001. SCOPE AND AUTHORITY. This chapter provides special requirements for registrants using therapeutic radiation machines. The regulations are authorized by and implement the Nebraska Radiation Control Act, Nebraska Revised Statute (Neb. Rev. Stat.) §§ 71-3501 to 71-3520. The requirements of this 180 NAC 20 are in addition to, and not in substitution for applicable requirements in 180 NAC 1, 2, 3, 4, 6, 10, 15, and 18.

002. DEFINITIONS. The following definitions apply to this chapter.

002.01 ABSORBED DOSE RATE. Absorbed dose rate is the absorbed dose per unit time, for machines with timers, or dose monitor unit per unit time for linear accelerators.

002.02 AIR KERMA (K). Air kerma (K) is the kinetic energy released in air by ionizing radiation. Kerma is determined as the quotient of dE, where dE is the sum of the initial kinetic energies of all ionizing particles liberated by uncharged ionizing particles in air of mass dM. The International System of Units (SI) unit of air kerma is joule per kilogram and the special name for the unit of kerma is the gray (Gy).

002.03 BARRIER. Barrier has the same meaning as stated in protective barrier.

002.04 BEAM AXIS. Beam axis is the axis of rotation of the beam limiting device.

002.05 BEAM-LIMITING DEVICE. Beam-limiting device is a field defining collimator, integral to the therapeutic radiation machine, which provides a means to restrict the dimension of the useful beam.

002.06 BEAM MONITORING SYSTEM. Beam monitoring system is a system designed and installed in the radiation head to detect and measure the radiation present in the useful beam.

002.07 BEAM SCATTERING FOIL. Beam scattering foil is a thin piece of material, usually metallic, placed in the beam to scatter a beam of electrons in order to provide a more uniform electron distribution in the useful beam.

002.08 BENT BEAM LINEAR ACCELERATOR. Bent beam linear accelerator is a linear accelerator geometry in which the accelerated electron beam must change direction by passing through a bending magnet.
002.09 CHANGEABLE FILTERS. Changeable filters are any filter, exclusive of inherent filtration, which can be removed from the useful beam through any electronic, mechanical, or physical process.

002.10 CONTACT THERAPY SYSTEM. Contact therapy system is a therapeutic machine with a short target to skin distance (TSD), usually less than 5 centimeters.

002.11 CONVENTIONAL SIMULATOR. Conventional simulator is any x-ray system designed to reproduce the geometric conditions of the radiation therapy equipment.

002.12 DETECTOR. Detector has the same meaning as radiation detector.

002.13 DOSE MONITOR UNIT (DMU). Dose monitor unit (DMU) means the unit response from the beam monitoring system from which the absorbed dose can be calculated.

002.14 ELECTRONIC BRACHYTHERAPY. Electronic brachytherapy is the method of radiation therapy where an electrically generated source of ionizing radiation is placed in or near the tumor or target tissue to deliver therapeutic radiation dosage.

002.15 ELECTRONIC BRACHYTHERAPY DEVICE. Electronic brachytherapy device is a system used to produce and deliver therapeutic radiation including the x-ray tube, the control mechanism, the cooling system, and the power source.

002.16 ELECTRONIC BRACHYTHERAPY SOURCE. Electronic brachytherapy source is an x-ray tube component used in an electronic brachytherapy device.

002.17 EXTERNAL BEAM RADIATION THERAPY. External beam radiation therapy is a therapeutic irradiation in which the source of radiation is at a distance from the body.

002.18 FIELD-FLATTENING FILTER. Field-flattening filter is a filter used to homogenize the absorbed dose rate over the radiation field.

002.19 FILTER. Filter is material placed in the useful beam to change beam quality in therapeutic radiation machines subject to 180 NAC 20-006.

002.20 GANTRY. Gantry is part of a radiation therapy system supporting and allowing movements of the radiation head about a center of rotation.

002.21 HALF-VALUE LAYER (HVL). Half-value layer (HVL) is the thickness of a specified material which attenuates x-radiation or gamma radiation to an extent such that the air kerma rate, exposure rate or absorbed dose rate is reduced to one-half of the value measured without the material at the same point.

002.22 INTERLOCK. Interlock is a device preventing the start or continued operation of equipment unless certain predetermined conditions prevail.
002.23 **INTERRUPTION OF IRRADIATION.** Interruption of irradiation is the stopping of irradiation with the possibility of continuing irradiation without resetting of operating conditions at the control panel.

002.24 **IRRADIATION.** Irradiation is the exposure of a living being or matter to ionizing radiation.

002.25 **ISOCENTER.** Isocenter is the center of the sphere through which the useful beam axis passes while the gantry moves through its full range of motions.

002.26 **KILOVOLT (kV) OR KILOELECTRON VOLT (keV).** Kilovolt (kV) or kiloelectron volt (keV) is the energy equal to that acquired by a particle with one electron charge in passing through a potential difference of one thousand volts in a vacuum. The current convention is to use kilovolts (kV) for photons and kiloelectron volt (keV) for electrons.

002.27 **LEAD EQUIVALENT.** Lead equivalent is the thickness of the material in question affording the same attenuation, under specified conditions, as lead.

002.28 **LEAKAGE RADIATION.** Leakage radiation is radiation emanating from the radiation therapy system except for the useful beam.

002.29 **LIGHT FIELD.** Light field is the area illuminated by light, simulating the radiation field.

002.30 **MA.** The term mA is milliampere (mA).

002.31 **MEGAVOLT (MV) OR MEGAELECTRON VOLT (MeV).** Megavolt (MV) or megaelectron volt (MeV) is the energy equal to that acquired by a particle with one electron charge in passing through a potential difference of one million volts in a vacuum. The current convention is to use megavolt (MV) for photons and megaelectron volt (MeV) for electrons.

002.32 **MISADMINISTRATION.** Misadministration is an event that meets the criteria in 180 NAC 20-005.02.

002.33 **MOBILE ELECTRONIC BRACHYTHERAPY SERVICE.** Mobile electronic brachytherapy service is the transportation of an electronic brachytherapy device to provide electronic brachytherapy at an address that is not the address of record.

002.34 **MONITOR UNIT (MU).** Monitor unit (MU) has the same meaning as dose monitor unit.

002.35 **MOVING BEAM RADIATION THERAPY.** Moving beam radiation therapy is radiation therapy with any planned displacement of radiation field or patient relative to each other, or with any planned change of absorbed dose distribution. Includes arc, skip, conformal, intensity modulation, and rotational therapy.

002.36 **NOMINAL TREATMENT DISTANCE.** Nominal treatment distance is:
(A) For electron irradiation, the distance from the scattering foil, virtual source, or exit window of the electron beam to the entrance surface of the irradiated object along the central axis of the useful beam; or
(B) For x-ray irradiation, the virtual source or target to isocenter distance along the central axis of the useful beam. For non-isocentric equipment, this distance must be that specified by the manufacturer.

002.37  PATIENT. Patient is an individual subjected to machine produced external beam radiation for the purposes of medical therapy.

002.38  PEAK TUBE POTENTIAL. Peak tube potential is the maximum value of the potential difference across the x-ray tube during an exposure.

002.39  PERIODIC QUALITY ASSURANCE CHECK. Periodic quality assurance check is a procedure which is performed to ensure that a previous calibration continues to be valid.

002.40  PHANTOM. Phantom is an object behaving in essentially the same manner as tissue, with respect to the absorption or scattering of the ionizing radiation in question.

002.41  PRESCRIBED DOSE. Prescribed dose is the total dose and dose per fraction as documented in the written directive. The prescribed dose is an estimation from measured data from a specified therapeutic machine using assumptions that are clinically acceptable for that treatment technique and historically consistent with the clinical calculations previously used for patients treated with the same clinical technique.

002.42  PRIMARY DOSE MONITORING SYSTEM. Primary dose monitoring system is the system which will monitor the useful beam during irradiation and which will terminate irradiation when a pre-selected number of dose monitor units have been delivered.

002.43  PROTECTIVE BARRIER. Protective barrier is a barrier of radiation absorbing material or materials used to reduce radiation exposure. The types of protective barriers are as follows:

002.43(A)  PRIMARY PROTECTIVE BARRIER. Primary protective barrier is the material, excluding filters, placed in the useful beam.

002.43(B)  SECONDARY PROTECTIVE BARRIER. Secondary protective barrier is the material which attenuates stray radiation.

002.44  RADIATION DETECTOR. Radiation detector is a device which, in the presence of radiation provides, by either direct or indirect means, a signal or other indication suitable for use in measuring one or more quantities of incident radiation.

002.45  RADIATION FIELD. Radiation field has the same meaning as stated in useful beam.

002.46  RADIATION HEAD. Radiation head is the structure from which the useful beam emerges.
002.47 **RADIOLOGICAL MEDICAL PHYSICIST.** Radiological medical physicist is an individual qualified according to 180 NAC 15-004.01.

002.48 **REDUNDANT BEAM MONITORING SYSTEM.** Redundant beam monitoring system is a combination of two independent dose monitoring systems in which each system is designed to terminate irradiation according to a pre-selected number of dose monitor units.

002.49 **SCATTERED RADIATION.** Scattered radiation is an ionizing radiation emitted by interaction of ionizing radiation with matter, the interaction being accompanied by a change in direction of the radiation. Scattered primary radiation means that scattered radiation which has been deviated in direction only by materials irradiated by the useful beam.

002.50 **SECONDARY DOSE MONITORING SYSTEM.** Secondary dose monitoring system is a system which will terminate irradiation in the event of failure of the primary dose monitoring system.

002.51 **SHADOW TRAY.** Shadow tray is a device attached to the radiation head to support auxiliary beam blocking material.

002.52 **SHUTTER.** Shutter is a device attached to the tube housing assembly which can totally intercept the useful beam and which has a lead equivalency not less than that of the tube housing assembly.

002.53 **SIMULATOR OR RADIATION THERAPY SIMULATION SYSTEM.** Simulator or radiation therapy simulation system is any x-ray system intended for localizing the volume to be exposed during radiation therapy and establishing the position and size of the therapeutic irradiation field. See conventional simulator and virtual simulator.

002.54 **SOURCE.** Source is the region or material from which the radiation emanates.

002.55 **SOURCE SKIN DISTANCE (SSD).** The source skin distance (SSD) has the same meaning as target skin distance.

002.56 **STATIONARY BEAM RADIATION THERAPY.** Stationary beam radiation therapy is radiation therapy without displacement of one or more mechanical axes relative to the patient during irradiation.

002.57 **STRAIGHT RADIATION.** Stray radiation equals the sum of leakage and scattered radiation.

002.58 **TARGET.** Target is that part of an x-ray tube or accelerator onto which a beam of accelerated particles is directed to produce ionizing radiation or other particles.

002.59 **TARGET SKIN DISTANCE (TSD).** Target skin distance (TSD) is the distance measured along the beam axis from the center of the front surface of the x-ray target or electron virtual source to the surface of the irradiated object or patient.
002.60  TENTH VALUE LAYER (TVL). Tenth value layer (TVL) is the thickness of a specified material which attenuates x-radiation or gamma radiation to an extent such that the air kerma rate, exposure rate, or absorbed dose rate is reduced to one tenth of the value measured without the material at the same point.

002.61  TERMINATION OF IRRADIATION. Termination of irradiation is the stopping of irradiation in a fashion which will not permit continuance of irradiation without the resetting of operating conditions at the control panel.

002.62  THERAPEUTIC RADIATION MACHINE. Therapeutic radiation machine is an x-ray or electron producing equipment designed and used for external beam radiation therapy. For the purpose of these regulations, therapeutic radiation machine includes, but is not limited to devices used to administer electronic brachytherapy.

002.63  TUBE. Tube is an x-ray tube, unless otherwise specified.

002.64  TUBE HOUSING ASSEMBLY. Tube housing assembly is the tube housing with tube installed. It includes high voltage or filament transformers and other appropriate elements when such are contained within the tube housing.

002.65  USEFUL BEAM. Useful beam is radiation emanating from the tube housing port or the radiation head and passing through the aperture of the beam limiting device when the exposure controls are in a mode to cause the therapeutic radiation machine to produce radiation.

002.66  VIRTUAL SIMULATOR. Virtual simulator is a computed tomography (CT) unit used in conjunction with relevant software which recreates the treatment machine; and that allows import, manipulation, display, and storage of images from computed tomography (CT) or other imaging modalities.

002.67  VIRTUAL SOURCE. Virtual source is a point from which radiation appears to originate.

002.68  WEDGE FILTER. Wedge filter is a filter which effects continuous change in transmission over all or a part of the useful beam.

002.69  WRITTEN DIRECTIVE. Written directive is an order in writing for the administration of radiation to a specific patient or human research subject, as specified in 180 NAC 20-005.01(A).

002.70  X-RAY TUBE. X-ray tube is any electron tube which is designed to be used primarily for the production of x-rays.

003.  GENERAL ADMINISTRATIVE REQUIREMENTS FOR REGISTRANTS OF THERAPEUTIC RADIATION MACHINES. The general administrative requirements for registrants using therapeutic radiation machines are as follows.
003.01 ADMINISTRATIVE CONTROLS. The registrant must be responsible for directing the operation of the therapeutic radiation machines that have been registered with the Department. The registrant or the registrant's agent must ensure that the requirements of this chapter are met in the operation of the therapeutic radiation machine or machines.

003.02 PROHIBITION. A therapeutic radiation machine that does not meet the provisions of these regulations cannot be used for irradiation of patients.

003.03 TRAINING FOR EXTERNAL BEAM RADIATION THERAPY USERS. The registrant of any therapeutic radiation machine subject to 180 NAC 20-006 or 20-007 must require the user to be a physician who is licensed in the State of Nebraska and who:

(1) Is certified in:
   (i) Radiology, combined diagnostic and therapeutic radiology program, therapeutic radiology or radiation oncology, by the American Board of Radiology;
   (ii) Radiation oncology by the American Osteopathic Board of Radiology;
   (iii) Radiology, with specialization in radiotherapy, as a British "Fellow of the Faculty of Radiology" or "Fellow of the Royal College of Radiology"; or
   (iv) Therapeutic radiology by the Canadian Royal College of Physicians and Surgeons; or

(2) Is in the active practice of therapeutic radiology, and has completed 200 hours of instruction in basic radiation techniques applicable to the use of an external beam radiation therapy unit, 500 hours of supervised work experience, and a minimum of three years of supervised clinical experience.
   (i) To satisfy the requirement for instruction, the classroom and laboratory training must include:
      (a) Radiation physics and instrumentation;
      (b) Radiation protection;
      (c) Mathematics pertaining to the use and measurement of ionization radiation; and
      (d) Radiation biology; and
   (ii) To satisfy the requirement for supervised work experience, training must be under the supervision of an individual meeting the requirements of 180 NAC 20-003.03 and must include:
      (a) Review of the full calibration measurements and periodic quality assurance checks;
      (b) Evaluation of prepared treatment plans and calculation of treatment times and patient treatment settings;
      (c) Using administrative controls to prevent misadministrations; and
      (d) Implementing emergency procedures to be followed in the event of the abnormal operation of an external beam radiation therapy unit or console; and
   (iii) To satisfy the requirement for a period of supervised clinical experience, training must include one year in a formal training program approved by the Residency Review Committee for Radiology of the Accreditation Council for Graduate Medical Education or the Committee on Postdoctoral Training of the American Osteopathic Association and an additional two years of clinical experience in therapeutic radiology under the supervision of an individual meeting the requirements of 180 NAC 20-003.03. The supervised clinical experience must include:
(a) Examining individuals and reviewing their case histories to determine their suitability for external beam radiation therapy treatment, and any limitations or contraindications;
(b) Selecting proper dose and how it is to be administered;
(c) Calculating the external beam radiation therapy doses and collaborating with the user in the review of patients' progress and consideration of the need to modify originally prescribed doses or treatment plans, or both as warranted by patients' reaction to radiation; and
(d) Post administration follow up and review of case histories.

003.03(A) TRAINING DOCUMENTATION. The names and training of all current users of therapeutic radiation machines must be kept on file at the facility. Information on former users of therapeutic radiation machines must be maintained for at least two years beyond the last date they were authorized as a user of the therapeutic radiation machine at that facility.

003.04 TRAINING FOR RADIOLOGICAL MEDICAL PHYSICIST. The registrant for any therapeutic radiation machine subject to 20 NAC 20-006 or 20-007 must require the individual to meet the training requirements of 180 NAC 15-004.01.

003.05 QUALIFICATIONS OF OPERATORS. Individuals who will be operating therapeutic radiation machines for medical use must meet the following requirements.

(A) Individuals who will be operating a therapeutic radiation machine for medical use must be American Registry of Radiologic Technologists (ARRT) Registered Radiation Therapy Technologists. Individuals who are not American Registry of Radiologic Technologists (ARRT) Registered Radiation Therapy Technologists must submit evidence that they have satisfactorily completed a radiation therapy technologist training program that complies with the requirements of the Joint Review Committee on Education in Radiologic Technology, "Standards for Accredited Educational Program in Radiologic Sciences – Effective January 1, 2002", Joint Review Committee on Education in Radiologic Technology, January 1996; Revised 2001. This document is available for viewing at the Department of Health and Human Services, Division of Public Health, Office of Radiological Health, 301 Centennial Mall South, 3rd Floor, Lincoln, Nebraska 68509.

(B) The names and training of all personnel currently operating a therapeutic radiation machine must be kept on file at the facility. Information on former operators must be retained for at least two (2) years beyond the last date they were authorized to operate a therapeutic radiation machine at that facility.

003.06 WRITTEN SAFETY PROCEDURES. Written safety procedures and rules must be developed by a radiological medical physicist and must be available in the control area of a therapeutic radiation machine. The written safety procedures must include any restrictions required for the safe operation of the each therapeutic radiation machine. The operator must be able to demonstrate familiarity with these procedures.

003.07 LIMITATIONS. Individuals must not be exposed to the useful beam except for medical therapy purposes and unless the exposure has been ordered in writing by a licensed
practitioner of the healing arts. This provision prohibits deliberate exposure of an individual for training, demonstration, or other non-healing arts purposes.

003.08 COMPLIANCE WITH QUALITY MANAGEMENT PROGRAM. All individuals associated with the operation of a therapeutic radiation machine must be instructed in and comply with the provisions of the registrant's quality management program. In addition to the requirements of this chapter, these individuals are also subject to the requirements of 180 NAC 4-005, 4-009 and 4-021.

003.09 INFORMATION AND MAINTENANCE RECORD AND ASSOCIATED INFORMATION. The registrant must maintain the following information in a separate file or package for each therapeutic radiation machine for inspection by the Department:
(A) A report of acceptance testing;
(B) The records of all surveys, calibrations, and periodic quality assurance checks of the therapeutic radiation machine required by this chapter, to include the name or names of the individual or individuals who performed the activities;
(C) The records of maintenance or modifications, or both, performed on the therapeutic radiation machine, to include the name or names of the individual or individuals who performed the services; and
(D) A signature of person authorizing the return of therapeutic radiation machine to clinical use after service, repair, or upgrade.

003.10 RECORDS RETENTION. All records required by this chapter must be retained until disposal is authorized by the Department, unless another retention period is authorized in this chapter. All required records must be retained in an active file from the time of generation, until at least the next Department inspection. Any required record generated prior to the last Department inspection may be microfilmed or otherwise archived as long as a complete copy of the record can be retrieved until the Department authorizes final disposal.

004. GENERAL TECHNICAL REQUIREMENTS FOR REGISTRANTS OF THERAPEUTIC RADIATION MACHINES. The general technical requirements for registrants using therapeutic radiation machines are as follows.

004.01 PROTECTION SURVEYS. Radiation protection surveys must be conducted as follows.

004.01(A) RADIATION SURVEY INSTRUMENT. The registrant must ensure that radiation protection surveys of all new facilities, and existing facilities not previously surveyed, are performed with an operable radiation measurement survey instrument calibrated according to 180 NAC 20-008. The radiation protection survey must be performed by, or under the direction of, a radiological medical physicist. The radiological medical physicist must verify, with the therapeutic radiation machine in a "BEAM-ON" condition, with the largest clinically available treatment field and with and without a scattering phantom in the useful beam of radiation:
(i) Radiation levels in restricted areas are not likely to cause personnel exposures in excess of the limits specified in 180 NAC 4-005.01; and
(ii) Radiation levels in unrestricted areas do not exceed the limits specified in 180 NAC 4-013.01 and 180 NAC 4-013.02.
004.01(B) ADDITIONAL REQUIREMENTS. In addition to the requirements of 180 NAC 20-004.01(A), a radiation protection survey must also be performed prior to any subsequent medical use and:
(i) After making any change in the treatment room shielding;
(ii) After making any change in the location of the therapeutic radiation machine within the treatment room;
(iii) After relocating the therapeutic radiation machine; or
(iv) Before using the therapeutic radiation machine in a manner that could result in increased radiation levels in areas outside the external beam radiation therapy treatment room.

004.01(C) VIOLATIONS. The survey record must indicate all instances where the registrant, in the opinion of the radiological medical physicist, is in violation of applicable regulations. The survey record must also include:
(i) The date of the measurements;
(ii) The reason the survey is required;
(iii) The manufacturer's name;
(iv) The model number and serial number of the therapeutic radiation machine;
(v) The instrument or instruments used to measure radiation levels;
(vi) A plan of the areas surrounding the treatment room that were surveyed;
(vii) The measured dose rate at several points in each area expressed in microsieverts or millirems per hour;
(viii) The calculated maximum level of radiation over a period of one week for each restricted and unrestricted area; and
(ix) The signature of the individual responsible for conducting the survey.

004.01(D) PROHIBITION OF USE. If the results of the surveys required by 180 NAC 20-004.01(A) or (B) indicate any radiation levels in excess of the respective limit specified in 180 NAC 20-004.01(A), the registrant must lock the control in the "OFF" position and not use the unit:
(i) Except as may be necessary to repair, replace, or test the therapeutic radiation machine, the therapeutic radiation machine shielding, or the treatment room shielding; or
(ii) Until the registrant has received a specific exemption from the Department.

004.02 MODIFICATION OF RADIATION THERAPY UNIT OR ROOM BEFORE BEGINNING A TREATMENT PROGRAM. If the survey required by 180 NAC 20-004.01 indicates that an individual in an unrestricted area may be exposed to levels of radiation greater than those permitted by 180 NAC 4-013.01 and 4-013.02, before beginning the treatment program the registrant must:
(A) Either equip the unit with beam direction interlocks or add additional radiation shielding to ensure compliance with 180 NAC 4-013.01 and 4-013.02;
(B) Perform the survey required by 180 NAC 20-004.01 again; and
(C) Include in the report required by 180 NAC 20-004.04 the results of the initial survey, a description of the modification made to comply with 180 NAC 20-004.01(A), and the results of the second survey; or
(D) Request and receive a registration amendment under 180 NAC 4-013.04 that authorizes radiation levels in unrestricted areas greater than those permitted by 180 NAC 4-013.01 and 4-013.02.

004.03 DOSIMETRY EQUIPMENT. The requirements for use of dosimetry equipment are as follows.

004.03(A) CALIBRATED DOSIMETRY SYSTEM. The registrant must have a calibrated dosimetry system available for use. The system must have been calibrated by the National Institute for Standards and Technology (NIST) or by an American Association of Physicists in Medicine (AAPM) Accredited Dosimetry Calibration Laboratory (ADCL). The calibration must have been performed within the previous 24 months and after any servicing that may have affected system calibration. The dosimetry system must have been calibrated at an energy or energy range appropriate for the radiation being measured.

004.03(B) QUALITY ASSURANCE CHECK MEASUREMENTS. The registrant must have available for use a dosimetry system for quality assurance check measurements. To meet this requirement, the system may be compared with a system that has been calibrated according to 180 NAC 20-004.03(A). This comparison must have been performed within the previous 12 months and after any servicing that may have affected the system calibration. The quality assurance check system may be the same system used to meet the requirement in 180 NAC 20-004.03(A).

004.03(C) RECORDS. The registrant must maintain a record of each dosimetry system calibration, intercomparison, and comparison for the duration of the registration. For each calibration, intercomparison, or comparison, the record must include:

(i) The date;
(ii) The model numbers and serial numbers of the instruments that were calibrated, inter-compared, or compared as required by 180 NAC 20-004.03(A) and (B); the correction factors that were determined;
(iii) The names of the individuals who performed the calibration, intercomparison, or comparison; and
(iv) Evidence that the intercomparison was performed by, or under the direct supervision and in the physical presence of, a radiological medical physicist.

005. QUALITY MANAGEMENT PROGRAM. Each registrant or applicant subject to 180 NAC 20-006 or 20-007 must develop, implement, and maintain a quality management program to ensure that radiation will be administered as directed by the user.

005.01 SCOPE AND APPLICABILITY. The quality management program must address, as a minimum, the following specific objectives.

005.01(A) WRITTEN DIRECTIVES. A written directive must:

(i) Be dated and signed by a user prior to the administration of radiation. If, because of the patient’s condition, a delay in the order to provide a written revision to an existing written directive would jeopardize the patient's health, an oral revision to an existing written directive will be acceptable, provided that the oral revision is
documented as soon as possible in writing in the patient’s record and a revised
written directive is signed by a user within 48 hours of the oral revision;
(ii) Contain the patient or human research subject’s name, the type and energy of the
beam, the total dose, dose per fraction, treatment site, and the number of fractions;
(iii) Be dated and signed by a user prior to the administration of the external beam
dose or the next fractional dose for a written revision to an existing written directive; and
(iv) Be retained by the registrant for three years.

005.01(B) PROCEDURES FOR ADMINISTRATION. The registrant must develop,
implement, and maintain written procedures to ensure that:
(i) Prior to the administration of each course of radiation treatments, the patient’s or
human research subject’s identity is verified, by more than one method, as the
individual named in the written directive;
(ii) Each administration is in according to the written directive;
(iii) Therapeutic radiation machine final plans of treatment and related calculations are
in according to the respective written directives by:
   (1) Checking both manual and computer generated dose calculations to verify they
       are correct and in according to the written directive; and
   (2) Verifying that any computer-generated calculations are correctly transferred
       into the consoles of authorized therapeutic medical units;
(iv) Any unintended deviation from the written directive is identified and evaluated, and
    appropriate action is taken; and
(v) The registrant retains a copy of the procedures for administration for the duration
    of the registration.

005.02 REPORTS AND NOTIFICATIONS OF MISADMINISTRATIONS. The requirements
for reports and notifications of misadministrations are as follows.

005.02(A) EVENTS RESULTING FROM PATIENT OR HUMAN RESEARCH SUBJECT
INTERVENTION. A registrant must report any event resulting from intervention of a patient
or human research subject in which the administration of therapeutic radiation machine
radiation results, or will result in, unintended permanent functional damage to an organ or
a physiological system as determined by a physician.

005.02(B) OTHER EVENTS. Other than events that result from intervention by a patient
or human research subject, a registrant must report any event in which the administration
of an external beam radiation dose:
(i) Involves the wrong patient, wrong treatment modality, or wrong treatment site;
(ii) Causes the calculated weekly administered dose to differ from the weekly
prescribed dose by more than 30 percent; or
(iii) Causes the calculated total administered dose to differ from the total prescribed
dose by more than 20 percent.

005.02(C) TELEPHONE NOTIFICATION. The registrant must notify the Department by
telephone no later than the next business day after the discovery of a misadministration.
005.02(D) WRITTEN REPORT. The registrant must submit a written report to the Department within 30 days after the discovery of a misadministration. The written report must include:

(i) The registrant’s name;
(ii) The name of the prescribing physician;
(iii) A brief description of the event;
(iv) Why the event occurred;
(v) The effect, if any, on the individual or individuals who received the administration;
(vi) Actions, if any, that have been taken, or are planned to prevent recurrence; and
(vii) Certification that the registrant notified the individual, or the individual’s responsible relative or guardian, and if not, why notification was not made.

005.02(E) IDENTIFICATION OF THE INDIVIDUAL PROHIBITED IN REPORT. The report must not contain the individual’s name or any other information that could lead to the identification of the individual.

005.02(F) NOTIFICATION TO THE REFERRING PHYSICIAN AND INDIVIDUAL. The registrant must provide notification of the event to the referring physician and also notify the individual who is the subject of the misadministration no later than 24 hours after its discovery, unless the referring physician personally informs the registrant that they will inform the individual or, based on their medical judgment, telling the individual would be harmful. The registrant is not required to notify the individual without first consulting the referring physician. If the referring physician or the affected individual cannot be reached within 24 hours, the registrant must make the appropriate notifications as soon as possible after that. The registrant must not delay any appropriate medical care for the individual, including any necessary remedial care as a result of the misadministration, because of any delay in notification. To meet the requirements of 180 NAC 20-005.02(E), the notification of the individual who is the subject of the misadministration may be made to that individual’s responsible relative or guardian. If a verbal notification is made, the registrant must inform the individual, or responsible relative or guardian that a written description of the event can be obtained from the registrant upon request. The registrant must provide such a written description if requested.

005.02(G) RIGHTS AND DUTIES. Aside from the notification requirement, nothing in this subsection affects any rights or duties of registrants and physicians in relation to each other, to individuals affected by the misadministration, or to that individual’s responsible relative or guardians.

005.02(H) COPY OF RECORD TO REFERRING PHYSICIAN. The registrant must retain a record of misadministration according to 180 NAC 20-005.03. A copy of the required record must be provided to the referring physician, if other than the registrant, within 15 days after discovery of the misadministration.

005.03 RECORDS OF MISADMINISTRATIONS. A registrant must retain a record of misadministration reported according to 180 NAC 20-005.02 for three years.

006. THERAPEUTIC RADIATION MACHINES OF LESS THAN 500 KILOVOLTS (kV). The requirements for therapeutic radiation machines of less than 500 kilovolts (kV) are as follows.
006.01 LEAKAGE RADIATION. When the x-ray tube is operated at its maximum rated tube current for the maximum kilovolts (kV), the leakage air kerma rate must not exceed the value specified at the distance specified for that classification of therapeutic radiation machine as follows.

006.01(A) 0-50 KILOVOLT (KV) SYSTEMS. The leakage air kerma rate measured at any position 5 centimeters from the tube housing assembly must not exceed 1 milligray (mGy) or 100 millirad (mrad) in any one hour.

006.01(B) GREATER THAN 50 AND LESS THAN 500 KILOVOLT (KV) SYSTEMS. The leakage air kerma rate measured at a distance of 1 meter from the target in any direction must not exceed 1 centigray (cGy) or 1 rad in any 1 hour. This air kerma rate measurement may be averaged over areas no larger than 100 square centimeters. In addition, the air kerma rate at a distance of 5 centimeters from the surface of the tube housing assembly must not exceed 30 centigray (cGy) or 30 rad per hour.

006.01(C) DETERMINATION OF LEAKAGE RADIATION. For each therapeutic radiation machine, the registrant must determine, or obtain from the manufacturer, the leakage radiation existing at the positions specified in 180 NAC 20-006.01(A) and 20-006.01(B) for the specified operating conditions. Records on leakage radiation measurements must be maintained at the installation for inspection by the Department.

006.02 PERMANENT BEAM LIMITING DEVICES. Permanent diaphragms or cones used for limiting the useful beam must provide at least the same degree of attenuation as required for the tube housing assembly.

006.03 ADJUSTABLE OR REMOVABLE BEAM LIMITING DEVICES. Adjustable or removable beam limiting devices must:

(A) Not transmit more than 5 percent of the useful beam for the most penetrating beam used for all adjustable or removable beam limiting devices, diaphragms, cones or blocks; and

(B) Indicate the position and shape of the radiation field by a light beam when adjustable beam limiting devices are used.

006.04 FILTER SYSTEM. The filter system must be marked so that:

(A) Filters cannot be accidentally displaced at any possible tube orientation;

(B) An interlock system prevents irradiation if the proper filter is not in place for equipment installed after July 11, 2009;

(C) The air kerma rate escaping from the filter slot does not exceed 1 centigray (cGy) or 1 rad per hour at 1 meter under any operating conditions; and

(D) Each filter is marked regarding its material of construction and its thickness.

006.05 TUBE IMMOBILIZATION. The x-ray tube and housing must be immobilized as follows.

006.05(A) X-RAY TUBE. The x-ray tube must be mounted so it cannot accidentally turn or slide with respect to the housing aperture.
006.05(B) TUBE HOUSING ASSEMBLY. The tube housing assembly must be capable of being immobilized for stationary portal treatments.

006.06 SOURCE MARKING. The tube housing assembly must be marked so that it is possible to determine the location of the source to within 5 millimeters. That marking must be readily accessible for use during calibration procedures.

006.07 BEAM BLOCK. Contact therapy tube housing assemblies must have a removable shield of material, equivalent in attenuation to 0.5 millimeters of lead at 100 kilovolts (kV) that can be positioned over the entire useful beam exit port during periods when the beam is not in use.

006.08 TIMER. A suitable irradiation control device must be provided to terminate the irradiation after a pre-set time interval. The timer must:
   (A) Be provided at the treatment control panel. The timer must have a display and must have a preset time selector and an elapsed time or time remaining indicator;
   (B) Be a cumulative timer that activates with an indication of "BEAM-ON" and retains its reading after irradiation is interrupted or terminated. After irradiation is terminated and before irradiation can be reinitiated, it must be necessary to reset the elapsed time indicator;
   (C) The timer must terminate irradiation when a pre-selected time has elapsed, if any dose monitoring system present has not previously terminated irradiation;
   (D) Permit accurate pre-setting and determination of exposure times as short as 1 second;
   (E) Not permit an exposure if the timer is set at zero;
   (F) Not activate until the shutter is opened when irradiation is controlled by a shutter mechanism unless calibration includes a timer error correction to compensate for mechanical lag; and
   (G) Be accurate to within 1 percent of the selected value or 1 second, whichever is greater.

006.09 CONTROL PANEL FUNCTIONS. The control panel, in addition to the displays required by other provisions in 180 NAC 20-006, must have:
   (A) An indication of whether electrical power is available at the control panel and if activation of the x-ray tube is possible;
   (B) An indication of whether x-rays are being produced;
   (C) A means for indicating x-ray tube potential and current;
   (D) A means for terminating an exposure at any time;
   (E) A locking device which will prevent unauthorized use of the therapeutic radiation machine; and
   (F) A positive display of specific filter or filters in the beam for therapeutic radiation machines installed after July 11, 2009.

006.10 MULTIPLE TUBES. When a control panel can energize more than one x-ray tube:
   (A) It must be possible to activate only one x-ray tube at any time;
   (B) There must be an indication at the control panel identifying which x-ray tube is activated; and
(C) There must be an indication at the tube housing assembly when that tube is energized.

006.11 TARGET TO SKIN DISTANCE (TSD). There must be a means of determining the central axis target to skin distance (TSD) to within 1 centimeter and to reproduce this measurement to within 2 millimeters from then on.

006.12 SHUTTERS. Unless it is possible to bring the x-ray output to the prescribed exposure parameters within 5 seconds after the x-ray "ON" switch is energized, the beam must be attenuated by a shutter having a lead equivalency not less than that of the tube housing assembly. In addition, after the unit is at operating parameters, the shutter must be controlled by the operator from the control panel. An indication of shutter position must appear at the control panel.

006.13 LOW FILTRATION X-RAY TUBES. Each therapeutic radiation machine equipped with a beryllium or other low filtration window must have the tube housing assembly clearly labeled and must be provided with a permanent warning device on the control panel that is activated when no additional filtration is present, to indicate that the dose rate is very high.

006.14 FACILITY DESIGN REQUIREMENTS FOR THERAPEUTIC RADIATION MACHINES CAPABLE OF OPERATING IN THE RANGE 50 KILOVOLTS (KV) TO 500 KILOVOLTS (KV). In addition to shielding adequate to meet requirements of 180 NAC 20-009, the treatment room must meet the following design requirements.

006.14(A) AURAL COMMUNICATION. Provision must be made for continuous two way aural communication between the patient and the operator at the control panel.

006.14(B) VIEWING SYSTEMS. Provision must be made to permit continuous observation of the patient during irradiation. The viewing system must be located so the operator can observe the patient from the control panel. The therapeutic radiation machine must not be used for patient irradiation unless at least one viewing system is operational.

006.15 ADDITIONAL REQUIREMENTS. Treatment rooms that contain a therapeutic radiation machine capable of operating above 150 kilovolts (kV) must meet the following additional requirements.

006.15(A) FIXED PROTECTIVE BARRIERS. All protective barriers must be fixed except for entrance doors or beam interceptors.

006.15(B) CONTROL PANEL. The control panel must be located outside the treatment room or in a totally enclosed booth, which has a ceiling, inside the room.

006.15(C) INTERLOCKS. Interlocks must be provided so all entrance doors, including doors to any interior booths, must be closed before treatment can be initiated or continued. If the radiation beam is interrupted by any door opening, it must not be possible to restore the machine to operation without closing the door and reinitiating irradiation by manual action at the control panel.
006.15(D) DOOR. When any door referred to in 180 NAC 20-006.15, item (C) is opened while the x-ray tube is activated, the air kerma rate at a distance of 1 meter from the source must be reduced to less than 1 milligray (mGy) or 100 millirad (mrad) per hour.

006.16 FULL CALIBRATION MEASUREMENTS. Full calibration measurements must be performed as follows.

006.16(A) FREQUENCY. Full calibration of a therapeutic radiation machine subject to 180 NAC 20-006 must be performed by, or under the direct supervision of, a radiological medical physicist:

(i) Before the first medical use following installation or reinstallation of the therapeutic radiation machine;

(ii) At intervals not exceeding one year; and

(iii) Before medical use under the following conditions:

(1) Whenever quality assurance check measurements indicate that the radiation output differs by more than 5 percent from the value obtained at the last full calibration and the difference cannot be reconciled; and

(2) Following any component replacement, major repair, or modification of components that could significantly affect the characteristics of the radiation beam.

(iv) Apart from the requirements of 180 NAC 20-006.16(A)(iii):

(1) Full calibration of therapeutic radiation machines with multi energy capabilities is required only for those modes or energies that are not within their acceptable range; and

(2) If the repair, replacement or modification does not affect all energies, full calibration must be performed on the affected energy that is in most frequent clinical use at the facility. The remaining energies may be validated with quality assurance check procedures compared to the criteria in 180 NAC 20-006.16(A), item (iii)(1).

006.16(B) REQUIRED MEASUREMENTS. To satisfy the requirement of 180 NAC 20-006.16(A), full calibration must include all measurements recommended for annual calibration by the National Council on Radiation Protection and Measurements (NCRP) Report 69, “Dosimetry of X-ray and Gamma Ray Beams for Radiation Therapy in the Energy Range 10 keV to 50 MeV” (1981). This document is available for viewing at the Department of Health and Human Services, Division of Public Health, Office of Radiological Health, 301 Centennial Mall South, 3rd Floor, Lincoln, Nebraska 68509.

006.16(C) CALIBRATION RECORDS. The registrant must maintain a record of each calibration for the duration of the registration. The record must include:

(i) The date of the calibration;

(ii) The manufacturer's name;

(iii) Model number and serial number for the therapeutic radiation machine if applicable;

(iv) The model numbers and serial numbers of the instruments used to calibrate the therapeutic radiation machine; and

(v) The signature of the radiological medical physicist responsible for performing the calibration.
006.17 PERIODIC QUALITY ASSURANCE CHECKS. The requirements for periodic quality assurance checks are as follows.

006.17(A) APPLICABILITY. Periodic quality assurance checks must be performed on therapeutic radiation machines subject to 180 NAC 20-006 that are capable of operation at greater than or equal to 50 kilovolts (kV).

006.17(B) QUALITY ASSURANCE CHECK CONTENT. To satisfy the requirements of 180 NAC 20-006.17(A), quality assurance checks must meet the following requirements:
   (i) Quality assurance checks must be performed according to written procedures established by the radiological medical physicist; and
   (ii) The quality assurance check procedures must specify the frequency the tests or measurements are to be performed. The quality assurance check procedures must specify that the quality assurance check will be performed during the calibration specified in 180 NAC 20-006.16(A). The acceptable tolerance for each parameter measured in the quality assurance check, when compared to the value for that parameter established in the calibration specified in 180 NAC 20-006.16(A), must be specified.

006.17(C) PARAMETER EXCEEDING TOLERANCE. The cause for a parameter exceeding a tolerance set by the radiological medical physicist must be investigated and corrected before the system is used for patient irradiation.

006.17(D) CHANGE IN OPERATING CHARACTERISTICS. Whenever a quality assurance check indicates a significant change in the operating characteristics of a system, as specified in the radiological medical physicist's quality assurance check procedures, the system must be recalibrated as required in 180 NAC 20-006.16(A).

006.17(E) DOSIMETRY SYSTEM. The registrant must use the dosimetry system described in 180 NAC 20-004.03(B), to make the quality assurance check required in 180 NAC 20-006.17(B).

006.17(F) RADIOLOGICAL MEDICAL PHYSICIST REVIEW. The registrant must have the radiological medical physicist review and sign the results of each radiation output quality assurance check within 1 month of the date that the check was performed.

006.17(G) REQUIRED INTERVALS. The registrant must ensure that quality assurance checks of therapeutic radiation machines subject to 180 NAC 20-006 are performed at intervals not to exceed 1 month.

006.17(H) PROHIBITION. Despite the requirements of 180 NAC 20-006.17(F) and (G), the registrant must ensure that no therapeutic radiation machine is used to administer radiation to humans unless the quality assurance checks required by 180 NAC 20-006.17(F) and (G) have been performed within the 30 day period immediately prior to that administration.

006.17(I) PROPER OPERATION. To satisfy the requirement of 180 NAC 20-006.17(B), safety quality assurance checks must ensure proper operation of:
(i) Electrical interlocks at each external beam radiation therapy room entrance;
(ii) The "BEAM-ON" and termination switches;
(iii) Beam condition indicator lights on the access door or doors, control console, and in the radiation therapy room;
(iv) Viewing systems; and
(v) If applicable, electrically operated treatment room doors from inside and outside the treatment room.

006.17(J) RECORD MAINTENANCE. The registrant must maintain a record of each quality assurance check required by 180 NAC 20-006.17(A) and (G) for three years. The record must include:
(i) The date of the quality assurance check;
(ii) The manufacturer's name, model number, and serial number of the therapeutic radiation machine;
(iii) The manufacture’s name, model number and serial number for the instrument or instruments used to measure the radiation output of the therapeutic radiation machine; and
(iv) The signature of the individual who performed the periodic quality assurance check.

006.18 OPERATING PROCEDURES. The requirements for operation procedures are as follows.

006.18(A) PROHIBITION. The therapeutic radiation machine must not be used for irradiation of patients unless the requirements of 180 NAC 20-006.16 and 20-006.17 have been met.

006.18(B) SECURED IF UNATTENDED. Therapeutic radiation machines must not be left unattended unless secured according to 180 NAC 20-006.09, item (E).

006.18(C) MECHANICAL SUPPORT OR RESTRAINT. When a patient must be held in position for radiation therapy, mechanical supporting or restraining devices must be used.

006.18(D) HOLDER PROTECTION. The tube housing assembly must not be held by an individual during operation unless the assembly is designed to require such holding and the peak tube potential of the system does not exceed 50 kilovolts (kV). In those cases, the holder must wear protective gloves and a protective apron of not less than 0.5 millimeters lead equivalency at 100 kilovolts (kV).

006.18(E) OPERATING AND EMERGENCY PROCEDURES. A copy of the current operating and emergency procedures must be maintained at the therapeutic radiation machine control console.

006.18(F) INDIVIDUALS OTHER THAN THE PATIENT. No individual other than the patient must be in the treatment room during exposures from therapeutic radiation machines operating above 150 kilovolts (kV). At energies less than or equal to 150 kilovolts (kV), any individual, other than the patient, in the treatment room must be protected by a barrier sufficient to meet the requirements of 180 NAC 4-005.
007. THERAPEUTIC RADIATION MACHINES – PHOTON THERAPY SYSTEMS 500 KILOVOLTS (KV) AND ABOVE AND ELECTRON THERAPY SYSTEMS 500 KILOELECTRON VOLTS (keV) and ABOVE. The requirements for therapeutic radiation machines – photon therapy systems kilovolts (kV) and above and electron therapy systems 500 kiloelectron volts (keV) and above are as follows.

007.01 LEAKAGE RADIATION OUTSIDE THE MAXIMUM USEFUL BEAM IN PHOTON AND ELECTRON MODES. The requirements governing leakage radiation outside the maximum useful beam in photon and electron modes are as follows.

007.01(A) ABSORBED DOSE. The absorbed dose due to leakage radiation, excluding neutrons, at any point outside the maximum sized useful beam, but within a circular plane of radius 2 meters which is perpendicular to and centered on the central axis of the useful beam at the nominal treatment distance, must not exceed a maximum of 0.2 percent and an average of 0.1 percent of the absorbed dose on the central axis of the beam at the nominal treatment distance. Measurements must be averaged over an area not exceeding 100 square centimeters at a minimum of 16 points uniformly distributed in the plane.

007.01(B) LEAKAGE RADIATION. Except for the area defined in 180 NAC 20-007.01(A), the absorbed dose due to leakage radiation, excluding neutrons, at 1 meter from the electron path between the electron source and the target or electron window must not exceed 0.5 percent of the absorbed dose on the central axis of the beam at the nominal treatment distance. Measurements must be averaged over an area not exceeding 100 square centimeters.

007.01(C) NEUTRON ABSORBED DOSE. For equipment manufactured after July 11, 2009, the neutron absorbed dose outside the useful beam must not exceed manufacturer’s specifications.

007.01(D) LEAKAGE RADIATION MEASUREMENTS. For each therapeutic radiation machine, the registrant must determine, or obtain from the manufacturer, the leakage radiation existing at the positions specified in 180 NAC 20-007.01(A) through (C) for the specified operating conditions. Records on leakage radiation measurements must be maintained for inspection by the Department.

007.02 LEAKAGE RADIATION THROUGH BEAM LIMITING DEVICES. The requirements governing leakage radiation through beam limiting devices are as follows.

007.02(A) PHOTON RADIATION. All adjustable or interchangeable beam limiting devices must attenuate the useful beam so that at the nominal treatment distance, the maximum absorbed dose anywhere in the area shielded by the beam limiting device or devices must not exceed 2 percent of the maximum absorbed dose on the central axis of the useful beam measured in a 100 square centimeters radiation field, or maximum available field size if less than 100 square centimeters.

007.02(B) ELECTRON RADIATION. All adjustable or interchangeable electron applicators must attenuate the radiation, including, but not limited to, photon radiation generated by electrons incident on the beam limiting device and electron applicator and
other parts of the radiation head, so that the absorbed dose in a plane perpendicular to the central axis of the useful beam at the nominal treatment distance must not exceed:

(i) A maximum of 2 percent and average of 0.5 percent of the absorbed dose on the central axis of the useful beam at the nominal treatment distance. This limit must apply beyond a line 7 centimeters outside the periphery of the useful beam; and

(ii) A maximum of 10 percent of the absorbed dose on the central axis of the useful beam at the nominal treatment distance. This limit must apply beyond a line 2 centimeters outside the periphery of the useful beam.

007.03 MEASUREMENT OF LEAKAGE RADIATION. The requirements for measurement of leakage radiation are as follows.

007.03(A) PHOTON RADIATION. Measurements of leakage radiation through the beam limiting devices must be made with the beam limiting devices closed and any residual aperture blocked by a minimum of two tenth value layers of suitable absorbing material. In the case of overlapping beam limiting devices, the leakage radiation through each set must be measured independently at the depth of maximum dose. Measurements must be made, using a radiation detector, of an area not exceeding 10 square centimeters.

007.03(B) ELECTRON RADIATION. Measurements of leakage radiation through the electron applicators must be made with the electron beam directed into the air and using a radiation detector of an area up to, but not exceeding, one square centimeter suitably protected against radiation that has been scattered from material beyond the radiation detector. Measurements must be made using one centimeter of water equivalent build up material.

007.04 FILTERS AND WEDGES. The requirements governing filters and wedges are as follows.

007.04(A) WEDGE FILTERS. Each wedge filter that is removable from the system must be clearly marked with an identification number. For removable wedge filters, the nominal wedge angle must appear on the wedge, or wedge tray if permanently mounted to the tray. If the wedge or wedge tray is significantly damaged, the wedge transmission factor must be redetermined.

007.04(B) ABSORBED DOSE. If the absorbed dose rate information required by 180 NAC 20-007.01 relates exclusively to operation with a field flattening filter or beam scattering foil in place, such foil or filter must be removable only by the use of tools.

007.04(C) WEDGE FILTERS, INTERCHANGEABLE FIELD FLATTENING FILTERS, OR INTERCHANGEABLE BEAM SCATTERING FOILS. For equipment installed after July 11, 2009 that utilizes wedge filters, interchangeable field flattening filters, or interchangeable beam scattering foils, the following apply:

(i) Irradiation must not be possible until a selection of a filter, or a selection to use "no filter", has been made at the treatment control panel, either manually or automatically;

(ii) An interlock system must be provided to prevent irradiation if the filter selected is not in the correct position;
(iii) A display must be provided at the treatment control panel showing the wedge filter or filters, interchangeable field flattening filter or filters, or interchangeable beam scattering foil or foils in use; and
(iv) An interlock must be provided to prevent irradiation if any filter or beam scattering foil selection operation carried out in the treatment room does not agree with the filter or beam scattering foil selection operation carried out at the treatment control panel.

007.05 STRAY RADIATION IN THE USEFUL BEAM. For equipment installed after July 11, 2009, the registrant must determine during acceptance testing, or obtain from the manufacturer, data sufficient to ensure that x-ray stray radiation in the useful electron beam, absorbed dose at the surface during x-ray irradiation, and stray neutron radiation in the useful x-ray beam do not exceed manufacturer’s specifications.

007.06 BEAM MONITORS. All therapeutic radiation machines subject to 180 NAC 20-007 must be provided with redundant beam monitoring systems. The sensors for these systems must be fixed in the useful beam during treatment to indicate the dose monitor unit rate.

007.06(A) INDEPENDENTLY POWERED INTEGRATING DOSE METERS. Equipment installed after July 11, 2009 must be provided with at least two independently powered integrating dose meters. Alternatively, common elements may be used if the production of radiation is terminated upon failure of any common element.

007.06(B) RADIATION DETECTOR. Equipment installed on or before July 11, 2009, must be provided with at least one radiation detector. This detector must be incorporated into a useful beam monitoring system;

007.06(C) DETECTOR AND SYSTEM. The detector and the system into which that detector is incorporated must meet the following requirements:
   (i) Each detector must be removable only with tools and, if movable, must be interlocked to prevent incorrect positioning;
   (ii) Each detector must form part of a beam monitoring system from whose readings in dose monitor units the absorbed dose at a reference point can be calculated;
   (iii) Each beam monitoring system must be capable of independently monitoring, interrupting, and terminating irradiation;
   (iv) For equipment installed after July 11, 2009, the design of the beam monitoring systems must ensure that the:
       (1) Malfunctioning of one system must not affect the correct functioning of the other system or systems; and
       (2) Failure of either system must terminate irradiation or prevent the initiation of radiation; and
   (v) Each beam monitoring system must have a legible display at the treatment control panel. For equipment installed after July 11, 2009, each display must:
       (1) Maintain a reading until intentionally reset;
       (2) Have only one scale and no electrical or mechanical scale multiplying factors;
       (3) Utilize a design so that increasing dose is displayed by increasing numbers; and
(4) In the event of power failure, the beam monitoring information required in 180 NAC 20-007.06(C)(v)(3) displayed at the control panel at the time of failure must be retrievable in at least one system for a 20 minute period of time.

007.07  BEAM SYMMETRY. The requirements for beam symmetry are as follows.

007.07(A)  AUXILIARY DEVICES. Bent beam linear accelerators with a beam flattening filter or filters subject to 180 NAC 20-007 must be provided with an auxiliary device or devices to monitor beam symmetry.

007.07(B)  DETECTION OF ASYMMETRY. The device or devices referenced in 180 NAC 20-007.07(A) must be able to detect field asymmetry greater than 10 percent.

007.07(C)  TERMINATE IRRADIATION. The device or devices referenced in 180 NAC 20-007.07(A) must be configured to terminate irradiation if the specifications in 180 NAC 20-007.07(B) cannot be maintained.

007.08  SELECTION AND DISPLAY OF DOSE MONITOR UNITS. The requirements for selection and display of dose monitor units are as follows.

007.08(A)  SELECTION OF DOSE MONITORING UNITS. Irradiation must not be possible until a new selection of a number of dose monitor units has been made at the treatment control panel.

007.08(B)  DISPLAY OF DOSE MONITORING UNITS. The pre-selected number of dose monitor units must be displayed at the treatment control panel until reset manually for the next irradiation.

007.08(C)  RESETTING OF DOSIMETER DISPLAY. After termination of irradiation, it must be necessary to reset the dosimeter display before subsequent treatment can be initiated.

007.08(D)  RESET OF PRE-SELECTED DOSE MONITOR UNITS. For equipment installed after July 11, 2009 after termination of irradiation, it must be necessary for the operator to reset the pre-selected dose monitor units before irradiation can be initiated.

007.09  AIR KERMA RATE OR ABSORBED DOSE RATE. For equipment installed after July 11, 2009, a system must be provided from where the readings of the air kerma rate or absorbed dose rate at a reference point can be calculated. The radiation detectors specified in 180 NAC 20-007.06 may form part of this system. In addition:

(A) The dose monitor unit rate must be displayed at the treatment control panel;

(B) If the equipment can deliver, under any conditions, an air kerma rate or absorbed dose rate at the nominal treatment distance more than twice the maximum value specified by the manufacturer, a device must be provided which terminates irradiation when the air kerma rate or absorbed dose rate exceeds a value twice the specified maximum. A record of the dose rate that the irradiation will be terminated must be maintained by the registrant;
(C) If the equipment can deliver, under any fault condition or conditions, an air kerma rate or absorbed dose rate at the nominal treatment distance more than ten times the maximum value specified by the manufacturer, a device must be provided to prevent the air kerma rate or absorbed dose rate anywhere in the radiation field from exceeding twice the specified maximum value and to terminate irradiation if the excess absorbed dose at the nominal treatment distance exceeds 4 gray (Gy) or 400 rad; and

(D) For each therapeutic radiation machine the registrant must determine, or obtain from the manufacturer, the maximum value or values specified in 180 NAC 20-007.09(B) and (C), for the specified operating conditions. Records of these maximum value or values must be maintained for inspection by the Department.

007.10 TERMINATION OF IRRADIATION BY THE BEAM MONITORING SYSTEM OR SYSTEMS DURING STATIONARY BEAM RADIATION THERAPY. The requirements for termination of irradiation by the beam monitoring system or systems during stationary beam radiation therapy are as follows.

007.10(A) PRE-SELECTED NUMBER OF DOSE MONITOR UNITS. Each primary system must terminate irradiation when the pre-selected number of dose monitor units has been detected by the system.

007.10(B) SECONDARY DOSE MONITORING SYSTEM. If the original design of the equipment included a secondary dose monitoring system, that system must be capable of terminating irradiation when not more than 15 percent or 40 dose monitor units above the pre-selected number of dose monitor units set at the control panel has been detected by the secondary dose monitoring system.

007.10(C) CONTROL PANEL INDICATION. For equipment installed after July 11, 2009, an indicator on the control panel must show which monitoring system has terminated irradiation.

007.11 TERMINATION OF IRRADIATION. It must be possible to terminate irradiation and equipment movement or go from an interruption condition to termination condition at any time from the operator's position at the treatment control panel.

007.12 INTERRUPTION OF IRRADIATION. If a therapeutic radiation machine has an interrupt mode, it must be possible to interrupt irradiation and equipment movements at any time from the treatment control panel. Following an interruption it must be possible to restart irradiation by operator action without any reselection of operating conditions. If any change is made of a pre-selected value during an interruption, irradiation and equipment movements must be automatically terminated.

007.13 TIMER. A suitable irradiation control device must be provided to terminate the irradiation after a pre-set time interval as follows:

(A) A timer must be provided which has a display at the treatment control panel. The timer must have a pre-set time selector and an elapsed time indicator;

(B) The timer must be a cumulative timer that activates with an indication of "BEAM-ON" and retains its reading after irradiation is interrupted or terminated. After irradiation is
terminated and before irradiation can be reinitiated, it must be necessary to reset the elapsed time indicator; and
(C) The timer must terminate irradiation when a pre-selected time has elapsed, if the dose monitoring systems have not previously terminated irradiation.

007.14 SELECTION OF RADIATION TYPE. Equipment capable of both x-ray therapy and electron therapy must meet the following additional requirements:
(A) Irradiation must not be possible until a selection of radiation type, x-rays or electrons, has been made at the treatment control panel;
(B) The radiation type selected must be displayed at the treatment control panel before and during irradiation;
(C) An interlock system must be provided to ensure that the equipment can principally emit only the radiation type that has been selected;
(D) An interlock system must be provided to prevent irradiation with x-rays, except to obtain an image, when electron applicators are fitted;
(E) An interlock system must be provided to prevent irradiation with electrons when accessories specific for x-ray therapy are fitted; and
(F) An interlock system must be provided to prevent irradiation if any selected operations carried out in the treatment room do not agree with the selected operations carried out at the treatment control panel.

007.15 SELECTION OF ENERGY. Equipment capable of generating radiation beams of different energies must meet the following requirements:
(A) Irradiation must not be possible until a selection of energy has been made at the treatment control panel;
(B) The nominal energy value selected must be displayed at the treatment control panel until reset manually for the next irradiation. After termination of irradiation, it must be necessary to reset the nominal energy value selected before subsequent treatment can be initiated; and
(C) Irradiation must not be possible until the appropriate flattening filter or scattering foil for the selected energy is in its proper location.

007.16 SELECTION OF STATIONARY BEAM RADIATION THERAPY OR MOVING BEAM RADIATION THERAPY. Therapeutic radiation machines capable of both stationary beam radiation therapy and moving beam radiation therapy must meet the following requirements:
(A) Irradiation must not be possible until a selection of stationary beam radiation therapy or moving beam radiation therapy has been made at the treatment control panel;
(B) The mode of operation must be displayed at the treatment control panel;
(C) An interlock system must be provided to ensure that the equipment can operate only in the mode that has been selected;
(D) An interlock system must be provided to prevent irradiation if any selected parameter in the treatment room does not agree with the selected parameter at the treatment control panel;
(E) Moving beam radiation therapy must be controlled to obtain the selected relationships between incremental dose monitor units and incremental movement. For equipment installed after July 11, 2009:
(i) An interlock system must be provided to terminate irradiation if the number of dose monitor units delivered in any 10 degrees of rotation or 1 centimeter of linear motion differs by more than 20 percent from the selected value;

(ii) Where the angle terminates the irradiation in moving beam radiation therapy, the dose monitor units delivered must differ by less than 5 percent from the dose monitor unit value selected;

(iii) An interlock must be provided to prevent motion of more than 5 degrees or 1 centimeter beyond the selected limits during moving beam radiation therapy;

(iv) An interlock must be provided to require that a selection of direction be made at the treatment control panel in all units that are capable of both clockwise and counter clockwise moving beam radiation therapy; and

(v) Moving beam radiation therapy must be controlled with both primary position sensors and secondary position sensors to obtain the selected relationships between incremental dose monitor units and incremental movement;

(F) Where the beam monitor system terminates the irradiation in moving beam radiation therapy, the termination of irradiation must be as required by 180 NAC 20-007.10; and

(G) For equipment installed after July 11, 2009, an interlock system must be provided to terminate irradiation if movement:

(i) Occurs during stationary beam radiation therapy; or

(ii) Does not start or stop during moving beam radiation therapy unless such stoppage is a pre-planned function.

007.17 FACILITY DESIGN REQUIREMENTS FOR THERAPEUTIC RADIATION MACHINES OPERATING ABOVE 500 kilovolts (kV). In addition to shielding adequate to meet the requirements of 180 NAC 20-009, the facility design must include the following:

(A) All protective barriers must be fixed, except for access doors to the treatment room or movable beam interceptors;

(B) In addition to other requirements specified in this chapter, the control panel must also:

(i) Be located outside the treatment room;

(ii) Provide an indication of whether electrical power is available at the control panel and if activation of the radiation is possible;

(iii) Provide an indication of whether radiation is being produced; and

(iv) Include an access control or locking device that will prevent unauthorized use of the therapeutic radiation machine;

(C) Windows, mirrors, closed circuit television, or an equivalent viewing system must be provided to permit continuous observation of the patient following positioning and during irradiation and must be located so the operator may observe the patient from the treatment control panel. The therapeutic radiation machine must not be used for patient irradiation unless at least one viewing system is operational;

(D) Provision must be made for continuous two way aural communication between the patient and the operator at the control panel. The therapeutic radiation machine must not be used for irradiation of patients unless continuous two way aural communication is possible;

(E) Treatment room entrances must be provided with warning lights in a readily observable position near the outside of all access doors that indicate when the useful beam is "ON" and when it is "OFF";
(F) Interlocks must be provided so all access controls are activated before treatment can be initiated or continued. If the radiation beam is interrupted by any access control, it must not be possible to restore the machine to operation without resetting the access control and reinitiating irradiation by manual action at the control panel;

(G) If the shielding material in any protective barrier requires the presence of a beam interceptor to ensure compliance with 180 NAC 4-013.01 and 4-013.02, interlocks must be provided to prevent the production of radiation, unless the beam interceptor is in place, whenever the useful beam is directed at the designated barrier or barriers;

(H) At least one emergency power cutoff switch must be located in the radiation therapy room and must terminate all equipment electrical power including radiation and mechanical motion. This switch is in addition to the termination switch required by 180 NAC 20-007.11. All emergency power cutoff switches must include a manual reset so that the therapeutic radiation machine cannot be restarted from the unit's control console without resetting the emergency cutoff switch;

(I) All safety interlocks must be designed so that any defect or component failure in the safety interlock system prevents or terminates operation of the therapeutic radiation machine; and

(J) Surveys for residual activity must be conducted on all therapeutic radiation machines capable of generating photon and electron energies above 10 megavolts (MV) prior to machining, removing, or working on therapeutic radiation machine components which may have become activated due to photo-neutron production.

007.18 RADILOGICAL MEDICAL PHYSICIST SUPPORT. The requirements for radiological medical physicist support are as follows.

007.18(A) RADILOGICAL MEDICAL PHYSICIST RESPONSIBILITIES. The services of a radiological medical physicist is required for registrants using therapeutic radiation machines with energies of 500 kilovolts (kV) and above. The radiological medical physicist must be responsible for:

(i) Full calibration or calibrations required by 180 NAC 20-007.20 and protection surveys required by 180 NAC 20-004.01;

(ii) Supervision and review of dosimetry;

(iii) Beam data acquisition and transfer for computerized dosimetry, and supervision of its use;

(iv) Quality assurance, including quality assurance check review required by 180 NAC 20-007.21(E);

(v) Consultation with the user in treatment planning, as needed; and

(vi) Performing calculations and assessments regarding misadministrations.

007.18(B) RADILOGICAL MEDICAL PHYSICIST CONTACT INFORMATION. If the radiological medical physicist is not a full time employee of the registrant, the operating procedures required by 180 NAC 20-007.19(F) must also specifically address how the radiological medical physicist is to be contacted for problems or emergencies, as well as the specific actions, if any, to be taken until the radiological medical physicist can be contacted.

007.19 OPERATING PROCEDURES. The requirements for operating procedures are as follows.
007.19(A) **INDIVIDUALS OTHER THAN THE PATIENT.** No individual, other than the patient, must be in the treatment room during treatment or during any irradiation for testing or calibration purposes.

007.19(B) **PROHIBITION.** Therapeutic radiation machines must not be made available for medical use unless the requirements of 180 NAC 20-004.01, 20-007.20 and 20-007.21 have been met.

007.19(C) **SECURED IF UNATTENDED.** Therapeutic radiation machines, when not in operation, must be secured to prevent unauthorized use.

007.19(D) **RADIATION FIELD POSITION AND SHAPE INDICATION.** When adjustable beam limiting devices are used, the position and shape of the radiation field must be indicated by a light field.

007.19(E) **MECHANICAL SUPPORT OR RESTRAINING DEVICES.** If a patient must be held in position during treatment, mechanical supporting or restraining devices must be used; and

007.19(F) **OPERATING AND EMERGENCY PROCEDURES.** A copy of the current operating and emergency procedures must be maintained at the therapeutic radiation machine control console.

007.20 **ACCEPTANCE TESTING, COMMISSIONING, AND FULL CALIBRATION MEASUREMENTS.** The requirements for acceptance testing, commissioning, and full calibration measurements are as follows.

007.20(A) **PERFORMANCE.** Acceptance testing, commissioning, and full calibration of a therapeutic radiation machine subject to 180 NAC 20-007 must be performed by, or under the supervision of, a radiological medical physicist.

007.20(B) **INSTALLATION OR REINSTALLATION.** Acceptance testing and commissioning must be performed according to "AAPM Code of Practice for Radiotherapy Accelerators: Report of AAPM Radiation Therapy Task Group 45" and manufacturers specifications, and must be conducted before the first medical use following installation or reinstallation of the therapeutic radiation machine. This document is available for viewing at the Department of Health and Human Services, Division of Public Health, Office of Radiological Health, 301 Centennial Mall South, 3rd Floor, Lincoln, Nebraska 68509.

007.20(C) **FULL CALIBRATION MEASUREMENTS.** Full calibration must include measurement of all applicable parameters required by Table II of "Comprehensive QA for Radiation Oncology: Report of AAPM Radiation Therapy: AAPM Report No. 46," prepared by Committee Task Group 40 and must be performed according to "AAPM Code of Practice for Radiotherapy Accelerators: AAPM Report No. 47" prepared by Radiation Therapy Task Group 45. These documents are available for viewing at the Department of Health and Human Services, Division of Public Health, Office of Radiological Health, 301 Centennial Mall South, 3rd Floor, Lincoln, Nebraska 68509. Although it may not be necessary to complete all elements of a full calibration at the same time, all applicable
parameters, for all energies, must be completed at intervals not exceeding twelve calendar months, unless a more frequent interval is required in Table II.

007.20(D) FREQUENCY. The radiological medical physicist must perform all elements of a full calibration necessary to determine that all parameters are within acceptable limits:

(i) Whenever quality assurance check measurements indicate that the radiation output differs by more than 5 percent from the value obtained at the last full calibration and the difference cannot be reconciled. Therapeutic radiation machines with multi energy or multi-mode capabilities must require measurements for only those modes or energies that are not within their acceptable range; and

(ii) Following any component replacement, major repair, or modification of components that could significantly affect the characteristics of the radiation beam. If the repair, replacement or modification does not affect all modes or energies, measurements must be performed on the effected mode or energy that is in most frequent clinical use at the facility. The remaining energies or modes may be validated with quality assurance check procedures against the criteria in 180 NAC 20-007.20(D)(i).

007.20(E) CALIBRATION RECORDS. The registrant must maintain a record of each calibration in an auditable form for the duration of the registration. The record must include:

(i) The date of the calibration;

(ii) The manufacturer’s name, model number, and serial number for the therapeutic radiation machine;

(iii) The model numbers and serial numbers of the instruments used to calibrate the therapeutic radiation machine; and

(iv) The signature of the radiological medical physicist responsible for performing the calibration.

007.21 PERIODIC QUALITY ASSURANCE CHECKS. The requirements for periodic quality assurance checks are as follows.

007.21(A) PERIODIC QUALITY ASSURANCE CHECKS INTERVALS. Periodic quality assurance checks must be performed on all therapeutic radiation machines subject to 180 NAC 20-007 at intervals not to exceed those specified in "Comprehensive QA for Radiation Oncology: AAPM Report No. 46," prepared by AAPM Radiation Therapy Committee Task Group 40; or other procedure.

007.21(B) QUALITY ASSURANCE CHECK CONTENTS. To satisfy the requirement of 180 NAC 20-007.21(A), quality assurance checks must include a determination of central axis radiation output and a representative sampling of periodic quality assurance checks contained in "Comprehensive QA for Radiation Oncology: AAPM Report No. 46" prepared by Radiation Therapy Committee Task Group 40. The registrant may use an alternative protocol that has been submitted to the Department for approval. Representative sampling must include all applicable referenced periodic quality assurance checks in an interval not to exceed 12 consecutive calendar months.

007.21(C) DOSIMETRY SYSTEM. The registrant must use a dosimetry system that has been intercompared within the previous 12 months with the dosimetry systems described
in 180 NAC 20-004.03(A) to make the periodic quality assurance checks required in 180 NAC 20-007.21(B).

007.21(D)  **WRITTEN PROCEDURES.** The registrant must perform periodic quality assurance checks required by 180 NAC 20-007.21(A) according to written procedures established by the radiological medical physicist;

007.21(E)  **REVIEW OF RESULTS.** The registrant must review the results of each periodic radiation output check according to the following procedures:

(i) The user and radiological medical physicist must be immediately notified if any parameter is not within its acceptable tolerance. The therapeutic radiation machine must not be made available for subsequent medical use until the radiological medical physicist has determined that all parameters are within their acceptable tolerances;

(ii) If all quality assurance check parameters appear to be within their acceptable range, the quality assurance check must be reviewed and signed by either the user or radiological medical physicist within 3 treatment days; and

(iii) The radiological medical physicist must review and sign the results of each radiation output quality assurance check at intervals not to exceed one month.

007.21(F)  **QUALITY ASSURANCE CHECK FREQUENCY.** Therapeutic radiation machines subject to 180 NAC 20-007 must have applicable safety quality assurance checks listed in "Comprehensive QA for Radiation Oncology: AAPM Report No. 46" prepared by AAPM Radiation Therapy Committee Task Group 40. The registrant may use an alternative protocol that has been submitted to the Department for approval. The checks must be performed at intervals not to exceed 1 week.

007.21(G)  **PROPER OPERATION.** To satisfy the requirement of 180 NAC 20-007.21(F), safety quality assurance checks must ensure proper operation of:

(i) Electrical interlocks at each external beam radiation therapy room entrance;

(ii) Proper operation of the "BEAM-ON", interrupt and termination switches;

(iii) Beam condition indicator lights on the access doors, control console, and in the radiation therapy room;

(iv) Viewing systems; and

(v) Electrically operated treatment room door or doors from inside and outside the treatment room.

007.21(H)  **PROMPTLY REPAIR.** The registrant must promptly repair any system identified in 180 NAC 20-007.21(G) that is not operating properly.

007.21(I)  **QUALITY ASSURANCE CHECK RECORDS.** The registrant must maintain a record of each quality assurance check required by 180 NAC 20-007.021(A) and (G) for three years. The record must include:

(i) The date of the quality assurance check;

(ii) The manufacturer's name, model number, and serial number of the therapeutic radiation machine;
(iii) The manufacturer's name, model number, and serial number for the instrument or instruments used to measure the radiation output of the therapeutic radiation machine; and
(iv) The signature of the individual who performed the periodic quality assurance check.

007.21(J) QUALITY ASSURANCE CHECKS FOR INTENSITY MODULATED RADIATION THERAPY (IMRT) SYSTEMS. Quality assurance checks for intensity modulated radiation therapy (IMRT) systems must:
(i) Include commissioning and testing of the treatment planning and delivery systems, routine quality assurance of the delivery system, and patient-specific validation of treatment plans;
(ii) Be performed according to "Guidance document on delivery, treatment planning, and clinical implementation of IMRT: Report of the IMRT subcommittee of the AAPM radiation therapy committee: AAPM Report No. 82". This document is available for viewing at the Department of Health and Human Services, Division of Public Health, Office of Radiological Health, 301 Centennial Mall South, 3rd Floor, Lincoln, Nebraska 68509; and
(iii) Be performed according to the manufacturer's specifications.

008. CALIBRATION OF RADIATION SURVEY INSTRUMENTS. The requirements for calibration of survey instruments are as follows.

008.01 CALIBRATION FREQUENCY. The registrant must ensure that the survey instruments used to show compliance with this chapter have been calibrated before first use, at intervals not to exceed 12 months, and following repair.

008.02 CALIBRATION METHOD. To satisfy the requirements of 180 NAC 20-008.01, the registrant must:
(A) Calibrate all required scale readings up to 10 milliSievert (mSv) or 1000 millirem (mrem) per hour with an appropriate radiation source that is traceable to the National Institute of Standards and Technology (NIST); and
(B) Calibrate at least two points on each scale to be calibrated. These points should be at approximately 1/3 and 2/3 of full-scale.

008.03 PERFORMANCE. To satisfy the requirements of 180 NAC 20-008.02, the registrant must:
(A) Consider a point as calibrated if the indicated dose rate differs from the calculated dose rate by not more than 10 percent; and
(B) Consider a point as calibrated if the indicated dose rate differs from the calculated dose rate by not more than 20 percent if a correction factor or graph is conspicuously attached to the instrument.

008.04 RADIATION SURVEY INSTRUMENT CALIBRATION RECORDS. The registrant must retain a record of each calibration required in 180 NAC 20-008.01 for three years. The record must include:
(A) A description of the calibration procedure; and
(B) A description of the source used, the certified dose rates from the source, the rates indicated by the instrument being calibrated, the correction factors deduced from the calibration data, the signature of the individual who performed the calibration, and the date of calibration.

008.05 CALIBRATION SERVICES. The registrant may obtain the services of individuals licensed by the Department, the U. S. Nuclear Regulatory Commission, an Agreement State, or a Licensing State to perform calibrations of survey instruments. Records of calibrations that contain information required by 180 NAC 20-008.04 must be maintained by the registrant.

009. SHIELDING AND SAFETY DESIGN REQUIREMENTS. The design requirements for shielding and safety are as follows.

009.01 PRIMARY AND SECONDARY BARRIERS. Each therapeutic radiation machine subject to this chapter must be provided with primary or secondary barriers as necessary to ensure compliance with 180 NAC 4-005 and 4-013.

009.02 NEW INSTALLATION. Facility design information for all new installations of a therapeutic radiation machine or installations of a therapeutic radiation machine of higher energy into a room not previously approved for that energy, must be submitted for Department approval prior to actual installation of the therapeutic radiation machine. The minimum facility design information that must be submitted is contained in Appendix 20-A.

010. QUALITY ASSURANCE FOR RADIATION THERAPY SIMULATION SYSTEMS. The quality assurance requirements for radiation therapy simulation systems are as follows.

010.01 ACCEPTANCE TESTING AND PERIODIC VERIFICATION OF PERFORMANCE. Quality assurance for a conventional or virtual simulator must include acceptance testing and periodic verification of system performance.

010.02 METHOD. Quality assurance for radiation therapy simulation systems must be performed according to "Comprehensive QA for Radiation Oncology: Report of AAPM Radiation Therapy Committee Task Group No. 40: AAPM Report No. 46" for a conventional simulator.

010.02(A) VIRTUAL SIMULATOR. For radiation therapy virtual simulator's, quality assurance must be performed according to "Quality assurance for computed tomography simulators and the computed tomography-simulation process: Report of the AAPM Radiation Therapy Committee Task Group No. 66: AAPM Report No. 83". This document is available for viewing at the Department of Health and Human Services, Division of Public Health, Office of Radiological Health, 301 Centennial Mall South, 3rd Floor, Lincoln, Nebraska 68509.

011. ELECTRONIC BRACHYTHERAPY. This section addresses electronic brachytherapy.

011.01 APPLICABILITY. Electronic brachytherapy devices are subject to the requirements of 180 NAC 20-011, and are exempt for the requirements of 180 NAC 20-006.
011.01(A) PROHIBITION. An electronic brachytherapy device that does not meet the requirements of 180 NAC 20-011 must not be used for irradiation of patients.

011.01(B) APPROVAL FOR USE. An electronic brachytherapy device must only be utilized for human use applications specifically approved by the U.S. Food and Drug Administration (FDA) unless participating in a research study approved by the registrant’s Institutional Review Board (IRB).

011.02 POSSESSION OF SURVEY INSTRUMENTS. Each location authorized to use an electronic brachytherapy device according to 180 NAC 20-011 must possess appropriately calibrated portable monitoring equipment. As a minimum, the equipment must include a portable radiation measurement survey instrument capable of measuring dose rates over the range 10 microSievert ($\mu$Sv) or 1 millirem (mrem) per hour to 10 milliSievert (mSv) or 1000 millirem (mrem) per hour. The survey instrument or instruments must be operable and calibrated in accordance to 180 NAC 20-008 for the applicable electronic brachytherapy source energy.

011.03 FACILITY DESIGN REQUIREMENTS FOR ELECTRONIC BRACHYTHERAPY DEVICES. In addition to shielding adequate to meet requirements of 180 NAC 20-009, the treatment room must meet the following design requirements.

011.03(A) SIMULTANEOUS OPERATION. If applicable, provision must be made to prevent simultaneous operation of more than one therapeutic radiation machine in a treatment room.

011.03(B) ENTRANCE. Access to the treatment room must be controlled by a door at each entrance.

011.03(C) CONTINUOUS AURAL COMMUNICATION AND VISUAL OBSERVATION. Each treatment room must have provisions to permit continuous aural communication and visual observation of the patient from the treatment control panel during irradiation. The electronic brachytherapy device must not be used for patient irradiation unless the patient can be observed.

011.03(D) SHIELDING FOR DEVICES OPERATING BELOW 50 KILOVOLTS (KV). For electronic brachytherapy devices capable of operating below 50 kilovolts (kV), radiation shielding for the staff in the treatment room must be available, either as a portable shield or as localized shielded material around the treatment site.

011.03(E) SHIELDING FOR DEVICES OPERATING ABOVE 150 KILOVOLTS (KV). For electronic brachytherapy devices capable of operating at greater than 150 kilovolts (kV):

(i) The control panel must be located outside the treatment room; and

(ii) Electrical interlocks must be provided for all door or doors to the treatment room that must:

(1) Prevent the operator from initiating the treatment cycle unless each treatment room entrance door is closed;

(2) Cause the source to be shielded when an entrance door is opened; and
(3) Prevent the source from being exposed following an interlock interruption until all treatment room entrance doors are closed and the source on-off control is reset at the console.

011.04 ELECTRICAL SAFETY FOR ELECTRONIC BRACHYTHERAPY DEVICES. The requirements for electrical safety for electronic brachytherapy devices are as follows.

011.04(A) ELECTRICAL AND MAGNETIC INTERFERENCE. The high voltage transformer must be electrically isolated to prevent electrical and magnetic interference with the surrounding environment and ancillary equipment.

011.04(B) PROTECTIVE HOUSING. The high voltage transformer must be isolated from personnel, the operator, and the environment by a protective housing that can only be accessed through a cover requiring a tool for access or with electrical interlocks to prevent operation while open.

011.04(C) SAFETY LABELS. The high voltage transformer must have appropriate safety labels warning personnel of potential electrical shock or heat related injuries.

011.05 CONTROL PANEL FUNCTIONS. In addition to the displays required by other provisions of 180 NAC 20-011, the control panel must:

(A) Provide an indication of whether electrical power is available at the control panel and if activation of the electronic brachytherapy source is possible;
(B) Provide an indication of whether x-rays are being produced; and
(C) Provide a means for indicating electronic brachytherapy source potential and current;
   (i) Provide the means for terminating an exposure at any time; and
   (ii) Include an access control or locking device that will prevent unauthorized use of the electronic brachytherapy device.

011.06 TIMER. A suitable irradiation control device or timer must be provided to terminate the irradiation after a pre-set time interval or integrated charge on a dosimeter-based monitor.

011.06(A) TIMER INDICATIONS. A timer must be provided at the treatment control panel. The timer must indicate the planned setting and the time elapsed or remaining.

011.06(B) PREVENT EXPOSURE IF SET AT ZERO. The timer must not permit an exposure if set at zero.

011.06(C) CUMULATIVE DEVICE. The timer must be a cumulative device that activates with an indication of "BEAM-ON" and retains its reading after irradiation is interrupted or terminated. After irradiation is terminated and before irradiation can be reinitiated, it must be necessary to reset the elapsed time indicator.

011.06(D) IRRADIATION TERMINATION. The timer must terminate irradiation when a pre-selected time has elapsed, if any dose monitoring system has not previously terminated irradiation.
011.06(E) SHORT TIMER SETTING. The timer must permit setting of exposure times as short as 0.1 second.

011.06(F) TIMER ACCURACY. The timer must be accurate to within one percent of the selected value or 0.1 second, whichever is greater.

011.07 RADIOLOGICAL MEDICAL PHYSICIST SUPPORT. The services of a radiological medical physicist is required for registrants using electronic brachytherapy devices.

011.07(A) RADIOLOGICAL MEDICAL PHYSICIST RESPONSIBILITIES. The radiological medical physicist is responsible for:

(i) Evaluation of the output from the electronic brachytherapy source;
(ii) Generation of the necessary dosimetric information;
(iii) Supervision and review of treatment calculations prior to initial treatment of any treatment site;
(iv) Establishing the periodic and day-of-use quality assurance checks and reviewing the data from those checks as required in 20-011.10;
(v) Consultation with the user in treatment planning, as needed; and
(vi) Performing calculations and assessments regarding patient treatments that may constitute a misadministration.

011.07(B) RADIOLOGICAL MEDICAL PHYSICIST CONTACT INFORMATION. If the radiological medical physicist is not a full-time employee of the registrant, the operating procedures required by 20-011.08 must also specifically address how the radiological medical physicist is to be contacted for problems or emergencies, as well as the specific actions, if any, to be taken until the radiological medical physicist can be contacted.

011.08 OPERATING PROCEDURES. The requirements for operating procedures are as follows.

011.08(A) INDIVIDUALS PRESENT IN THE TREATMENT ROOM. Only individuals approved by the user or radiological medical physicist may be present in the treatment room during treatment.

011.08(B) PROHIBITION. Electronic brachytherapy devices must not be made available for medical use unless the requirements of 180 NAC 20-004.01, 20-011.09 and 20-010.10 have been met.

011.08(C) SECURE WHEN UNATTENDED. The electronic brachytherapy device must be inoperable, either by hardware or password, when unattended by qualified staff or service personnel.

011.08(D) MONITOR TO PREVENT UNSHIELDED EXPOSURE. During operation, the electronic brachytherapy device operator must monitor the position of all persons in the treatment room, and all persons entering the treatment room, to prevent entering persons from unshielded exposure from the treatment beam.
011.08(E) MECHANICAL SUPPORTING OR RESTRAINING DEVICES. If a patient must be held in position during treatment, mechanical supporting or restraining devices must be used.

011.08(F) WRITTEN PROCEDURES. Written procedures must be developed, implemented, and maintained for responding to an abnormal situation. These procedures must include:

(i) Instructions for responding to equipment failures and the names of the individuals responsible for implementing corrective actions; and

(ii) The names and telephone numbers of the users and the radiological medical physicist to be contacted if the device or console operates abnormally.

011.08(G) PHYSICAL LOCATION OF OPERATING AND EMERGENCY PROCEDURES. A copy of the current operating and emergency procedures must be physically located at the electronic brachytherapy device control console. If the control console is integral to the electronic brachytherapy device, the required procedures must be kept where the operator is located during electronic brachytherapy device operation.

011.08(H) CONTACT INFORMATION. Instructions must be posted at the electronic brachytherapy device control console to inform the operator of the names and telephone numbers of the users and radiological medical physicist to be contacted if the device or console operates abnormally.

011.09 SAFETY PRECAUTIONS FOR ELECTRONIC BRACHYTHERAPY DEVICES. The safety precautions for electronic brachytherapy devices are as follows.

011.09(A) MONITORING. A radiological medical physicist must determine which persons in the treatment room require monitoring when the beam is energized.

011.09(B) PHYSICALLY PRESENT DURING INITIATION OF TREATMENTS. A user and a radiological medical physicist must be physically present during the initiation of all patient treatments involving the electronic brachytherapy device.

011.09(C) PHYSICALLY PRESENT DURING CONTINUATION OF TREATMENTS. A radiological medical physicist and either a user, a physician, or electronic brachytherapy device operator, under the supervision of a user, who has been trained in the operation and emergency response for the electronic brachytherapy device, must be physically present during continuation of all patient treatments involving the electronic brachytherapy device.

011.09(D) USE OF A RADIATION SURVEY METER. When shielding is required by 180 NAC 20-011.03D the electronic brachytherapy device operator must use a survey meter to verify proper placement of the shielding immediately upon initiation of treatment. Alternatively, a radiological medical physicist must designate shield locations sufficient to meet the requirements of 180 NAC 4-005 for any individual, other than the patient, in the treatment room.
011.09(E) PERSONNEL SHIELDING. All personnel in the treatment room are required to remain behind shielding during treatment. A radiological medical physicist must approve any deviation from this requirement and must designate alternative radiation safety protocols, compatible with patient safety, to provide an equivalent degree of protection.

011.10 ELECTRONIC BRACHYTHERAPY SOURCE CALIBRATION MEASUREMENTS. The requirements for electronic brachytherapy source calibration measurements are as follows.

011.10(A) PERFORMANCE OF ELECTRONIC BRACHYTHERAPY SOURCE OUTPUT CALIBRATION. Calibration of the electronic brachytherapy source output for an electronic brachytherapy device subject to 180 NAC 20-011 must be performed by, or under the direct supervision of, a radiological medical physicist.

011.10(B) CALIBRATION FREQUENCY. Calibration of the electronic brachytherapy source output must be made for each electronic brachytherapy source, after any repair affecting the x-ray beam generation, and when indicated by the electronic brachytherapy source quality assurance checks.

011.10(C) DOSIMETRY SYSTEM. Calibration of the electronic brachytherapy source output must utilize a dosimetry system described in 180 NAC 20-004.03(A).

011.10(D) ELECTRONIC BRACHYTHERAPY SOURCE OUTPUT CALIBRATION. Calibration of the electronic brachytherapy source output must include, as applicable, determination of:

(i) The output within two percent of the expected value, if applicable, or determination of the output if there is no expected value;
(ii) Timer accuracy and linearity over the typical range of use;
(iii) Proper operation of back-up exposure control devices;
(iv) Evaluation that the relative dose distribution about the source is within five percent of that expected; and
(v) Source positioning accuracy to within one millimeter within the applicator;

011.10(E) CALIBRATION PROTOCOL. Calibration of the x-ray source output required by 180 NAC 20-010.09(A) through (F) must be those in current published recommendations from a recognized national professional association with expertise in electronic brachytherapy. The manufacturer’s calibration protocol must be followed.

011.10(F) CALIBRATION RECORDS. The registrant must maintain a record of each calibration in an auditable form for the duration of the registration. The record must include:

(i) The date of the calibration;
(ii) The manufacturer’s name, model number, and serial number for the electronic brachytherapy device and a unique identifier for its electronic brachytherapy source;
(iii) The model numbers and serial numbers of the instrument or instruments used to calibrate the electronic brachytherapy device; and
(iv) The name and signature of the radiological medical physicist responsible for performing the calibration.
011.11 PERIODIC AND DAY-OF-USE QUALITY ASSURANCE CHECKS FOR ELECTRONIC BRACHYTHERAPY DEVICES. Periodic and day-of-use quality assurance checks for electronic brachytherapy devices are as follows.

011.11(A) QUALITY ASSURANCE CHECK FREQUENCY. Quality assurance checks must be performed on each electronic brachytherapy device subject to 180 NAC 20-011:
(i) At the beginning of each day-of-use;
(ii) Each time the device is moved to a new room or site. A site includes each day-of-use at each operating location for a self-contained electronic brachytherapy unit transported in a van or trailer; and
(iii) After each x-ray tube installation.

011.11(B) PROCEDURES FOR PERIODIC QUALITY ASSURANCE CHECKS. The registrant must perform periodic quality assurance checks required by 180 NAC 20-011.11(A) in procedures established by the radiological medical physicist.

011.11(C) QUALITY ASSURANCE CHECK CONTENT. To satisfy the requirements of 180 NAC 20-011.11(A), radiation output quality assurance checks must include, at a minimum:
(i) Verification that output of the electronic brachytherapy source falls within three percent of expected values, as appropriate for the device, as determined by:
   (1) Output as a function of time, or
   (2) Output as a function of setting on a monitor chamber.
(ii) Verification of the consistency of the dose distribution to within three percent of that found during calibration required by 180 NAC 20-011.10; and
(iii) Validation of the operation of positioning methods to ensure that the treatment dose exposes the intended location within one millimeter.

011.11(D) DOSIMETRY SYSTEM. The registrant must use a dosimetry system that has been intercompared within the previous twelve months with the dosimetry system described in 180 NAC 20-004.03(A) to make the quality assurance checks required in 180 NAC 20-011.11(C).

011.11(E) REVIEW OF QUALITY ASSURANCE CHECK RESULTS. The registrant must review the results of each radiation output quality assurance check according to the following procedures:
(i) A user and radiological medical physicist must be immediately notified if any parameter is not within its acceptable tolerance. The electronic brachytherapy device must not be made available for subsequent medical use until the radiological medical physicist has determined that all parameters are within their acceptable tolerances;
(ii) If all radiation output quality assurance check parameters appear to be within their acceptable range, the quality assurance check must be reviewed and signed by either the user or radiological medical physicist within two days; and
(iii) The radiological medical physicist must review and sign the results of each radiation output quality assurance check at intervals not to exceed thirty days.
011.11(F) SAFETY DEVICE QUALITY ASSURANCE CHECKS. To satisfy the requirements of 180 NAC 20-011.11(A), safety device quality assurance checks must, at a minimum, assure:

(i) Proper operation of radiation exposure indicator lights on the electronic brachytherapy device and on the control console;
(ii) Proper operation of viewing and intercom systems in each electronic brachytherapy facility, if applicable;
(iii) Proper operation of radiation monitors, if applicable;
(iv) The integrity of all cables, catheters or parts of the device that carry high voltages; and
(v) Connecting guide tubes, transfer tubes, transfer-tube-applicator interfaces, and treatment spacers are free from any defects that interfere with proper operation.

011.11(G) SAFETY DEVICE QUALITY ASSURANCE CHECK RESULTS. If the results of the safety device quality assurance checks required in 180 NAC 20-011.11(F) indicate the malfunction of any system, a registrant must secure the control console in the OFF position and not use the electronic brachytherapy device except as may be necessary to repair, replace, or check the malfunctioning system.

011.11(H) QUALITY ASSURANCE CHECK PROTOCOL. Quality assurance checks required by 20-011.11(A) through (F) must be those in current published recommendations from a recognized national professional association with expertise in electronic brachytherapy, or the manufacturer’s quality assurance check protocol.

011.11(I) RECORD MAINTENANCE. The registrant must maintain a record of each quality assurance check required by 180 NAC 20-011.11(C) and 20-011.11(F) in an auditable form for three years and must include:

(i) The date of the quality assurance check;
(ii) The manufacturer’s name, model number, and serial number for the electronic brachytherapy device;
   (1) The name and signature of the individual who performed the periodic quality assurance check; and
   (2) The name and signature of the radiological medical physicist who reviewed the quality assurance check; and
(iii) For radiation output quality assurance checks required by 180 NAC 20-011.11(C) the record must also include:
   (1) The unique identifier for the electronic brachytherapy source and the manufacturer’s name; and
   (2) The model number and serial number for the instrument or instruments used to measure the radiation output of the electronic brachytherapy device.

011.12 THERAPY-RELATED COMPUTER SYSTEMS. The registrant must perform acceptance testing on the treatment planning system of electronic brachytherapy related computer systems according to current published recommendations from a recognized national professional association with expertise in electronic brachytherapy, when available. In the absence of an acceptance testing protocol published by a national professional association, the manufacturer’s acceptance testing protocol must be followed.
011.12(A) ACCEPTANCE TESTING. Acceptance testing must be performed by, or under the direct supervision of, radiological medical physicist. At a minimum, the acceptance testing must include, as applicable, verification of:
   (i) The source-specific input parameters required by the dose calculation algorithm;
   (ii) The accuracy of dose, dwell time, and treatment time calculations at representative points;
   (iii) The accuracy of isodose plots and graphic displays;
   (iv) The accuracy of the software used to determine radiation source positions from radiographic images; and
   (v) If the treatment-planning system is different from the treatment-delivery system, the accuracy of electronic transfer of the treatment delivery parameters to the treatment delivery unit from the treatment planning system.

011.12(B) APPLICATOR POSITION INDICATORS. The position indicators in the applicator must be compared to the actual position of the source or planned dwell positions, as appropriate, at the time of commissioning.

011.12(C) INDEPENDENT VERIFICATION. Prior to each patient treatment regimen, the parameters for the treatment must be evaluated and approved by the user and the radiological medical physicist for accuracy through means independent of that used for the determination of the parameters.

011.13 TRAINING. The requirements for training are as follows.

011.13(A) TRAINING ON OPERATING PROCEDURES. A registrant must provide instruction, initially and at least annually, to all individuals who operate the electronic brachytherapy device, as appropriate to the individual's assigned duties, in the operating procedures identified in 180 NAC 20-011.08. If the interval between patients exceeds one year, retraining of the individuals must be provided.

011.13(B) DEVICE SPECIFIC TRAINING. In addition to the requirements of 180 NAC 20-003.03 for therapeutic radiation machine users and 180 NAC 20-003.04 for radiological medical physicists, these individuals must also receive device specific instruction initially from the manufacturer, and annually from either the manufacturer or other qualified trainer. The training must be of a duration recommended by a recognized national professional association with expertise in electronic brachytherapy, when available. In the absence of any training protocol recommended by a national professional association, the manufacturer’s training protocol must be followed. The training must include, but not be limited to:
   (i) Device-specific radiation safety requirements;
   (ii) Device operation;
   (iii) Clinical use for the types of use approved by the U.S. Food and Drug Administration (FDA);
   (iv) Emergency procedures, including an emergency drill; and
   (v) The registrant’s quality assurance program.

011.13(C) RECORDS. A registrant must retain a record of individuals receiving instruction required by 180 NAC 20-011.13(A) and (B) for three years. The record must include:
(i) A list of the topics covered;
(ii) The date of the instruction;
(iii) The name or names of the attendee or attendees; and
(iv) The name or names of the individual or individuals who provided the instruction.

011.14 MOBILE ELECTRONIC BRACHYTHERAPY SERVICE. This section addresses the requirements for registrants providing mobile electronic brachytherapy service.

011.14(A) SURVEY INSTRUMENT FUNCTION CHECK. All survey instruments must be checked before medical use at each address of use or on each day-of-use, whichever is more restrictive.

011.14(B) ACCOUNT FOR ELECTRONIC BRACHYTHERAPY SOURCE BEFORE DEPARTURE. Each electronic brachytherapy source must be accounted for in the electronic brachytherapy device before departure from the client's address.

011.14(C) QUALITY ASSURANCE CHECKS. At each location on each day-of-use, all of the required quality assurance checks specified in 180 NAC 20-011.11 must be performed to assure proper operation of the device.

012. OTHER USES OF ELECTRONICALLY PRODUCED RADIATION TO DELIVER THERAPEUTIC RADIATION DOSAGE. A person must not utilize any device which is designed to electrically generate a source of ionizing radiation to deliver therapeutic radiation dosage, and which is not regulated under any existing category of therapeutic radiation machine, until the following requirements have been met.

012.01 SUBMISSION REQUIREMENTS. The applicant or registrant must submit:
   (A) A detailed description of the device and its intended application or applications;
   (B) Facility design requirements, including shielding and access control;
   (C) Documentation of appropriate training for users and radiological medical physicist or physicists;
   (D) Methodology for measurement of dosages to be administered to patients or human research subjects;
   (E) Documentation regarding calibration, maintenance, and repair of the device, as well as instruments and equipment necessary for radiation safety
   (F) Radiation safety precautions and instructions; and
   (G) Other information requested by the Department in its review of the application.

012.02 WRITTEN APPROVAL. The applicant or registrant must receive written approval from the Department to utilize a device specified in 180 NAC 20-012. The device may be used only as specified in this chapter and specific conditions the Department considers necessary for the medical use of the device.
INFORMATION ON RADIATION SHIELDING REQUIRED FOR PLAN REVIEWS

I. ALL THERAPEUTIC RADIATION MACHINES MUST SUBMIT THE FOLLOWING INFORMATION:
   (A) Basic information including: name and telephone number of the individual responsible for preparation of the shielding plan; name and telephone number of the supervisor at the registrant's facility; and the street address, including room number of the therapeutic radiation machine. The plan should also indicate whether this is a new structure or a modification to existing structure or structures;
   (B) All wall, floor, and ceiling areas struck by the useful beam must have primary barriers; and
   (C) Secondary barriers must be provided in all wall, floor, and ceiling areas not having primary barriers.

II. THERAPEUTIC RADIATION MACHINES UP TO 150 KILOVOLTS (KV) - PHOTONS ONLY. In addition to the information listed in Section I, registrants using therapeutic radiation machines that produce only photons with a maximum energy less than or equal to 150 kilovolts (kV) must submit shielding plans which contain, at a minimum, the following additional information:
   (A) Equipment specifications, including the manufacturer and model number of the therapeutic radiation machine, as well as the maximum technique factors;
   (B) Maximum design workload for the machine including total weekly radiation output, expressed in gray or rad, or air kerma at 1 meter, total beam-on time per day or week, the average treatment time per patient, along with the anticipated number of patients to be treated per day or week;
   (C) A facility blueprint or drawing indicating: scale, 0.25 inch = 1 foot is typical; direction of North; normal location of the therapeutic radiation machine's radiation port or ports; the port's travel and traverse limits; general direction or directions of the useful beam; locations of any windows and doors; and the location of the therapeutic radiation machine control panel. If the control panel is located inside the therapeutic radiation machine treatment room, the location of the operator's booth must be noted on the plan and the operator's station at the control panel must be behind a protective barrier sufficient to ensure compliance with 180 NAC 4-005;
   (D) The structural composition and thickness or lead or concrete equivalent of all walls, doors, partitions, floor, and ceiling of the room or rooms concerned;
   (E) The type of occupancy of all adjacent areas inclusive of space above and below the room or rooms concerned. If there is an exterior wall, show distance to the closest area or areas where it is likely that individuals may be present; and
   (F) At least one example calculation which shows the methodology used to determine the amount of shielding required for each physical condition, primary and secondary or leakage barriers, restricted and unrestricted areas, entry door or doors and shielding material in the facility:
      (i) If commercial software is used to generate shielding requirements, also identify the software used and the version or revision date.
      (ii) Submit quality control sample calculations to verify the result obtained with the software.
III. THERAPEUTIC RADIATION MACHINES OVER 150 KILOVOLTS (KV). In addition to the information listed in Section I, registrants using therapeutic radiation machines that produce photons with a maximum energy in excess of 150 kilovolts (kV) or electrons must submit shielding plans which contain, as a minimum, the following additional information:

(A) Equipment specifications including the manufacturer and model number of the therapeutic radiation machine, and gray (rad) at the isocenter and the energy or energies and type or types of radiation produced, photon or electron. The target to isocenter distance must be specified;

(B) Maximum design workload for the machine including total weekly radiation output, expressed in gray or rad at 1 meter, total beam-on time per day or week, the average treatment time per patient, along with the anticipated number of patients to be treated per day or week;

(C) Facility blueprint or drawing [including both floor plan and elevation views] indicating relative orientation of the therapeutic radiation machine, scale, 0.25 inch = 1 foot is the typical scale, type or types, thickness and minimum density of shielding material or materials, direction of North, the locations and size of all penetrations through each shielding barrier, ceiling, walls and floor, as well as details of the door or doors and maze;

(D) The structural composition and thickness or concrete equivalent of all walls, doors, partitions, floor, and ceiling of the room or rooms concerned;

(E) The type of occupancy of all adjacent areas inclusive of space above and below the room or rooms concerned. If there is an exterior wall, show distance to the closest area or areas where it is likely that individuals may be present;

(F) Description of all assumptions that were used in shielding calculations including, but not limited to, design energy, room may be designed for 6 megavolt (MV) unit although only a 4 megavolt (MV) unit is currently proposed, work load, presence of integral beam stop in unit, occupancy and use or uses of adjacent areas, fraction of time that useful beam will intercept each permanent barrier, walls, floor and ceiling, and expected radiation exposure in both restricted and unrestricted areas; and

(G) At least one example calculation which shows the methodology used to determine the amount of shielding required for each physical condition, primary and secondary leakage barriers, restricted and unrestricted areas, small angle scatter, entry door or doors and maze, and shielding material in the facility:

(i) If commercial software is used to generate shielding requirements, also identify the software used and the version or revision date; and

(ii) If the software used to generate shielding requirements is not in the open literature, also submit quality control sample calculations to verify the result obtained with the software.

IV. NEUTRON SHIELDING. In addition to the information listed in Section III, registrants using therapeutic radiation machines that are capable of operating above 10 megavolts (MV) must submit shielding plans which contain, as a minimum, the following additional information:

(A) The structural composition, thickness, minimum density and location of all neutron shielding material;

(B) Description of all assumptions that were used in neutron shielding calculations including, but not limited to, neutron spectra as a function of energy, neutron fluence rate, absorbed dose and dose equivalent, due to neutrons, in both restricted and unrestricted areas;
(C) At least one example calculation which shows the methodology used to determine the amount of neutron shielding required for each physical condition, restricted and unrestricted areas, entry door or doors and maze, and neutron shielding material utilized in the facility:
   (i) If commercial software is used to generate shielding requirements, also identify the software used and the version or revision date; and
   (ii) If the software used to generate shielding requirements is not in the open literature, also submit quality control sample calculations to verify the results obtained with the software; and

(D) The method or methods and instrumentation that will be used to verify the adequacy of all neutron shielding installed in the facility.

V. REFERENCES. References for shielding designs are as follows and are available for viewing at the Department of Health and Human Services, Division of Public Health, Office of Radiological Health, 301 Centennial Mall South, 3rd Floor, Lincoln, Nebraska 68509:
   (C) NCRP Report 144, "Radiation Protection for Particle Accelerator Facilities" (2003).