Table of Contents

Introduction ................................................................................................................................. 1
1.0 Policy..................................................................................................................................... 1
2.0 Responsibilities ......................................................................................................................... 1
    2.1 Radiation Safety Committee ............................................................................................... 1
    2.2 Radiation Safety Officer ........................................................................................................ 2
    2.3 Authorized Principal Investigator ....................................................................................... 3
3.0 Regulatory Requirements ........................................................................................................... 3
    3.1 Occupation Exposure Limits ............................................................................................... 4
    3.2 Exposure Limits to Minors .................................................................................................... 4
    3.3 Occupational Exposure Records .......................................................................................... 4
    3.4 Exposure Limits to Pregnant Employees .............................................................................. 5
    3.5 Exposure Limits to the General Public .................................................................................. 5
    3.6 Leak Testing Sealed Sources ............................................................................................... 5
    3.7 monitoring Radiation ............................................................................................................ 5
    3.8 Dosimetry Requirements ...................................................................................................... 5
    3.9 Engineering Controls ........................................................................................................... 6
    3.10 Posting Requirements ........................................................................................................ 6
4.0 Exposure Action Levels ............................................................................................................ 6
5.0 Obtaining an Authorization ........................................................................................................ 7
6.0 Amending an Authorization ..................................................................................................... 7
7.0 Terminating Authorization ....................................................................................................... 7
8.0 Radiation Safety Training ......................................................................................................... 7
9.0 Radiation Dosimetry ................................................................................................................ 8
10.0 Bioassay ................................................................................................................................. 8
    10.1 Thyroid Scan ..................................................................................................................... 8
    10.2 Urine Analysis .................................................................................................................... 8
    10.3 Summation of Dose ............................................................................................................ 9
11.0 Posting and Labeling for Radioactive Material Use ................................................................ 9
12.0 Obtaining Radioactive Material .............................................................................................. 9
    12.1 Purchasing Material .......................................................................................................... 10
    12.2 Other Institutions .............................................................................................................. 10
    12.3 Delivery ............................................................................................................................ 10
    12.4 Disposing of Packing Material .......................................................................................... 10
13.0 Inventorying Radioactive Material .......................................................................................... 10
    13.1 Incoming Material .............................................................................................................. 10
    13.2 Radioactive Waste ............................................................................................................. 11
14.0 Radiation and Contamination Monitoring .............................................................................. 11
    14.1 Survey and Wipe Test Frequency ....................................................................................... 11
    14.2 Meter Surveys .................................................................................................................... 11
    14.3 Wipe Tests ........................................................................................................................ 12
    14.4 Radiation Limits ................................................................................................................ 12
    14.5 Contamination Limits ....................................................................................................... 12
15.0 External Exposure Control ....................................................................................................... 13
    15.1 Time, Distance and Shielding ............................................................................................. 13
    15.2 Storage of Material and Waste .......................................................................................... 13
<table>
<thead>
<tr>
<th>Section</th>
<th>Title</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>16.0</td>
<td>Contamination Control</td>
<td>13</td>
</tr>
<tr>
<td>16.1</td>
<td>Contamination</td>
<td>13</td>
</tr>
<tr>
<td>16.2</td>
<td>Fume Hoods</td>
<td>14</td>
</tr>
<tr>
<td>17.0</td>
<td>Internal Exposure Control</td>
<td>14</td>
</tr>
<tr>
<td>17.1</td>
<td>Contamination Control</td>
<td>14</td>
</tr>
<tr>
<td>17.2</td>
<td>Airborne Contamination</td>
<td>14</td>
</tr>
<tr>
<td>17.3</td>
<td>No Eating, Drinking or Smoking</td>
<td>14</td>
</tr>
<tr>
<td>17.4</td>
<td>Absorption Through Skin</td>
<td>14</td>
</tr>
<tr>
<td>18.0</td>
<td>Working with Common Radioisotopes</td>
<td>14</td>
</tr>
<tr>
<td>18.1</td>
<td>General Requirements</td>
<td>14</td>
</tr>
<tr>
<td>18.2</td>
<td>Low Energy Beta Emitters</td>
<td>15</td>
</tr>
<tr>
<td>18.3</td>
<td>High Energy Beta Emitters</td>
<td>15</td>
</tr>
<tr>
<td>18.4</td>
<td>X-ray and Gamma Emitters</td>
<td>15</td>
</tr>
<tr>
<td>19.0</td>
<td>Radioactive Waste</td>
<td>16</td>
</tr>
<tr>
<td>19.1</td>
<td>Dry Solid</td>
<td>16</td>
</tr>
<tr>
<td>19.2</td>
<td>Liquid Scintillation Vials</td>
<td>16</td>
</tr>
<tr>
<td>19.3</td>
<td>Liquid Waste</td>
<td>16</td>
</tr>
<tr>
<td>19.4</td>
<td>Animal Carcasses</td>
<td>17</td>
</tr>
<tr>
<td>19.5</td>
<td>Mixed Waste</td>
<td>17</td>
</tr>
<tr>
<td>19.6</td>
<td>Decayed Waste</td>
<td>17</td>
</tr>
<tr>
<td>20.0</td>
<td>Emergency Procedures</td>
<td>18</td>
</tr>
<tr>
<td>20.1</td>
<td>Minor Spills</td>
<td>18</td>
</tr>
<tr>
<td>20.2</td>
<td>Major Spills</td>
<td>19</td>
</tr>
<tr>
<td>20.3</td>
<td>Personal Contamination</td>
<td>19</td>
</tr>
<tr>
<td>20.4</td>
<td>Radioactive Dust, Mist, etc</td>
<td>19</td>
</tr>
<tr>
<td>20.5</td>
<td>Personnel Injury</td>
<td>19</td>
</tr>
<tr>
<td>20.6</td>
<td>Unauthorized Removal of Material</td>
<td>19</td>
</tr>
<tr>
<td>20.7</td>
<td>Contact Information</td>
<td>20</td>
</tr>
<tr>
<td>Appendix A.</td>
<td>Forms</td>
<td>21</td>
</tr>
<tr>
<td>Appendix B.</td>
<td>Radioactive Hazard Group</td>
<td>27</td>
</tr>
<tr>
<td>Appendix C.</td>
<td>Notice to Employees</td>
<td>30</td>
</tr>
<tr>
<td>Appendix D.</td>
<td>Emergency Procedures for Radioisotope Users</td>
<td>32</td>
</tr>
<tr>
<td>Appendix E.</td>
<td>Instruction Concerning Prenatal Radiation Exposure</td>
<td>34</td>
</tr>
<tr>
<td>Appendix F.</td>
<td>Rules of The City of New York – Decay-In-Storage</td>
<td>47</td>
</tr>
</tbody>
</table>
Introduction

Essential elements of the College's Radiation Safety Program are presented in this Radiation Safety Manual. The safety program has been created to assist radiation users to work with radiation sources in a manner that meets their safety responsibilities in as efficient and non-intrusive manner as possible.

Radiation safety philosophy and regulatory requirements focus on the objective of ensuring that all radiation exposures are to be reduced to levels that are as far below regulatory limits as can reasonably be achieved. The College strongly supports this "As Low As Reasonably Achievable" (ALARA) safety goal. The policies and procedures found in this manual were designed to promote and achieve this goal.

The Albert Einstein College of Medicine is authorized by the City of New York's Bureau of Radiological Health to use radioactive material in education, research and development activities. The College's Radiation Safety Committee authorizes individuals to possess and use radioactive material. Prospective users must submit an application to use radioactive material to the Radiation Safety Officer for review and consideration of approval by the Radiation Safety Committee. If approved the user must abide by all regulations pertaining to the use of radioactive material set forth by the City of New York and the Albert Einstein College of Medicine.

1.0 Policy

The Albert Einstein College of Medicine of Yeshiva University (Einstein) is committed to providing a healthful and safe work environment for all employees, students and visitors. Einstein has established the Environmental Health and Safety Department and faculty committees to provide a healthful and safe work environment. Einstein delegates to the Department of Environmental Health and Safety, the Radiation Safety Officer and the Radiation Safety Committee, the task of implementing and maintaining an appropriate radiation safety program that ensures licensed activities involving radiation are conducted properly. The Radiation Safety Committee represents management when reviewing licensed activities. Any problems relating to Radiation Safety which the Committee cannot address are referred to upper management.

2.0 Responsibilities

2.1 Radiation Safety Committee: The Radiation Safety Committee (RSC) is responsible for establishing policies governing the procurement, use, storage and disposal of radioactive material and radiation producing devices. The Committee includes individuals experienced in the use of radioactive material in research at the College. The Committee consists of a Chairman, Radiation Safety Officer, representatives of management and Principal Investigators knowledgeable in the use of radioactive material. The Committee will:
   1. Meet as often as necessary to conduct business, but no less than quarterly
   2. Conduct periodic reviews and audits of the Radiation Safety Program and devote sufficient time, to review quarterly compliance inspections of authorized laboratories, reports from the RSO, results of The New York Bureau of Radiological Health
inspections and written safety procedures. Examples of program reviews include, but are not limited to, the following:

- Establish procedures to ensure compliance with the rules and regulations for Radiation Safety and recommend corrective actions for problems.
- Periodic review of protocol or user permits issued by the RSC (e.g., review of each permit at three year intervals).
- Review of letters of agreement with offsite emergency response agencies.
- Review of procedures for controlling and maintaining inventories, procurement of radioactive material, individual user and institutional cumulative possession limits, transfer of radioactive materials within the institution, and transfer of radioactive material to other persons or licensees.
- Review of audit findings (of RSC-approved users and facilities) by the radiation safety office staff.
- Conduct radiation safety evaluations of proposed users and uses.
- Develop procedures and criteria for annual training and testing each category of worker.
- Establish methods for maintaining records of the committees proceedings and radiation safety evaluations of proposed users and uses of radioactive materials.

2.2 Radiation Safety Officer: Einstein makes, through the Department of Environmental Health and Safety (EH&S), the Radiation Safety Officer (RSO) responsible for the day-to-day activities of the Radiation Protection Program established by the Radiation Safety Committee. The RSO communicates with the RSC and senior management regarding program implementation and compliance status and is available to provide advice and assistance on all radiological safety matters. The RSO reports directly to the Senior Director of the Department of EH&S.

The RSO is responsible for radiation protection at the College. This includes the general surveillance of all activities involving radioactive material and all areas where they are used. Other responsibilities include ensuring compliance with activities that are associated with New York City regulations and license conditions. The Radiation Safety staff provide a wide range of radiation protection services such as personnel monitoring, waste disposal, maintenance of required records, and consultation on the safe use of radioactive materials.

All applications for internal radioactive material licenses and license amendments including changes in location, procedures, and possession limits, are reviewed by the RSO. The RSO submits applications to the RSC for its consideration of approval. The RSO may approve amendments to existing authorizations for the Radiation Safety Committee. Any projects that are found to be a threat to health or property may be immediately suspended by the RSO.

Other responsibilities include the investigation of over exposures, accidents, spills, loss or thefts, unauthorized receipt or transfer, disposal, and any other deviation from approved radiation safety practice. The RSO is also responsible for implementing all written policies and procedures relating to radiation safety and implementing corrective actions as necessary.

The Radiation Safety Officer will:

1. Maintain surveillance of all activities involving the use of radioactive material (i.e. monitoring and surveying all areas in which radioactive materials are used).
2. Regulate compliance activities that support the rules, regulations and license conditions authorized by the City of New York.
3. Monitor and maintain the use, storage and disposal of radioactive material.
4. Provide necessary information on all aspects of radiation protection to personnel at all levels of responsibility.
5. Oversee the proper delivery and receipt of all shipments of radioactive material arriving to or leaving from the institution.
6. Distribute and process personnel radiation monitoring equipment.
7. Determine the need for and conducting of bioassays, monitor personnel radiation exposure for trends or high exposures, question individuals and their supervisors about radiation exposures approaching maximum permissible amounts and recommended appropriate remedial action.
8. Conduct training programs and otherwise instruct personnel in the proper procedures of working with radioactive material. Provide refresher training and make changes in procedures and equipment as required.
9. Supervise and coordinate the radioactive waste program, including effluent monitoring and record keeping of waste storage and disposal records.
10. Provide for the storage of radioactive materials that are not in current use.
11. Perform, or arrange for, leak tests on all sealed sources.
12. Ensure the timely calibration of all radiation survey instruments.
13. Maintain an inventory of all radioisotopes at the institution and limit the quantity of radionuclides to the amounts authorized by the license.
14. Immediately suspend any activity that is found to be a threat to public health, safety or property.
15. Supervise decontamination and recovery operations.
16. Maintain additional records not specifically addressed above; for example, records on receipts, transfers and surveys.
17. Hold periodic meetings with the RSC and provide reports to licensee management.
18. Maintain the in-house licensing program for radioisotopes.
19. Respond to and remediate all radiological emergencies.
20. Assist the RSC in the performance of its duties.

2.3 Licensed Principal Investigator: An Authorized Principal Investigator is a Principal Investigator who has received in-house approval to use radioactive material by the RSC. Licensed Principal Investigators are responsible for ensuring that students and staff using radioactive materials under their license are trained in safe laboratory practices, are familiar with the terms of the license and are complying with College policies and applicable regulations. The Radiation Safety Office offers periodic training sessions to assist the Licensed Principal Investigator in this regard.

3.0 Regulatory Requirements

The City of New York’s Bureau of Radiological Health established rules for the procurement, use and disposal of radioactive material, which can be found in the Rules of the City of New York, the Health Code, Part B, Article 175. These rules require that the College use, to the extent practicable, procedures and engineering controls to ensure that ALARA practices are maintained. In addition, the rules require that the College develop, document and implement a radiation protection program that is commensurate with the scope and extent of the program.
The College must also designate an RSO with the authority to implement the radiation protection program and establish a RSC to administer the program.

What follows are applicable regulations affecting the use of radioactive material and radiation producing devices at Einstein. For additional information refer to the Rules of the City of New York, the Health Code, Part B, Article 175. (Copy can be obtained from Radiation Safety at extension 2243, or online at http://24.97.137.100/nyc/rcny/entered.htm)

3.1 Occupational Exposure Limits: The annual occupational dose to any individual shall not exceed the following:
   a. A total effective dose equivalent to the whole body of 5 rem.
   b. The sum of the deep dose equivalent and the committed dose equivalent to any individual organ or tissue of 50 rem.
   c. An eye dose equivalent of 15 rem.
   d. A skin or extremity dose equivalent of 50 rem.

While occupational exposure at a medical research institution is negligible, the possibility that an exposure could occur does exist. For this reason, all research staff, custodial service and certain engineering staff, mailroom and receiving staff may wear dosimeters.

Exposures may occur to the hands of individuals working with larger quantities of high energy beta and gamma radiation. This results from the researcher’s hand being in close proximity to the P-32 container. For this reason a ring dosimeter is required for individuals using large quantities of P-32 and I-125. (See section 9.0).

3.2 Exposure Limits to Minors: The annual occupational dose to minors is limited to 10% of the occupational dose for adult workers listed in 3.1.
   a. A total effective dose equivalent to the whole body of 0.5 rem
   b. The sum of the deep dose equivalent and the committed dose equivalent to any individual organ or tissue of 5 rem
   c. An eye dose equivalent of 1.5 rem
   d. A skin or extremity dose equivalent of 5 rem

Typically, minors should not work with radioactive material. However, if it is absolutely necessary that they do, the individual should be wearing a monthly badge rather than the quarterly badge. This will permit better monitoring of the minor’s exposure to ensure compliance with the more restrictive limits.

3.3 Occupational Exposure Records: For each individual who may enter the licensees or registrant’s restricted area and is likely to receive, in a year, an occupational dose requiring monitoring, the licensee or registrant shall:
   a. Determine the occupational radiation dose received during the current year; and
   b. Request, in writing, the records of lifetime cumulative occupational radiation dose.

Individuals applying for a badge who have documented exposure at another institution must sign a form authorizing the release of exposure records to Einstein. See “Dosimetry Requirements” section 3.8 for more information on obtaining dosimetry.
3.4 Exposure Limits to Pregnant Employees: The dose to the embryo/fetus during the term of the pregnancy shall not exceed 500 mrem due to occupational exposure for a declared pregnant employee (see Appendix E). The licensee shall review exposure history and adjust worker conditions to ensure that the monthly dose to a declared pregnant employee does not exceed 50 mrem. The dose to the embryo/fetus shall be the sum of the dose to the deep dose equivalent to the pregnant employee and the dose to the embryo/fetus from radionuclides in the embryo/fetus and radionuclides in the declared pregnant employee. If by the time the employee declares pregnancy to the licensee the dose to the embryo/fetus has exceeded 450 mrem, the licensee shall be deemed in compliance with the Code if the additional dose to the embryo/fetus does not exceed 50 mrem during the remainder of the pregnancy.

A researcher who suspects or knows she is pregnant should request a fetal badge from radiation safety. This includes researchers who work with radioactive material and those who work in radioisotope laboratories. Radiation Safety will provide instructions regarding fetal exposure to radiation and issue the individual a monthly badge to track fetal exposure for the term of the pregnancy.

3.5 Exposure Limits to the General Public: The total effective dose to individual members of the public shall not exceed 100 mrem in a year. The dose in any unrestricted area from external sources shall not exceed 2 mrem in any one hour.

These exposure limits apply to any individual not affiliated with Albert Einstein College of Medicine who enters a laboratory; a vendor, a visitor, a relative, etc.

3.6 Leak Testing Sealed Sources: All beta/gamma and neutron sealed sources (greater than 100 microcuries) in active use will be tested for leakage at intervals not to exceed six months. All sealed sources in use (greater than 10 microcuries) designed for the purpose of emitting alpha particles will be tested at intervals not to exceed three months. Ni-63 foil sources in use (greater than 100 microcuries) will be tested at intervals not to exceed six months. Test for leakage for sealed sources shall be capable of detecting the presence of 0.005 uCi of radioactive material on a test sample.

3.7 Monitoring Radiation: Surveys shall be conducted to determine radiation levels, concentrations or quantities of radioactive material and the potential for radiological hazards that could be present. Instruments and equipment used for quantitative radiation measurements are to be calibrated at least every 12 months. See “Radiation and Contamination Monitoring” section 14.0 for monitoring requirements at Einstein.

3.8 Dosimetry Requirements: The Licensee is required to supply personal monitoring devices to individuals working with radioactive material if:
   a. An adult is likely to receive, in 1 year from an external source, a dose in excess of 10% the limits in 3.1.
   b. A minor or declared pregnant employee who is likely to receive, in 1 year from an external source, a dose in excess of 10% the limits in 3.2 and 3.3 respectively.
   c. An individual entering a high, or very high radiation area.

All dosimetry issued by the EH&S Office must be provided by a company certified by the National Voluntary Laboratory Accreditation Program (NVLAP) of the National Institute of Standards and Technology.
Personal dosimeters are worn for a period of three months (1 quarter). The dosimeters are returned to Radiation Safety at the end of the quarter when the laboratories receive their new badges. All dosimeters are then forwarded to a vendor who will process them to determine individual exposures, document the exposures, and send copies to the Radiation Safety Office. Records of individual exposures are kept on file by Radiation Safety and are available to the individual upon written request. In addition, Radiation Safety provides annual exposure records for each individual wearing a personal dosimeter. See section 9.0 “Radiation Dosimetry” for information on obtaining a dosimeter.

3.9 Engineering Controls: The Licensee must use procedural or engineering controls, such as containment or ventilation to control the concentration of radioactive material in air. For a research laboratory this may involve the use of a fume hood or glove box for work with I-125 or tritiated water.

3.10 Posting Requirements: The Licensee is required to post areas and rooms in which there is used or stored radioactive material in excess of quantities specified in the “Rules of the City of New York”.

The Licensed Principal Investigator is also required to label containers of radioactive material in which the amount of material exceeds quantities specified the Rules of the City of New York. The label must contain information regarding the isotope and an estimate of the amount of material in the container and the date the amount was estimated. See section 11.0 for posting and labeling requirements at Einstein.

4.0 Exposure Action Levels

As indicated in section 3.0, the City of New York sets annual occupational exposure limits that must not be exceeded. In order to ensure that these limits are adhered to the College sets action levels as follows:

**Level I**
Effective exposure above 125 mRem and below 375 mRem in a period of three months:
- The person is notified in writing of the amount of radiation he/she has received during the period in question.

**Level II**
Effective exposure above 375 mRem in a period of three months:
- The person is notified in writing of the exposure
- The person is visited by the Radiation Safety Officer who will make a report of their findings
- The employee will:
  1. describe how they received the exposure
  2. discuss ways of reducing future exposures

All radiation exposure should be kept as low as reasonable achievable (ALARA).
5.0 Obtaining an Authorization

The Principal Investigator wishing to use radioactive material must submit an application form with a copy of his/her CV and a map of the laboratory (see appendix A). The authorization request is forwarded to the RSO who will review it and forward it to members of the RSC if he finds it complete and accurate. The Committee members will review the authorization request and notify the RSO of their decision on the request. The approval may include stipulations that the Principal Investigator uses additional precautions or provides additional information. The authorization is good for 3 years after which the RSO will notify the Principal Investigator that they must renew their authorization in order to continue to use the material. The Principal Investigator will be asked if he/she wishes to renew the authorization, terminate it, or place the authorization on an inactive status. If the Principal Investigator prefers to go on inactive status they will need to notify the RSO in writing or e-mail to reinstate their license.

6.0 Amendment to Authorization

The Principal Investigator may wish to change the conditions of his/her authorization, which may include the addition of a radioisotope, the increase in total limits for a radioisotope, to add a laboratory or a new procedure. An amendment must be submitted to the RSO requesting a change to the Principal Investigator’s authorization (see appendix A).

7.0 Terminating Authorization to Use Radioactive Material

In the event that a Principal Investigator terminates their license to use radioactive material they are responsible to do the following:

- Notify the RSO of the termination and make arrangements to turn over all radioactive material and waste.
- Survey and decontaminate all potentially contaminated equipment and document the results of the final survey indicating the equipment is acceptable for release.
- Perform a thorough survey of the laboratory and decontaminate as required to bring contamination levels below 200 dpm/100 cm² beta/gamma.
- Document the results of the survey and provide a copy to the RSO.
- Dispose of all radioactive material properly.

8.0 Radiation Safety Training

Individuals who wish to work with radioactive material must receive hands-on training in working with the material in the laboratory and attend the Radiation Safety Training offered by the RSO. An individual knowledgeable in the use of the particular radioisotope and experimental technique to be used by the new researcher should provide the training to new users. The RSO will offer Radiation Safety Training on a monthly basis or as needed. The Principal Investigator is responsible for ensuring that all new staff members attend the training.
9.0 Radiation Dosimetry

Radiation Safety issues dosimetry to new research staff upon request by the Principal Investigator. The staff member must complete a “Dosimeter Request and/or Deletion” form (See Appendix A) and forward it to Radiation Safety. They may call x2243 to have the dosimeter mailed to them. The researchers are issued dosimeters, which are to be worn for three months on the wearer’s chest. At the end of the three months the dosimeter should be returned to Radiation Safety upon receipt of a new replacement. It is important that the used badges are returned promptly after receiving the new badges to ensure they are delivered to the vendor in a timely manner for reading. Dosimeters can still be read after a long period of time, therefore, if an old dosimeter is found please send it to the Safety Office for processing.

Individuals issued dosimetry will need to complete and sign a form letter to his/her previous employer requesting his/her exposure records.

The most frequent exposure to a researcher is to their hands from high energy beta emitters such as P-32, gamma radiation from I-125 or exposures from x-ray equipment. Therefore, an individual working with these radiation sources should wear a finger dosimeter on his/her hand. If the researcher is working with P-32 or I-125 in amounts greater than 500 uCi, they are required to wear a ring badge. It should be worn on the hand potentially receiving the greatest exposure. A researcher may request a ring dosimeter in the same manner they request a personal dosimeter.

10.0 Bioassay

10.1 Thyroid Scans: Radioactive iodine in the form of NaI can be volatile and become airborne and it is important to monitor researchers for potential internal contamination when they are working with the material. When radioactive iodine is inhaled or absorbed through the skin it collects in the thyroid. Since I-125 and I-131 are gamma emitters they can be readily detected with a NaI detector placed adjacent to the thyroid. If a researcher anticipates working with radioactive iodine in amounts greater than 1 mCi, they should contact the RSO to schedule a thyroid scan. A baseline scan is conducted prior to the experimental procedure. Six to twenty four hours after the procedure the researcher should return to the Safety Office to receive a post experiment scan to determine if they have received an uptake to the thyroid.

10.2 Urinalysis: On rare occasions a urinalysis may be required for individuals working with large quantities of radioactive material. This is of particular importance when working with tritiated water in millicurie quantities.

If a researcher is working with radioactive material in excess of the quantities specified in Table 1 the RSO will require that a bioassay be performed.

Containment Device: If a researcher is using a containment device, such as a glove box or fume hood, the RSO will take this into consideration when setting requirements for bioassays.
Table 1: Action Levels for Performing Bioassays

<table>
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<tr>
<th>Hazard Group</th>
<th>Radioisotopes</th>
<th>Activity handled at One time (mCi)</th>
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<tbody>
<tr>
<td>1</td>
<td>Am-241, Ra-226</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>Ca-45, Cl-36, Na-22, I-125</td>
<td>10</td>
</tr>
<tr>
<td>3</td>
<td>C-14, Cr-51, P-32, S-35</td>
<td>100</td>
</tr>
<tr>
<td>4</td>
<td>H-3, Rb-87, U-238</td>
<td>1000</td>
</tr>
</tbody>
</table>

10.3 Summation of Dose: Any exposure resulting from an uptake of radioactive material as determined by the bioassay will be added to the external exposure to give the total effective dose equivalent for that individual.

11.0 Posting and Labeling for Radioactive Material

Laboratories in which radioactive materials are used must be posted with certain signs and labels. “Caution Radioactive Materials” warning labels must be posted on the entrance to the lab. The following locations and equipment should also be posted or labeled:

- laboratory benches on which radioactive material is used
- plexiglass shields or containers
- refrigerators/freezers used to store material
- potentially-contaminated equipment (glassware, pipettes, centrifuges, etc.)
- waste storage cabinets and containers
- material storage containers
- fume hoods in which radioactive material is used or stored

The laboratory should also be posted with emergency procedures that describe how to respond to spills, injuries, fires, and unauthorized removal of radioactive material (see appendix D).

In addition, the City of New York’s “Notice to Employees” which describes standards for protection against radiation must be posted in each laboratory (see Appendix C).

12.0 Obtaining Radioactive Material

To order radioactive material you must be a licensed Principal Investigator or work for a licensed Principal Investigator. The Principal Investigator must be authorized to receive the particular radioisotope and amount to be ordered.
12.1 **Purchasing Material:** When purchasing radioactive material, the researcher must include their name, Principal Investigator’s name, vendor providing the material, the catalog number, radioactive isotope, and amount of material being ordered. The delivery address for all radioactive material packages must be to the “Radiation Safety Officer” at, "The Department of Environmental Health and Safety, Forchheimer Room 800”. The order must be approved by the RSO prior to being placed by the Purchasing Department. The RSO is only responsible for approving the order, not for calling in the order to the vendor or tracking the order if not received.

12.2 **Other Institutions:** Anyone wishing to receive radioactive material from another institution must notify the RSO. The institution will require a copy of Einstein’s Radioactive Materials License and may request additional information. As with routine shipments of radioactive material, orders from other institutions must be addressed to the RSO at the EH&S Office and cleared by the Radiation Safety Staff before being delivered to the laboratory.

Anyone wishing to ship radioactive material must contact the RSO with the name of the individual receiving the material, the name of the institution, and the telephone number of the individual receiving the package. A copy of the radioactive materials license for the institution will be required before the package can be shipped.

12.3 **Delivery:** All radioactive material packages must be delivered to the EH&S Office at Forchheimer 800, and be processed by the Radiation Safety Staff. Once processed, EH&S will notify the laboratory that the package has arrived. The laboratory is responsible for picking up the package in a timely manner. Film badges and lab coats should be worn when picking up the radioactive package. EH&S is not responsible for damage to items that are not picked up by the end of the business day on which they arrived.

12.4 **Disposing of Packing Material:** After the package is brought to the laboratory, the packing material must be surveyed to determine that it is free of contamination. This should be done using the wipe test method. The activity on the package should not exceed 220 dpm/100 cm². All radioactive material labels must be defaced or removed. If the packing material is free of contamination it may be disposed as non-radioactive waste. However, if contamination is found on the packing material greater than the limit above, The RSO should immediately be contacted at x2243.

Always wear gloves when handling radioactive material containers.

13.0 **Inventorying Radioactive Material**

13.1 **Incoming Material:** All radioactive material must be inventoried to ensure that the Principal Investigator does not exceed their limits. A running total should be kept of each radioisotope that the Principal Investigator has on hand. The inventory must include the all radioactive material in stock solution, samples, radioactive waste, and disposed of through the sewer (See “Record of Radioactive Materials” form in appendix A).

When incoming packages of material are received the total quantity of material received should be documented on the inventory form with the date received.
13.2 Radioactive Waste: Radioactive waste including dry/solid waste material, animal carcasses/bedding and liquid waste must be included in the inventory. This may be accomplished through estimating the amount of material for a particular radioisotope in the waste stream. You may assume that a certain percentage goes into liquid waste, the scintillation vials and dry solid waste.

14.0 Radiation and Contamination Monitoring

The Principal Investigator has two standard techniques available to monitor for the presence of contamination; the portable survey instrument or the wipe test method. The portable survey instrument may utilize a GM detector, or a NaI detector. It is used to detect gross contamination on a surface or an object. However, it is not as sensitive when attempting to detect small amounts of contamination or low energy beta radiation as the wipe test method. Therefore the wipe test method is the method of choice for conducting a thorough survey of laboratories or determining if an object is contaminated.

14.1 Survey and Wipe Test Frequency: Monthly wipe tests are required to be performed in all laboratories that are using or possessing radioactive material. In addition, a survey of the designated work areas should be performed after each use of radioactive material. Monitoring should be done with a survey meter equipped with a Geiger Mueller (GM) or Sodium Iodide detector or wipe tested if tritium is involved. All wipe test results shall be recorded in dpm/100 cm². Table 2 provides guidance regarding the frequency for conducting wipe tests for large quantities of common radioisotopes.

14.2 Meter Surveys: The GM detector is the most commonly used instrument for conducting. However, a better instrument to use if working with Gamma emitters is a Sodium Iodide detector.

Table 2: Frequency of Wipe Test Survey According to Hazard Group and Activity (mCi)

<table>
<thead>
<tr>
<th>Hazard Group</th>
<th>Radioisotopes</th>
<th>Monthly</th>
<th>Biweekly</th>
<th>Weekly</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Am-241, Ra-226</td>
<td>&lt;0.1</td>
<td>&gt;0.1 and &lt;1.0</td>
<td>&gt;1.0</td>
</tr>
<tr>
<td>2</td>
<td>Ca-45, Cl-36, Na-22, I-125</td>
<td>&lt;1.0</td>
<td>&gt;1.0 and &lt;10.0</td>
<td>&gt;10.0</td>
</tr>
<tr>
<td>3</td>
<td>C-14, Cr-51, P-32, S-35</td>
<td>&lt;10.0</td>
<td>&gt;10.0 and &lt;100.0</td>
<td>&gt;100.0</td>
</tr>
<tr>
<td>4</td>
<td>H-3, Rb-87, U-238</td>
<td>&lt;100.0</td>
<td>&gt;100.0 and &lt;1000.0</td>
<td>&gt;1000.0</td>
</tr>
</tbody>
</table>

When beta (except H-3) and gamma emitters are used in the laboratory the Principal Investigator must conduct a survey using a portable, handheld meter. The survey should be conducted in the following manner:

a. Ensure that the instrument has been calibrated within the last 12 months,

b. Ensure that the batteries have an adequate charge,

c. Check the survey instrument with a known source of material to insure that it is
responding to radiation,

d. Monitor the area very slowly at about one centimeter above the surface being monitored.

e. Document the results of the survey, include the instrument make, model number, serial number, calibration date, and readings on the survey report.

The results should be documented on a survey map of the laboratory that indicates the location of laboratory benches, refrigerators/freezers, fume hoods, desks and other distinctive items in the laboratory. The observed radiation dose levels can be documented on the survey map at the locations where readings were taken.

14.3 **Wipe Tests:** Wipe tests are performed by wiping the areas of interest with a piece of absorbent material (i.e. filter paper or Q-tip) and then determining the removable activity in a liquid scintillation or gamma counter set to detect the suspected radionuclides. The wipe test method is more sensitive than instrument surveys and should especially be used when instrument surveys indicate possible contamination. This is the only practical method of monitoring for low energy beta emitters, such as H-3, C-14 and S-35. Wipe tests should be used for all surveys conducted for the purpose of identifying and/or documenting removable contamination levels.

14.4 **Radiation Limits:** External radiation levels should be kept to less than 0.1 mRem/hr at 30 centimeters from a source’s surface and to levels as low as reasonably achievable. For most energetic beta and gamma emitters roentgens, rads, and rems may be said to be equivalent.

14.5 **Contamination Limits:** The Radiation Safety Office records removable contamination levels in terms of disintegrations per minute (dpm) per 100 square centimeters. The limits required for decontaminating a surface are listed in Table 3:

<table>
<thead>
<tr>
<th>DPM/100 cm²</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;20 dpm</td>
<td>Unrestricted areas (hallways, offices, and labs not licensed for radioactive material)</td>
</tr>
<tr>
<td>&lt;200 dpm</td>
<td>Cleanup recommended to as low as practicable levels.</td>
</tr>
<tr>
<td>&lt;100,000 dpm</td>
<td>Cleanup to less than 1000 dpm 100 dpm beta/gamma or alpha and as far below as practicable is required</td>
</tr>
</tbody>
</table>

All post-cleanup contamination surveys must be documented to demonstrate the area was decontaminated.
15.0 External Exposure Control

15.1 **Time, Distance, and Shielding:** There are three mechanisms for reducing an individual’s exposure to radiation: time, distance and shielding.

One can minimize their exposure to radiation by limiting the amount of time in proximity of a radiation source. Therefore, it is a good idea to have a thorough knowledge of the experimental technique to be used when handling radioactive material. Performing “dry runs” of procedures with non-radioactive material prior to working with active material will allow a researcher gain experience with the methods involved. As a result, the individual to complete the task in a timely manner when working with the radioactive material and help to minimize their exposure.

Increasing the distance from a source can significantly reduce exposure to radiation. The exposure levels fall off at the inverse square of the distance from the source. Therefore, if one is standing one meter from a source that has a dose rate of 40 mrem/hour at one meter and moves back two meters from the source the dose rate will reduce to 10 mrem/hour.

\[
\text{Dose at } 2 \text{ m} = 40 \text{ mrem/hr} \times \left( \frac{1 \text{ m}}{2 \text{ m}} \right)^2 = 40 \text{ mrem/hr} \times \left( \frac{1}{4} \right) = 40 \text{ mrem/hr} \times 0.25 = 10 \text{ mrem/hr}
\]

Shielding can effectively reduce exposure to radiation depending on the type, quantity and energy of the radiation. The most common types of radiation found in a medical research laboratory are: beta, gamma and x-ray radiation. The energies for these types of radiation vary from a few KeV to several MeV. Lead is the typical shielding used for x-rays and gamma radiation. Thin sheets of lead are used for the radioactive materials used in the laboratory such as I-125 and Cr-51. Plexiglass is used for shielding common beta emitting radioisotopes such as P-32. The shielding can be purchased through a number of vendors.

15.2 **Storage of Material and Waste:** Radioactive material should be stored in areas of the laboratory located away from staff. Material stored in refrigerators/freezers may need to be stored in shielded containers. The best locations for waste are in the back corner of the laboratory or in cabinets under unoccupied lab benches. P-32 waste should be stored in plexiglass waste containers, while I-125 and Cr-51 may need to be stored in lead containers. Radioactive waste storage involves strict storage and record keeping procedures. Only short lived radioisotopes such as P-32 can be reasonably kept in laboratories for decay. All other waste isotopes should be collected in the appropriate container and turned over to Radiation Safety for decay.

16.0 Contamination Control

16.1 **Containment:** Radioactive material use should be restricted to designated areas of the laboratory to minimize the potential spread of contamination. The material should be used at a work station equipped with a tray and absorbent pad. This will help reduce the spread of contamination in the event that there is an accidental spill. A survey meter should be used to monitor for contamination before and after conducting research. Volatile radioactive material should be used in a fume hood to minimize the spread of airborne contamination.
16.2 Fume Hoods: When using a fume hood it is important to confirm that it is operational. Verify that the flow rate has been checked within the last year and is at least 100 cfm while the sash is at a height of 12 inches. If it has been verified, a sticker should be affixed to the hood providing the flow rate and the date it was checked. Confirm that air is flowing into the fume hood by holding tissue paper up to the edge of the sash and note if it is being drawn in.

17.0 Internal Contamination Control

17.1 Contamination Routes: Internal contamination can result from material entering the body through three different routes: it can be inhaled, ingested or absorbed through the skin. In order for it to be inhaled the material must become airborne. This can happen if the radioactive material is volatile, an aerosol or a dry dust. Material can be ingested by the transfer of material to food or to the hand, and then transferred to the mouth. Certain materials, such as I-125 in the form of NaI or tritiated water, can be absorbed through the skin. Radioactive material can also be absorbed through wounds, cuts or through the mucous membranes in the eyes.

17.2 Airborne Contamination: When working with volatile radioactive material it is necessary to use a fume hood. This will prevent the potential inhalation of airborne material by the researcher. It is also important to avoid the splattering of radioactive materials that have the potential to generate aerosol.

17.3 No Eating, Drinking or Smoking: It is important to enforce the requirement of no eating, drinking, smoking or application of make-up in a laboratory. This minimizes the potential risk of ingesting radioactive material or any other hazardous material in the laboratory. This is a strict rule. Food and beverages shall not be brought into the laboratory or stored in refrigerators/freezers in which radioactive material is stored. In addition, mouth pipetting is not allowed.

17.4 Absorption Through Skin: Always wear a lab coat, gloves and safety glasses when working with radioactive material. You should not work with radioactive material with an open wound or sore. If you are working with tritiated water or radioactive iodine in the form of NaI, wear double gloves to provide additional protection to your hands.

18.0 Working with Common Radioisotopes

18.1 General Requirements: Follow General Precautions for working with radioactive material

Preparation
- Designate and label areas where radioactive materials are used.
- Label all containers with radiation caution tape and specify isotope.
- No eating, drinking or smoking in the laboratory.
- No mouth pipetting of radioactive material.

Conducting the Research
- Use spill trays and absorbent covering.
- Use a fume hood or glove box when handling potentially volatile material.
• Wear lab coat, disposable gloves, and safety glasses.
• Wear gloves that are appropriate for the chemicals being handled.

Post Research
• Monitor and decontaminate surfaces.
• Dispose of radioactive waste in waste containers.
• Store radioactive material in refrigerator/freezer.

18.2 Low Energy Beta Emitters (H-3, C-14, S-35)
• Follow General Precautions for working with radioactive material.
• Dosimetry is not required when handling tritium, C-14 or S-35.
• Most research involving low energy beta emitters may be performed on a laboratory bench.
• Shielding is not required.
• Use the wipe test method to monitor for contamination.
• Urinalysis is required within 24 hours after working with large quantities of material.
• Dispose of radioactive waste in accordance with requirements in section 19.0.

18.3 High Energy Beta Emitters (P-32, Sr-90)
• Follow General Precautions for working with radioactive material.
• Whole body and ring dosimetry is required when using mCi quantities of material.
• Use Lucite shielding to keep exposures to less than 0.1 mRem/Hr.
• For larger quantities of material lead may be added to the outside surface of the Lucite shield.
• Avoid looking into or working over an un-shielded container of P-32.
• Conduct dry-run experiments to ensure dexterity and speed of handling P-32.
• Routinely monitor gloves for contamination and replace if contaminated.
• Urinalysis is required within 24 hours after working with 100 mCi or greater of P-32.
• Isolate waste in a labeled, shielded container.
• Dispose of radioactive waste in accordance with requirements in Section 19.0.

18.4 X-ray and Gamma Emitters (I-125, Cr-51)
• Follow General Precautions for working with radioactive material.
• Store in shielded containers.
• Whole body and ring dosimetry is required for work with large quantities (mCi).
• Minimize exposure with lead shielding.
• Use a NaI detector or liquid scintillation counter to detect gamma and x-ray emitters.
• Urinalysis is required within 24 hours after working with 100 mCi or greater.
• Dispose of radioactive waste in accordance with requirements in Section 19.0.
19.0 Radioactive Waste

19.1 Dry Solid Waste: Dry solid waste consists of potentially contaminated items such as paper, plastic and glass. It may include gloves, absorbent pads, pipette tips and empty containers. This waste type can be broken down into two categories: long-lived and short-lived material. The long-lived material includes all radioisotopes with half-lives greater than 90 days, such as tritium and C-14. This waste type must be disposed as radioactive waste through an approved radioactive waste vendor. It should be segregated from short-lived radioactive waste in a separate waste container and scheduled for removal by Radiation Safety Staff. Short-lived radioactive waste, with half-lives less than 90 days, can be held for decay and disposed as non-radioactive waste after 10 half-lives. For example, P-32 waste may be held in storage for 6 months and then disposed as medical waste. It is strongly recommended that the Principal Investigator only store P-32 waste in the laboratory for decay. All other waste containing decayable radioisotopes should be scheduled for removal and decay through Radiation Safety. Segregation of waste by radioisotope is still required even if waste is being decayed in the lab. Otherwise, the volume of the waste could overwhelm the storage capacity. If P-32 were mixed with S-35, all of the waste in the container would have to be decayed for the half-life of S-35 (3 years). If the P-32 waste was separate from the S-35 it could be disposed in 6 months while the S-35 is disposed in 3 years.

If a PI holds the waste for decay it is important that they provide the proper shielding for the waste containers and ensure that staff are not needlessly exposed to radiation.

19.2 Liquid Scintillation Vials: Liquid scintillation vials must be collected separately from dry solid waste. All vials are required to be collected in a designated Liquid Scintillation Vial waste drum. Once the drum is full the laboratory should submit a request for EH&S to remove it. Reasonable amounts of biodegradable scintillation counting fluid may be disposed down the sink as long as the sink disposal limits for radioactive material are observed.

19.3 Liquid Waste: Liquid waste consists of aqueous, non-hazardous radioactive liquids. This waste may contain any radioisotope typically used in research and may be disposed down the drain in accordance with the limits specified in Table 4. Liquid waste with activities that exceed sink disposal limits can be either held for decay in the lab or a pick-up request can be submitted for its removal. High activity liquid waste that poses a dose risk may also be collected by Radiation Safety Staff.
### Table 4: Limits for Sink Disposal

<table>
<thead>
<tr>
<th>Isotope</th>
<th>Limit/month/lab</th>
<th>Average/day/lab</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>uCi</td>
<td>uCi</td>
</tr>
<tr>
<td>P-32</td>
<td>360</td>
<td>12</td>
</tr>
<tr>
<td>S-35</td>
<td>300</td>
<td>10</td>
</tr>
<tr>
<td>I-125</td>
<td>360</td>
<td>12</td>
</tr>
<tr>
<td>Cr-51</td>
<td>1500</td>
<td>50</td>
</tr>
<tr>
<td>H-3</td>
<td>360</td>
<td>12</td>
</tr>
<tr>
<td>C-14</td>
<td>900</td>
<td>30</td>
</tr>
</tbody>
</table>

#### 19.4 Biological Waste

**Animal Carcasses:** Animal carcasses and tissue samples containing short-lived radioisotopes must be held in a freezer for decay. If the animal carcasses/samples contain tritium or C-14 in quantities less than 0.05 uCi/gram of animal tissue, they may be treated as non-radioactive and turned over to the Institute for Animal Studies for disposal. If the concentration of H-3 or C-14 is greater than 0.05 uCi/gram, the carcass/samples must be placed in a freezer and disposed as biological radioactive waste. All other radioisotopes in animal carcasses and tissue samples must be stored in a freezer for 10 half-lives then disposed as non-radioactive waste. If you are planning to use long-lived radioisotopes in animals, your experiments must be discussed with the Radiation Safety Officer.

**Animal Bedding:** Animal bedding contaminated with long-lived radioactive material such as H-3 or C-14 must be disposed as radioactive biological waste. The bedding must be stored in a freezer until collected by Radiation Safety Staff. As with animal carcasses and tissues, bedding containing short-lived radioisotopes may be held for decay for 10 half-lives then disposed as non-radioactive biological waste.

#### 19.5 Mixed Waste:

Mixed waste is hazardous waste that contains radioactive material. It may contain organic, corrosive or other hazardous compounds. This waste must be collected by Radiation Safety Staff for disposal. Short-lived mixed waste must be stored separately from long-lived waste so it can be decayed and disposed as non-radioactive hazardous waste. Due to regulatory restrictions and the high cost of disposal, all attempts must be made to minimize the generation of mixed waste.

#### 19.6 Decayed Waste:

Waste that contains short-lived isotopes and has been held in storage for 10 half lives is considered “fully decayed radioactive waste”. Decayed waste can be disposed as non-radioactive medical waste. There are regulatory requirements associated with the proper disposal of decayed waste and it is highly recommended that Radiation Safety be contacted to remove and dispose of the waste (see appendix F). Labs surveying and disposing of their own waste must keep appropriate disposal records on file (see appendix A).
These records will be reviewed quarterly by Radiation Safety Staff and will be required to be maintained as a condition of the Principal Investigator’s radioactive materials license. Ten half lives for the commonly used radioisotopes are listed below.

Table 5: Ten Half Lives for Common Radioisotopes

<table>
<thead>
<tr>
<th>Isotope</th>
<th>Half-life</th>
<th>Minimum Storage Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>P-32</td>
<td>14.3 days</td>
<td>143 Days/6 months</td>
</tr>
<tr>
<td>Cr-51</td>
<td>27.7 days</td>
<td>277 days/1 year</td>
</tr>
<tr>
<td>I-125</td>
<td>59.6 days</td>
<td>596 days/2 years</td>
</tr>
<tr>
<td>S-35</td>
<td>87.4 days</td>
<td>874 days/2.5 years</td>
</tr>
<tr>
<td>H-3</td>
<td>12.4 years</td>
<td>N/A*</td>
</tr>
<tr>
<td>C-14</td>
<td>5730 years</td>
<td>N/A*</td>
</tr>
</tbody>
</table>

* These isotopes may not be decayed.

Radiation Safety strongly recommends that only short-lived radioisotopes such as P-32 be kept in the laboratory for decay-in-storage. All other isotopes should be turned over to the Radiation Safety Officer for decay.

20.0 Emergency Procedures

Emergency procedures must be posted in each laboratory in which radioactive material is used (see Appendix D). What follows provides additional information for responding to an emergency situation.

20.1 Minor Spills: Spills that involve no radiation hazard to personnel with survey meter readings of less than or equal to 2.5 mr/hr, or for a spill of up to 100 uCi.
   a. Immediately notify all persons in the lab.
   b. Permit access to only the minimum number of persons necessary to deal with the spill.
   c. Confine the spill by placing paper towels or absorbing pads around the perimeter of the spill.
   d. Wear gloves, lab coat, and safety glasses when cleaning up the spill.
   e. Using a soapy solution or professional decon spray, clean the spill from the outside edge inward with paper towels.
   f. Avoid spreading the contamination beyond its original area.
   g. Dispose of all contaminated paper towels in a labeled waste container.
   h. Check the area for contamination using the wipe test method.
i. Repeat decontamination efforts and wipe test process until contamination levels are below acceptable limits*.

j. Dispose of paper towels, gloves and other potentially contaminated items as radioactive waste.

* If contamination remains after four or five attempts contact the Radiation Safety Officer.

### 20.2 Major Spills

Spills that present a potential radiation hazard to personnel.

a. Notify all persons not involved in the spill cleanup to vacate the room at once.

b. If the spill is liquid, and the hands are protected, right the container.

c. If the spill is on the skin, flush thoroughly.

d. For a spill on clothing, discard outer or protective layer of clothing, and keep contamination confined to the room in which the spill occurred.

e. Switch off all fans.

f. Vacate the room.

g. Notify the Radiation Safety Office as soon as possible.

### 20.3 Personal Contamination

a. Contact Radiation Safety

b. Record initial survey results taken with a GM detector in the log if the contaminant is a high-energy beta or gamma emitter.

c. Wash the affected area with lukewarm water and mild soap for 15 minutes.

d. Do not use an abrasive soap or brush as it could abrade the skin and create a pathway for the contamination to enter the body. If the contamination is in the eyes, use an eye wash station to flush the eyes.

e. Take additional measurements of the affected area with a survey meter.

f. If no contamination is found, discontinue decon process.

Otherwise repeat decon process up to two more times. If unsuccessful after 3 attempts, stop decontamination efforts.

### 20.4 Accidents Involving Radioactive Dusts, Mists, Fumes, Organic Vapors and Gases

a. Notify all other personnel to vacate the room immediately.

b. Hold breath and switch off all air circulating devices if time permits.

c. Vacate the room, close windows, lock door and post notice.

d. Notify the Radiation Safety Officer at once.

### 20.5 Injuries to Personnel involving Radioactive Materials

a. For serious injuries call 911 to obtain EMTs and an ambulance.

b. Administer first aid to the individual.

c. Notify the EMTs that the individual is potentially contaminated with radioactive material.

d. Wash minor wounds immediately under running water while spreading the edges of the wound.

e. Report all radiation accidents to the Radiation Safety Officer as soon as possible (wounds, overexposure, ingestion and inhalation).

### 20.6 Unauthorized Removal, Theft or Loss of Radiation Source

Notify the Radiation Safety Officer, who shall in turn notify the Department of Health, Office of Radiation Control.
20.7 **Emergency Contact Information:**

- Radiation Safety Officer (718) 430-2243
- Cell Phone (646) 523-5689
- Einstein Emergency - 24 hrs. x4111

(State the problems and identify location)
APPENDIX A
Radiation Safety Forms
Dear Radiation Safety Officer:

Mr/Mrs/Ms ____________, Social Security # ____________, who is presently associated with the Albert Einstein College of Medicine, has advised us that he/she was associated with your institution from ______ to ______ and while there, worked with radioactive materials and/or radiation producing machines. I would appreciate it if you would forward all pertinent previous radiation exposure data for this person at the above address.

This information is requested under the provision of Article 175, New York City Radiological Health Code, and Title 10 CFR Part 20, Regulations.

Thank you for your cooperation.

Respectfully,

Radiation Safety Officer

/\c

AUTHORIZATION TO OBTAIN RADIATION EXPOSURE RECORDS

I hereby authorize and request that my radiation exposure records be released to the Albert Einstein College of Medicine’s Radiation Safety Office.

__________ (Signature)
Dosimeter/Film Badge Request or Deletion Form

Date: __________

Check one: Please Issue_____ Please Delete______ Lost Badge________

Investigators Name ________________________________

Department ___________________________ Series __________

Building ______________________ Room __________ Extension ______

Employee or Student Name ______________________________ Badge No. ______

Social Security No. ________________________________

Date of Birth ___________________________ Male ______ Female ______

1. Did individual have a previous badge at AECOM? Yes _____ No ______

2. Has individual had Radiation Safety Training thru AECOM’s Safety Department? Yes _____ No ____ If “Yes” please provide date of training (Individual should receive training within 2 months of receiving badge)

3. Is individual a new employee at AECOM? Yes____ No ______

4. Did individual wear a dosimeter at his/her place of employment? Yes ____ No ____ If “Yes” complete attached letter to previous employer requesting exposure records
## Sink Disposal Log

<table>
<thead>
<tr>
<th>Date</th>
<th>Isotope</th>
<th>Amount (uCi)</th>
<th>Name</th>
</tr>
</thead>
</table>

### Caution

**Radioactive Materials**

- Any other radionuclide daily average 12 uCi
- The daily average can occasionally be exceeded providing that the monthly limit is met
- All sink disposals must be recorded

### Laboratory Sink Disposal

Maximum permissible amounts of liquid radioactive waste in water soluble form that may be disposed per lab.

<table>
<thead>
<tr>
<th>Isotope</th>
<th>Monthly Limit (uCi)</th>
<th>Daily Average (uCi)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P-32</td>
<td>300</td>
<td>12</td>
</tr>
<tr>
<td>S-35</td>
<td>300</td>
<td>10</td>
</tr>
<tr>
<td>I-125</td>
<td>350</td>
<td>12</td>
</tr>
<tr>
<td>Cr-51</td>
<td>1500</td>
<td>50</td>
</tr>
<tr>
<td>C-14</td>
<td>900</td>
<td>30</td>
</tr>
<tr>
<td>H-3</td>
<td>350</td>
<td>12</td>
</tr>
</tbody>
</table>
Albert Einstein College of Medicine

Record of Radioactive Material

Investigator: ___________________ Department: ___________________ Building: _____________ Room: ___________

Radiosotope: ___________________ Chemical Form: ___________________ Date Received: ___________

<table>
<thead>
<tr>
<th>Amount Received</th>
<th>Date of Use</th>
<th>Amount Used</th>
<th>Date of Waste Disposal</th>
<th>Manner of Disposal (list amount under appropriate category)</th>
<th>Balance Remaining</th>
</tr>
</thead>
<tbody>
<tr>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>Dry Solid Waste</td>
<td>XXXXXXX</td>
</tr>
<tr>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>Slack Disposal</td>
<td>XXXXXXX</td>
</tr>
<tr>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>Collected Liquid Waste</td>
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</tr>
<tr>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>Scintillation Vials</td>
<td>XXXXXXX</td>
</tr>
<tr>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>XXXXXXX</td>
<td>Animal Carcasses</td>
<td>XXXXXXX</td>
</tr>
</tbody>
</table>

...
### SURVEY RECORD OF DISPOSED RADIOACTIVE WASTE FROM DECAY-IN-STORAGE

<table>
<thead>
<tr>
<th>DATE</th>
<th>ISO TOPE</th>
<th>DECAY START DATE</th>
<th>WASTE VOLUME (Gallons)</th>
<th>SURVEY INSTRUMENT USED</th>
<th>INSTRUMENT READINGS (mR/hr)</th>
<th>BACKGROUND</th>
<th>SURFACE OF CONTAINER</th>
<th>NAME OF SURVEYOR</th>
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The information recorded on this form is required by Title 24 section 175.103 (c)(11) of the Rules of the City of New York.
APPENDIX B
Radioactive Hazard Group
Group IV: Low radiotoxicity

$^3$H  $^{58}$Co  $^{71}$Ge  $^{87}$Rb  $^{97}$Nb  $^{103}$Rh  $^{129}$I  $^{134}$Cs  $^{187}$Re  $^{197}$Pt  $^{235}$U
$^{15}$O  $^{59}$Ni  $^{85}$Kr  $^{91}$Y  $^{96}$Tm  $^{113}$In  $^{131}$Xe  $^{135}$Cs  $^{191}$Os  $^{232}$Th  $^{238}$U
$^{37}$A  $^{69}$Zn  $^{85}$Sr  $^{93}$Zr  $^{99}$Tc  $^{125}$I  $^{135}$Xe  $^{147}$Sm  $^{193}$Pt  NatTh  NatU

*From Safe Handling of Radionuclides, IAEA Safety Standards, 1973

### TABLE II: LIMITATION ON ACTIVITIES IN VARIOUS TYPES OF WORKING PLACE OR LABORATORY*

<table>
<thead>
<tr>
<th>Radiotoxicity of radionuclides</th>
<th>Minimum significant quantity (uCi)</th>
<th>Type C</th>
<th>Type B</th>
<th>Type A</th>
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</thead>
<tbody>
<tr>
<td>1. Very high</td>
<td>0.1</td>
<td>10 uCi or less</td>
<td>10 uCi - 10 mCi</td>
<td>10 mCi or more</td>
</tr>
<tr>
<td>2. High</td>
<td>1.0</td>
<td>100 uCi or less</td>
<td>100 uCi - 100 mCi</td>
<td>100 mCi or more</td>
</tr>
<tr>
<td>3. Moderate</td>
<td>10.0</td>
<td>1 mCi or less</td>
<td>1 mCi - 1 Ci</td>
<td>1 Ci or more</td>
</tr>
<tr>
<td>4. Low</td>
<td>100.0</td>
<td>10 mCi or less</td>
<td>10 mCi - 10 Ci</td>
<td>10 Ci or more</td>
</tr>
</tbody>
</table>

Type C, Type B and Type A have the meanings normally used in the classification of laboratories for handling radioactive materials. Type C is a good quality chemical laboratory. Type B is a specially designed radioisotope laboratory. Type A is a specially designed laboratory for handling large activities of radioactive materials. In the case of a conventional modern chemical laboratory with adequate ventilation and fume hoods, as well as polished, easily cleaned, non-absorbing surfaces, etc., it would be possible to increase the upper limits of activity for Type C laboratories towards the limits for Type B laboratories for toxicity groups 3 and 4.

*From Safe Handling of Radionuclides, IAEA Safety Standards, 1973
# LABORATORY CLASSIFICATION TABLE

RADIONUCLIDES CLASSIFIED ACCORDING TO RELATIVE RADIOTOXICITY PER UNIT ACTIVITY

**Group I:** Very high radiotoxicity

<table>
<thead>
<tr>
<th>Radionuclide</th>
<th>Radionuclide</th>
<th>Radionuclide</th>
<th>Radionuclide</th>
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<tr>
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**Group II:** High radiotoxicity

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**Group III:** Moderate radiotoxicity

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APPENDIX C
Notice To Employees
# NOTICE TO EMPLOYEES

## STANDARDS FOR PROTECTION AGAINST RADIATION

### YOUR EMPLOYER'S RESPONSIBILITY

The transfer, receipt, possession or use of all sources of ionizing radiation in the City of New York is controlled by the applicable rules, regulations and orders of either the New York State Departments of Labor or the New York City Department of Health. These agencies require either the registration or licensing of all significant radiation sources and they require your employer to post or otherwise make available to you a copy of the applicable regulations, license and registration and the operating procedures applying to the work in which you are engaged and to explain relevant provisions to you. These documents are made available in the office of the Radiation Safety Officer or from the licensee.

**R.S.O.**

Licensee: Albert Einstein College of Medicine of Yeshiva University

The applicable regulation in this installation is 24 RCNY Article 175.

### YOUR RESPONSIBILITY AS A WORKER

You should familiarize yourself with the provisions of the New York City Health Code and your radioactive materials license and the operating procedures which apply to the work in which you are engaged. You should observe these provisions for your own protection and the protection of your co-workers.

### WHAT IS COVERED BY THESE REGULATIONS

1. Limits on exposure to radiation and radioactive material in controlled areas;
2. Measures to be taken after accidental exposure;
3. Personnel monitoring, surveys and equipment;
4. Caution signs, labels, and safety interlock equipment
5. Exposure records and reports; and
6. Related matters.

### REPORTS ON YOUR EXPOSURE TO RADIATION

If you work where personnel monitoring equipment is required the New York City Department of Health requires your employer to provide you, upon request, a written report of your exposure to radiation both annually and at the time that you terminate employment.

### INSPECTIONS

All activities licensed or registered with the New York City Department of Health are subject to inspection. Any notice of violation involving radiological working conditions, any proposed imposition of civil penalty or order issued pursuant to the provisions of the New York City Health Code and any response from the licensee shall be posted within two (2) days after the receipt of the documents from the Department. The Licensee’s response if any, shall be posted within two (2) working days after dispatch from the licensee. Such documents shall remain posted for a minimum of five (5) working days or until the action correcting the violation has been completed, whichever is later.

### INQUIRIES

Inquiries dealing with matters outlined above can be directed to the:

New York City Department of Health
Bureau of Radiation Control
22 Cortlandt Street
34th Floor, CN 60
New York, New York 10007

### POSTING REQUIREMENTS

Copies of this notice must be posted where employees working in or frequenting any portion of controlled areas can observe a copy on the way to or from their place of employment.
APPENDIX D
Emergency Procedures for Radioisotope Users
EMERGENCY PROCEDURES FOR RADIOISO TOPE USERS

I. Minor Spills
   Involving no Radiation Hazard to Personnel: Survey Meter readings up to 2.5 mR/hr or for weak beta emitters such as H-3 or C-14 up to 100 uCi.
   1. Notify all persons in the room at once.
   2. Permit only the minimum number of persons necessary to deal with the spill into the areas.
   3. Confine the spill into the area using paper towels or absorbent pads.
   4. Liquid Spills: Don protective gloves; drop absorbent paper onto the spill.
   5. Dry Spills: Don protective gloves; dampen area of spill thoroughly making sure contamination is not spread.
   6. Using a soapy solution or professional Decon spray and paper towels, clean the spill from the outside perimeter of the spill inward.
   7. Avoid spreading the contamination beyond its original area.
   8. Check the area for contamination using a wipe test.
   9. Clean area until contamination levels are below acceptable limits.
   10. Dispose of contaminated items as radioactive waste.

II. Major Spills
    Involving Radiation Hazards to Personnel.
    1. Notify all persons not involved in spill to vacate the room at once.
    2. If the spill is liquid, and the hands are protected, right the container.
    3. If the spill is on the skin, wash the affected area for at least 15 minutes.
    4. For a spill on clothing, remove outer garments and place in disposal bag, and keep contamination confined to the room in which the spill occurred.
    5. Switch off fans.
    6. Vacate the room.
    7. Notify the Radiation Safety Office at x2243 as soon as possible.

III. Personal Contamination
    1. Wash the affected area with lukewarm water and a mild soap for at least 15 minutes.
    2. Don not use an abrasive cleaner or brush.
    3. Survey the affected area with a GM detector if the contamination is a high-energy beta, gamma or x-ray emitter. Otherwise, repeat the process three times to thoroughly clean the area.
    4. Report the contamination to the Radiation Safety Officer.

IV. Accidents Involving Radioactive Dusts, Mists, Fumes, Organic Vapors and Gases
    1. Notify all other persons to vacate the room immediately.
    2. Hold breath and switch off all air circulating devices, if time permits.
    3. Vacate the room, close windows, lock door, and post notice at entrance.
    4. Notify the Radiation Safety Officer at x2243 at once.

V. Injuries to Personnel Involving Radioactive Materials
    1. For serious injuries, call 4111 to obtain EMTs and an ambulance.
    2. Wash minor wounds immediately under running water for at least 15 minutes to remove contamination.
    3. Make the victim comfortable and administer basic first aid, if qualified.
    4. Inform the EMTs that the individual is potentially contaminated with radioactive material.
    5. Report all radiation accidents to the Radiation Safety Officer as soon as possible (wounds, overexposure, ingestion and inhalation).

VI. Persons to be Notified in the Event of an Emergency
    Radiation Safety Officer    (718) 430-2243
    Cell Phone                  (646) 523-5689

VII. 24 Hours Emergency Number  Dial x4111 and state the problem and your location. A Safety Officer will be paged.
APPENDIX E
Instruction Concerning Prenatal Radiation Exposure
INSTRUCTION CONCERNING PRENATAL RADIATION EXPOSURE

A. INTRODUCTION

The Code of Federal Regulations in 10 CFR Part 19, “Notices, Instructions and Reports to Workers; Inspection and Investigations,” in Section 19.12, “Instructions to Workers,” requires instruction in “the health protection problems associated with exposure to radiation and/or radioactive material, in precautions or procedures to minimize exposure, and in the purposes and functions of protective devices employed.” The instructions must be “commensurate with potential radiological health protection problems present in the workplace.”

The Nuclear Regulatory Commission’s (NRC’s) regulations on radiation protection are specified in 10 CFR Part 20, “Standards for Protection Against Radiation”; and 10 CFR 20.1208, “Dose to an Embryo/Fetus,” requires licensees to “ensure that the dose to an embryo/fetus during the entire pregnancy, due to occupational exposure of a declared pregnant woman, does not exceed 0.5 rem (5 mSv).” Section 20.1208 also requires licensees to “make efforts to avoid substantial variation above a uniform monthly exposure rate to a declared pregnant woman.” A declared pregnant woman is defined in 10 CFR 20.1003 as a woman who has voluntarily informed her employer, in writing, of her pregnancy and the estimated date of conception.

This regulatory guide is intended to provide information to pregnant women, and other personnel, to help them make decisions regarding radiation exposure during pregnancy. This Regulatory Guide 8.13 supplements Regulatory Guide 8.29, “Instruction Concerning Risks from Occupational Radiation Exposure” (Ref. 1), which contains a broad discussion of the risks from exposure to ionizing radiation.

Other sections of the NRC’s regulations also specify requirements for monitoring external and internal occupational dose to a declared pregnant woman. In 10 CFR 20.1502, “Conditions Requiring Individual Monitoring of External and Internal Occupational Dose,” licensees are required to monitor the occupational dose to a declared pregnant woman, using an individual monitoring device, if it is likely that the declared pregnant woman will receive, from external sources, a deep dose equivalent in excess of 0.1 rem (1 mSv). According to Paragraph (e) of 10 CFR 20.2106, “Records of Individual Monitoring Results,” the licensee must maintain

8.13-8.13-1
records of dose to an embryo/fetus if monitoring was required, and the records of dose to the embryo/fetus must be kept with the records of dose to the declared pregnant woman. The declaration of pregnancy must be kept on file, but may be maintained separately from the dose records. The licensee must retain the required form or record until the Commission terminates each pertinent license requiring the record.

The information collections in this regulatory guide are covered by the requirements of 10 CFR Parts 19 or 20, which were approved by the Office of Management and Budget, approval numbers 3150-0044 and 3150-0014, respectively. The NRC may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number.

B. DISCUSSION

As discussed in Regulatory Guide 8.29 (Ref. 1), exposure to any level of radiation is assumed to carry with it a certain amount of risk. In the absence of scientific certainty regarding the relationship between low dose exposure and health effects, and as a conservative assumption for radiation protection purposes, the scientific community generally assumes that any exposure to ionizing radiation may cause undesirable biological effects and that the likelihood of these effects increases as the dose increases. At the occupational dose limit for the whole body of 5 rem (50 mSv) per year, the risk is believed to be very low.

The magnitude of risk of childhood cancer following in utero exposure is uncertain in that both negative and positive studies have been reported. The data from these studies “are consistent with a lifetime cancer risk resulting from exposure during gestation which is two to three times that for the adult” (NCRP Report No. 116, Ref. 2). The NRC has reviewed the available scientific literature and has concluded that the 0.5 rem (5 mSv) limit specified in 10 CFR 20.1208 provides an adequate margin of protection for the embryo/fetus. This dose limit reflects the desire to limit the total lifetime risk of leukemia and other cancers associated with radiation exposure during pregnancy.

In order for a pregnant worker to take advantage of the lower exposure limit and dose monitoring provisions specified in 10 CFR Part 20, the woman must declare her pregnancy in writing to the licensee. A form letter for declaring pregnancy is provided in this guide or the licensee may use its own form letter for declaring pregnancy. A separate written declaration should be submitted for each pregnancy.

C. REGULATORY POSITION

1. Who Should Receive Instruction

Female workers who require training under 10 CFR 19.12 should be provided with the information contained in this guide. In addition to the information contained in Regulatory Guide 8.29 (Ref. 1), this information may be included as part of the training required under 10 CFR 19.12.

2. Providing Instruction

The occupational worker may be given a copy of this guide with its Appendix, an explanation of the
contents of the guide, and an opportunity to ask questions and request additional information. The information in this guide and Appendix should also be provided to any worker or supervisor who may be affected by a declaration of pregnancy or who may have to take some action in response to such a declaration.

Classroom instruction may supplement the written information. If the licensee provides classroom instruction, the instructor should have some knowledge of the biological effects of radiation to be able to answer questions that may go beyond the information provided in this guide. Videotaped presentations may be used for classroom instruction. Regardless of whether the licensee provides classroom training, the licensee should give workers the opportunity to ask questions about information contained in this Regulatory Guide 8.13. The licensee may take credit for instruction that the worker has received within the past year at other licensed facilities or in other courses or training.

3. Licensee's Policy on Declared Pregnant Women

The instruction provided should describe the licensee's specific policy on declared pregnant women, including how those policies may affect a woman's work situation. In particular, the instruction should include a description of the licensee's policies, if any, that may affect the declared pregnant woman's work situation after she has filed a written declaration of pregnancy consistent with 10 CFR 20.1208.

The instruction should also identify who to contact for additional information as well as identify who should receive the written declaration of pregnancy. The recipient of the woman's declaration may be identified by name (e.g., John Smith), position (e.g., immediate supervisor, the radiation safety officer), or department (e.g., the personnel department).

4. Duration of Lower Dose Limits for the Embryo/Fetus

The lower dose limit for the embryo/fetus should remain in effect until the woman withdraws the declaration in writing or the woman is no longer pregnant. If a declaration of pregnancy is withdrawn, the dose limit for the embryo/fetus would apply only to the time from the estimated date of conception until the time the declaration is withdrawn. If the declaration is not withdrawn, the written declaration may be considered expired one year after submission.

5. Substantial Variations Above a Uniform Monthly Dose Rate

According to 10 CFR 20.1208(b), "The licensee shall make efforts to avoid substantial variation above a uniform monthly exposure rate to a declared pregnant woman so as to satisfy the limit in paragraph (a) of this section," that is, 0.5 rem (5 mSv) to the embryo/fetus. The National Council on Radiation Protection and Measurements (NCRP) recommends a monthly equivalent dose limit of 0.05 rem (0.5 mSv) to the embryo/fetus once the pregnancy is known (Ref. 2). In view of the NCRP recommendation, any monthly dose of less than 0.1 rem (1 mSv) may be considered as not a substantial variation above a uniform monthly dose rate and as such will not require licensee justification. However, a monthly dose greater than 0.1 rem (1 mSv) should be justified by the licensee.

8.13-8.13-3
D. IMPLEMENTATION

The purpose of this section is to provide information to licensees and applicants regarding the NRC staff’s plans for using this regulatory guide.

Unless a licensee or an applicant proposes an acceptable alternative method for complying with the specified portions of the NRC’s regulations, the methods described in this guide will be used by the NRC staff in the evaluation of instructions to workers on the radiation exposure of pregnant women.

REFERENCES


APPENDIX

QUESTIONS AND ANSWERS CONCERNING PREGNATAL RADIATION EXPOSURE

1. Why am I receiving this information?

   The NRC's regulations (in 10 CFR 19.12, "Instructions to Workers") require that licensees instruct individuals working with licensed radioactive materials in radiation protection as appropriate for the situation. The instruction below describes information that occupational workers and their supervisors should know about the radiation exposure of the embryo/fetus of pregnant women.

   The regulations allow a pregnant woman to decide whether she wants to formally declare her pregnancy to take advantage of lower dose limits for the embryo/fetus. This instruction provides information to help women make an informed decision whether to declare a pregnancy.

2. If I become pregnant, am I required to declare my pregnancy?

   No. The choice whether to declare your pregnancy is completely voluntary. If you choose to declare your pregnancy, you must do so in writing and a lower radiation dose limit will apply to your embryo/fetus. If you choose not to declare your pregnancy, you and your embryo/fetus will continue to be subject to the same radiation dose limits that apply to other occupational workers.

3. If I declare my pregnancy in writing, what happens?

   If you choose to declare your pregnancy in writing, the licensee must take measures to limit the dose to your embryo/fetus to 0.5 rem (5 millisievert) during the entire pregnancy. This is one-tenth of the dose that an occupational worker may receive in a year. If you have already received a dose exceeding 0.5 rem (5 mSv) in the period between conception and the declaration of your pregnancy, an additional dose of 0.05 rem (0.5 mSv) is allowed during the remainder of the pregnancy. In addition, 10 CFR 20.1208, "Dose to an Embryo/Fetus," requires licensees to make efforts to avoid substantial variation above a uniform monthly dose rate so that all the 0.5 rem (5 mSv) allowed dose does not occur in a short period during the pregnancy.

   This may mean that, if you declare your pregnancy, the licensee may not permit you to do some of your normal job functions if those functions would have allowed you to receive more than 0.5 rem, and you may not be able to have some emergency response responsibilities.

4. Why do the regulations have a lower dose limit for the embryo/fetus of a declared pregnant woman than for a pregnant worker who has not declared?

   A lower dose limit for the embryo/fetus of a declared pregnant woman is based on a consideration of greater sensitivity to radiation of the embryo/fetus and the involuntary nature of the exposure. Several scientific advisory groups have recommended (References 1 and 2) that the dose to the embryo/fetus be limited to a fraction of the occupational dose limit.

   8.13-8.13-5
5. What are the potentially harmful effects of radiation exposure to my embryo/fetus?

The occurrence and severity of health effects caused by ionizing radiation are dependent upon the type and total dose of radiation received, as well as the time period over which the exposure was received. See Regulatory Guide 8.29, “Instruction Concerning Risks from Occupational Exposure” (Ref. 3), for more information. The main concern is embryo/fetal susceptibility to the harmful effects of radiation such as cancer.

6. Are there any risks of genetic defects?

Although radiation injury has been induced experimentally in rodents and insects, and in the experiments was transmitted and became manifest as hereditary disorders in their offspring, radiation has not been identified as a cause of such effect in humans. Therefore, the risk of genetic effects attributable to radiation exposure is speculative. For example, no genetic effects have been documented in any of the Japanese atomic bomb survivors, their children, or their grandchildren.

7. What if I decide that I do not want any radiation exposure at all during my pregnancy?

You may ask your employer for a job that does not involve any exposure at all to occupational radiation dose, but your employer is not obligated to provide you with a job involving no radiation exposure. Even if you receive no occupational exposure at all, your embryo/fetus will receive some radiation dose (on average 75 mrem (0.75 mSv)) during your pregnancy from natural background radiation.

The NRC has reviewed the available scientific literature and concluded that the 0.5 rem (5 mSv) limit provides an adequate margin of protection for the embryo/fetus. This dose limit reflects the desire to limit the total lifetime risk of leukemia and other cancers. If this dose limit is exceeded, the total lifetime risk of cancer to the embryo/fetus may increase incrementally. However, the decision on what level of risk to accept is yours. More detailed information on potential risk to the embryo/fetus from radiation exposure can be found in References 2-10.

8. What effect will formally declaring my pregnancy have on my job status?

Only the licensee can tell you what effect a written declaration of pregnancy will have on your job status. As part of your radiation safety training, the licensee should tell you the company’s policies with respect to the job status of declared pregnant women. In addition, before you declare your pregnancy, you may want to talk to your supervisor or your radiation safety officer and ask what a declaration of pregnancy would mean specifically for you and your job status.

In many cases you can continue in your present job with no change and still meet the dose limit for the embryo/fetus. For example, most commercial power reactor workers (approximately 93%) receive, in 12 months, occupational radiation doses that are less than 0.5 rem (5 mSv) (Ref. 11). The licensee may also consider the likelihood of increased radiation exposures from accidents and abnormal events before making a decision to allow you to continue in your present job.
If your current work might cause the dose to your embryo/fetus to exceed 0.5 rem (5 mSv), the licensee has various options. It is possible that the licensee can and will make a reasonable accommodation that will allow you to continue performing your current job, for example, by having another qualified employee do a small part of the job that accounts for some of your radiation exposure.

9. What information must I provide in my written declaration of pregnancy?

You should provide, in writing, your name, a declaration that you are pregnant, the estimated date of conception (only the month and year need be given), and the date that you give the letter to the licensee. A form letter that you can use is included at the end of these questions and answers. You may use that letter, use a form letter the licensee has provided to you, or write your own letter.

10. To declare my pregnancy, do I have to have documented medical proof that I am pregnant?

NRC regulations do not require that you provide medical proof of your pregnancy. However, NRC regulations do not preclude the licensee from requesting medical documentation of your pregnancy, especially if a change in your duties is necessary in order to comply with the 0.5 rem (5 mSv) dose limit.

11. Can I tell the licensee orally rather than in writing that I am pregnant?

No. The regulations require that the declaration must be in writing.

12. If I have not declared my pregnancy in writing, but the licensee suspects that I am pregnant, do the lower dose limits apply?

No. The lower dose limits for pregnant women apply only if you have declared your pregnancy in writing. The United States Supreme Court has ruled (in United Automobile Workers International Union v. Johnson Controls, Inc., 1991) that “decisions about the welfare of future children must be left to the parents who conceive, bear, support, and raise them rather than to the employers who hire those parents” (Reference 7). The Supreme Court also ruled that your employer may not restrict you from a specific job “because of concerns about the next generation.” Thus, the lower limits apply only if you choose to declare your pregnancy in writing.

13. If I am planning to become pregnant but am not yet pregnant and I inform the licensee of that in writing, do the lower dose limits apply?

No. The requirement for lower limits applies only if you declare in writing that you are already pregnant.

14. What if I have a miscarriage or find out that I am not pregnant?

If you have declared your pregnancy in writing, you should promptly inform the licensee in writing that you are no longer pregnant. However, if you have not formally declared your pregnancy in writing, you need not inform the licensee of your non-pregnant status.

15. How long is the lower dose limit in effect?

The dose to the embryo/fetus must be limited until you withdraw your declaration in writing or you

8.13-8.13-7
inform the licensee in writing that you are no longer pregnant. If the declaration is not withdrawn, the written declaration may be considered expired one year after submission.

16. If I have declared my pregnancy in writing, can I revoke my declaration of pregnancy even if I am still pregnant?

Yes, you may. The choice is entirely yours. If you revoke your declaration of pregnancy, the lower dose limit for the embryo/fetus no longer applies.

17. What if I work under contract at a licensed facility?

The regulations state that you should formally declare your pregnancy to the licensee in writing. The licensee has the responsibility to limit the dose to the embryo/fetus.

18. Where can I get additional information?

The references to this Appendix contain helpful information, especially Reference 3, NRC’s Regulatory Guide 8.29, “Instruction Concerning Risks from Occupational Radiation Exposure,” for general information on radiation risks. The licensee should be able to give this document to you.

For information on legal aspects, see Reference 7, “The Rock and the Hard Place: Employer Liability to Fertile or Pregnant Employees and Their Unborn Children—What Can the Employer Do?” which is an article in the journal Radiation Protection Management.

You may telephone the NRC Headquarters at (301) 415-7000. Legal questions should be directed to the Office of the General Counsel, and technical questions should be directed to the Division of Industrial and Medical Nuclear Safety.

You may also telephone the NRC Regional Offices at the following numbers: Region I, (610) 337-5000; Region II, (404) 562-4400; Region III, (630) 829-9500; and Region IV, (817) 860-8100. Legal questions should be directed to the Regional Counsel, and technical questions should be directed to the Division of Nuclear Materials Safety.
REFERENCES FOR APPENDIX


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3 Single copies of regulatory guides, both active and draft, and draft NUREG documents may be obtained free of charge by writing the Reproduction and Distribution Services Section, OCIO, USNRC, Washington, DC 20555-0001, or by fax to (301)415-2289, or by email to <DISTRIBUTION@NRC.GOV>. Active guides may also be purchased from the National Technical Information Service on a standing order basis. Details on this service may be obtained by writing NTIS, 5285 Port Royal Road, Springfield, VA 22161. Copies of active and draft guides are available for inspection or copying for a fee from the NRC Public Document Room at 2120 L Street NW, Washington, DC; the PDR’s mailing address is Mail Stop LL-6, Washington, DC 20555; telephone (202)634-3273; fax (202)634-3343.

8.13-8.13-9


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Copies are available at current rates from the U.S. Government Printing Office, P.O. Box 37082, Washington, DC 20402-9328 (telephone (202)512-1800); or from the National Technical Information Service by writing NTIS at 5285 Port Royal Road, Springfield, VA 22161. Copies are available for inspection or copying for a fee from the NRC Public Document Room at 2120 L Street NW., Washington, DC; the PDR's mailing address is Mail Stop LL-6, Washington, DC 20555; telephone (202)634-3273; fax (202)634-3343.

8.13-8.13-10
FORM LETTER FOR DECLARING PREGNANCY

This form letter is provided for your convenience. To make your written declaration of pregnancy, you may fill in the blanks in this form letter, you may use a form letter the licensee has provided to you, or you may write your own letter.

DECLARATION OF PREGNANCY

To:________________________

In accordance with the NRC's regulations at 10 CFR 20.1208, “Dose to an Embryo/Fetus,” I am declaring that I am pregnant. I believe I became pregnant in___________ (only the month and year need be provided).

I understand the radiation dose to my embryo/fetus during my entire pregnancy will not be allowed to exceed 0.5 rem (5 millisievert) (unless that dose has already been exceeded between the time of conception and submitting this letter). I also understand that meeting the lower dose limit may require a change in job or job responsibilities during my pregnancy.

(Your signature)

(Your name printed)

(Date)

8.13-8.13-11
REGULATORY ANALYSIS

A separate regulatory analysis was not prepared for this regulatory guide. A regulatory analysis prepared for 10 CFR Part 20, "Standards for Protection Against Radiation" (56 FR 23360), provides the regulatory basis for this guide and examines the costs and benefits of the rule as implemented by the guide. A copy of the “Regulatory Analysis for the Revision of 10 CFR Part 20” (PNL-6712, November 1988) is available for inspection and copying for a fee at the NRC Public Document Room, 2120 L Street NW, Washington, DC, as an enclosure to Part 20 (56 FR 23360).
APPENDIX F
Rules of the City of New York
Decay-In-Storage
Rules of the City of New York -

Title 24
Department of Health and Mental Hygiene

§175.103 Medical use of radioactive materials.

(c) General technical requirements.

(11) Decay-in-storage. (i) A licensee may hold radioactive material with a physical half-life of 65 days or less for decay-in-storage before disposal in ordinary trash and is exempt from the requirements of §175.104 of this Code if the licensee:

(A) holds radioactive material for decay a minimum of 10 half-lives;
(B) monitors radioactive material at the container surface before disposal as ordinary trash and determines that its radioactivity cannot be distinguished from natural background radiation levels with a radiation detection survey instrument set on its most sensitive scale and with no interposed shielding;
(C) removes or obliterates all radiation labels; and
(D) separates and monitors each generator column individually with all radiation shielding removed to ensure that its contents have decayed to natural background radiation levels before disposal.

(ii) For radioactive material disposed in accordance with §175.103 (c)(11)(i), the licensee shall retain a record of each disposal for 3 years. The record must include the date of the disposal, the date on which the radioactive material was placed in storage, the model and serial number of the survey instrument used, the background dose rate, the radiation dose rate measured at the surface of each waste container, and the name of the individual who performed the disposal.