Medical Center

Type of Action: Amendment

Date Issued: 3/21/08

License No.: GA 199-1

Amendment No.: 36

License Reviewer: IB

File No.: 12

Licensee: Phoebe Putney Memorial Hospital

Type of Action: Amendment

Date Issued: 4/30/08

License No.: GA 338-1

Amendment No.: 52

License Reviewer: KR

File No.: 13

Licensee: Grady Memorial Hospital Corp.

Type of Action: Renewal

Date Issued: 8/29/08

License No.: GA 258-2

Amendment No.: 11

License Reviewer: IB

File No.: 14

Licensee: Columbus Rad. Oncology Assoc.

Type of Action: Amendment

Date Issued: 8/17/05

License No.: GA 1256-1

Amendment No.: 10

License Reviewer: LP

Comment:

License amendment was issued without a required license condition.

File No.: 15

Licensee: Atlanta Oncology Associates

Type of Action: Renewal

Date Issued: 9/5/08

License No.: GA 1178-1

Amendment No.: 17

License Reviewer: TC

Comment:

License amendment was issued without a required license condition.

File No.: 16

Licensee: Memorial Health University Medical Center

Type of Action: Renewal

Date Issued: 6/10/08

License No.: GA 84-1

Amendment No.: 58

License Reviewer: CS

Comment:

License amendment was issued without a required license condition.

File No.: 17

Licensee: John Archibold Memorial Hospital

Type of Action: Amendment

Date Issued: 11/13/07

License No.: GA 78-1

Amendment No.: 39

License Reviewer: KR

Comment:

License amendment was issued without a required license condition.

File No.: 18

Licensee: Athens Regional Med. Center

Type of Action: Amendment

Date Issued: 12/11/07

License No.: GA 4-1

Amendment No.: 52

License Reviewer: KS

Comment:

License amendment was issued without a required license condition.

File No.: 19

Licensee: Medical Center of Central Georgia

Type of Action: Amendment

Date Issued: 9/18/08

License No.: GA 764-1

Amendment No.: 35

License Reviewer: KR

File No.: 20

Licensee: Oconee Regional Cancer Center  
Type of Action: Amendment  
Date Issued: 6/10/08

License No.: GA 1227-1  
Amendment No.: 18  
License Reviewer: KR

Comment:

License amendment was issued without a required license condition.

File No.: 21

Licensee: Radiology Associates of Macon  
Type of Action: Amendment  
Date Issued: 7/17/08

License No.: GA 1319-1  
Amendment No.: 06  
License Reviewer: KR

Comment:

License amendment was issued without a required license condition.

File No.: 22

Licensee: IBT  
Type of Action: Termination  
Date Issued: 10/17/06

License No.: GA 1366-1  
Amendment No.: 09  
License Reviewer: EJ

Comment:

The termination was issued without verifying that the licensee had properly disposed of their sealed sources.

File No.: 23

Licensee: Analytics, Inc.  
Type of Action: Amendment  
Date Issued: 10/25/05

License No.: GA 742-1  
Amendment No.: 24  
License Reviewer: EJ

File No.: 24

Licensee: Redmond Regional Medical Center  
Type of Action: Renewal  
Date Issued: 9/13/07

License No.: GA 165-1  
Amendment No.: 42  
License Reviewer: EJ

File No.: 25

Licensee: Northside Hospital  
Type of Action: Renewal  
Date Issued: 11/23/05

License No.: GA 748-1  
Amendment No.: 19  
License Reviewer: IB

File No.: 26

Licensee: Emory Eastside Medical Center  
Type of Action: Amendment  
Date Issued: 6/11/07

License No.: GA 728-1  
Amendment No.: 28  
License Reviewer: IB

File No.: 27

Licensee: Digirad Imaging

Type of Action: New

Date Issued: 6/11/08

License No.: GA 1529-3

Amendment No.: N/A

License Reviewer: IB

File No.: 28

Licensee: Chickasha of Georgia

Type of Action: Termination

Date Issued: 11/7/07

License No.: GA 1294-1

Amendment No.: 05

License Reviewer: EJ

Comment:

No evidence that sealed sources were not properly leak-tested prior to transfer or upon receipt of sources by the transferee.

File No.: 29

Licensee: Ft. Valley State University

Type of Action: Termination

Date Issued: 7/11/05

License No.: GA 460-1

Amendment No.: 32

License Reviewer: KS

Comment:

No evidence that sealed sources were not properly leak-tested prior to transfer or upon receipt of sources by the transferee.

File No.: 30

Licensee: State University of West Georgia

Type of Action: Termination

Date Issued: 10/18/07

License No.: GA 129-2

Amendment No.: 07

License Reviewer: SS

File No.: 31

Licensee: Wellstar Kennestone Hospital

Type of Action: Amendment

Date Issued: 2/28/07

License No.: GA 328-1

Amendment No.: 58

License Reviewers: LS

Comment:

License amendment was issued without a required license condition.

## APPENDIX E

### INCIDENT CASEWORK REVIEWS

NOTE: CASEWORK LISTED WITHOUT COMMENT IS INCLUDED FOR COMPLETENESS ONLY.

File No.: 1

Licensee: Medical Center of Central Georgia

Date of Incident: 9/22/04

Investigation Dates: 10/25-11/17/04

License No.: GA-364-1

NMED Log No.: 040838

Type of Incident: Lost Material

Type of Investigation: Telephone/Report Review/Site

File No.: 2

Licensee: GA Department of Transportation

Date of Incident: 12/16/2004

Investigation Date: 12/16/2004

License No.: GA-50-1

NMED Log No.: 050111

Type of Incident: Damaged Equipment

Type of Investigation: Telephone/Report Review

File No.: 3

Licensee: Southern Regional Medical Center

Date of Incident: 9/15/05

Investigation Date: 9/30/05

License No.: GA-1039-1

NMED Log No.: 050676

Type of Incident: Medical Event

Type of Investigation: Telephone/Report Review

File No.: 4

Licensee: Emory University

Dates of Incident: 10/5-6/05

Investigation Date: 10/11/05

License No.: GA-153-1

NMED Log No.: 050738

Type of Incident: Contamination Event

Type of Investigation: Telephone/Report Review

File No.: 5

Licensee: East Coast Isotopes, Inc.

Date of Incident: 12/21/05

Investigation Date: 12/21/05

License No.: SC-705

NMED Log No.: Not Reported

Type of Incident: Lost Material (Recovered)

Type of Investigation: Telephone

File No.: 6

Licensee: Gwinnitt Medical Center

Date of Incident: 1/10/06

Investigation Date: 1/17/06

License No.: GA-677-1

NMED Log No.: 060047

Type of Incident: Medical Event

Type of Investigation: Telephone/Report Review

File No.: 7

Licensee: Contour Engineering

Date of Incident: 4/26/06

Investigation Date: 4/26/06

License No.: GA-1398-1

NMED Log No.: 060306

Type of Incident: Damaged Portable Gauge

Type of Investigation: Telephone/Report Review

File No.: 8

Licensee: Miller Breweries East, Inc.  
Date of Incident: 3/20-25/06  
Investigation Date: 4/20/06

License No.: GA-564-1.13  
NMED Log No.: 060281  
Type of Incident: Loss of Control  
Type of Investigation: Telephone/Report Review

File No.: 9

Licensee: Gallet and Associates  
Date of Incident: 9/27/06  
Investigation Date: 10/5/06

License No.: GA-1316-1  
NMED Log No.: 070034  
Type of Incident: Damaged Equipment  
Type of Investigation: Telephone/Report Review

File No.: 10

Licensee: Sumpter Regional Hospital  
Date of Incident: 3/1/07  
Investigation Date: 3/4/07

License No.: GA-5-1  
NMED Log No.: 070173  
Type of Incident: Loss of Material  
Type of Investigation: Site

File No.: 11

Licensee: Nova Engineering  
Date of Incident: 3/27/07  
Investigation Date: 3/27/07

License No.: GA-1323-1  
NMED Log No.: 070182  
Type of Incident: Loss of Material  
Type of Investigation: Telephone/Report Review

File No.: 12

Licensee: GE Healthcare  
Date of Incident: 11/13/07  
Investigation Date: 11/13/07

License No.: NA  
NMED Log No.: 070753  
Type of Incident: Transportation  
Type of Investigation: Site/Report Review

File No.: 13

Licensee: Kaiser Permanente  
Date of Incident: 11/19/07  
Investigation Date: 1/25/08

License No.: GA-1276-1  
NMED Log No.: 080385  
Type of Incident: Medical Event  
Type of Investigation: Telephone/Report Review

File No.: 14

Licensee: Morehouse School of Medicine  
Date of Incident: 2/4/08  
Investigation Date: 2/6/08

License No.: GA-703-1  
NMED Log No.: NA  
Type of Incident: Security  
Type of Investigation: Telephone/LLEA Followup

File No.: 15

Licensee: Imperial Sugar  
Date of Incident: 2/7/08  
Investigation Date: 2/7-9/08

License No.: GA-917-1  
NMED Log No.: NA  
Type of Incident: Damage to Fixed Gauge  
Type of Investigation: Site

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File No.: 16

Licensee: Southern Regional Medical Center

Date of Incident: 5/30/08

Investigation Date: 5/30/2008

License No.: GA-1039-1

NMED Log No.: 080388

Type of Incident: Medical Event

Type of Investigation: Telephone/Report Review

File No.: 17

Licensee: Analytics, Inc.

Date of Incident: 8/22/08

Investigation Date: 8/28/08

License No.: GA-742-1

NMED Log No.: NA

Type of Incident: Leaking Source

Type of Investigation: Telephone/Report Review

## APPENDIX F

### SEALED SOURCE & DEVICE CASEWORK REVIEWS

NOTE: CASEWORK LISTED WITHOUT COMMENT IS INCLUDED FOR COMPLETENESS ONLY.

File No.: 1

Registry No.: GA-0716-D-110-S

Manufacturer: Scantech International Pty, Ltd.

Date Issued: 2/4/08

SS&D Type: Neutron Gauge

Model No.: ANALYZER-5

Action Type: New

Comment:

Thermal tests for the cast neutron shielding were not performed for a period of one hour as specified in ANSI N43.6-1997; thermal tests were instead performed for a period of only 30 minutes.

File No.: 2

Registry No.: GA-1077-D-101-S

Manufacturer: EADS SODURN North American Inc.

Date Issued: 4/4/07

SS&D Type: Neutron Gauge

Model No.: POLAB CNA Series

Action Type: Amendment

File No.: 3

Registry No.: GA-0269-D102-S

Manufacturer: Elekta Instrument AB

Date Issued: 2/28/07

SS&D Type: Medical Teletherapy

Model No.: 23004 & 24001

Action Type: Amendment

Comments:

- a) The amendment for a new sealed source was incorporated on the registration as a correction page when it should have been incorporated as a full amendment, an action that does not comply with the guidance in NUREG-1556, Vol. 3, Rev. 1.
- b) The amendment letter was not referenced at the conclusion of the certificate.

File No.: 4

Registry No.: GA-0716-D-104-S

Manufacturer: Scantech International Pty Ltd

Date Issued: 3/23/07

SS&D Type: Sulfur, Ash, Moisture Analyzer

Model No.: 9000 & 9000 Mk2

Action Type: Amendment

File No.: 5

Registry No.: GA-0269-D-104-S

Manufacturer: Elekta Instrument AB

Date Issued: 2/26/07

SS&D Type: Medical Teletherapy

Model No.: Leksell Gamma Knife Perflexion

Action Type: New

File No.: 6

Registry No.: GA-1138-D-105-S

Manufacturer: Hopewell Designs, Inc.

Date Issued: 7/26/05

SS&D Type: Dosimeter Irradiator

Model No.: GC88

Action Type: New

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File No.: 7

Registry No.: GA-1061-S-102-S

Manufacturer: International Brachytherapy SA

SS&D Type: Medical Brachytherapy

Model No.: OptiSeed<sup>103</sup> Model 1 032p

OptiStrand<sup>103</sup>

Date Issued: 2/16/05

Action Type: New

File No.: 8

Registry No.: GA-0645-D-103-S

Manufacturer: Theragenics Corporation

SS&D Type: Medical Brachytherapy

Model No.: TheraSight<sup>TM</sup> Ocular Brachytherapy

System Model 5000

Date Issued: 7/28/04

Action Type: New