X-RAY
REGULATORY GUIDE

VETERINARY X-RAY FACILITIES

May 1, 2016
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MDH Veterinary X-ray Regulatory Guide
INTRODUCTION TO THE REGULATORY GUIDE FOR VETERINARY REGISTRANTS

Minnesota Department of Health Mission, X-ray Program Mission
The mission of the Minnesota Department of Health (MDH) Radiation Control X-ray Unit is to protect and promote radiation safety through guidance and collaboration with the radiation community. Our vision is to reduce unnecessary radiation exposure from the use of ionizing radiation producing equipment.

Introduction
This guide is designed to describe the type and extent of a radiation safety/quality assurance program necessary for the safe use of x-ray equipment and compliance with Minnesota Rules, Chapter 4732 in veterinary facilities. The information in this guide is not a substitute for radiation safety/quality assurance training or for developing and implementing an effective radiation safety/quality assurance program. You should carefully study this guide and Minnesota Rules, Chapter 4732.


This guidance, instruction sheets, and additional information, is available on the X-ray Unit website as they are developed.

- http://www.health.state.mn.us/xray

Implementation
The information in this guide is intended to assist in compliance with Minnesota Rules, Chapter 4732. This guide provides one set of methods approved by MDH for meeting the regulations and represents the minimum acceptable standards. MDH has included many useful “Instruction Sheets” to assist you in creating your radiation safety/quality assurance program and complying with the Minnesota Rules, Chapter 4732. If you have questions, please contact the Radiation Control, X-ray Unit at (651)201-4545 or email at health.xray@state.mn.us.

Revisions to the MDH Regulatory Guide of Veterinary X-ray Facilities
MDH X-ray Unit is always striving to better the information that we provide to the veterinary registrants. This may include additions to the information presented in this guide. There may be occasion for revisions to this guide. These revisions are not changes to Minnesota Rules, Chapter 4732 and are intended to clarify or supplement what is already within the guide.

Any revisions to this guide will be documented in the Summary of Revisions at the end of this guide.

Note: Ionizing radiation producing equipment will be identified as “x-ray equipment” throughout this guidance document.
APPLICATION FOR REGISTRATION (4732.0200)

All facilities or individuals in possession of x-ray equipment must apply for registration using the current application process provided by the commissioner. Registration with payment must be completed by the facility and submitted to MDH. The Initial Registration application process may be found on our website.

*Note: Service providers are not responsible for registration of facility or x-ray equipment. Service providers are required to notify MDH when they deliver or install x-ray equipment in your facility. This is in addition to facility application requirements for registration and notification to MDH.*

ADDITIONAL REGISTRATION INFORMATION

- Registrants, who purchase replacement x-ray equipment, must notify MDH within 30 days of obtaining the replacement equipment.
- Registrants who purchase additional x-ray equipment must submit a registration using the current application process provided by the commissioner and submit a fee for each new tube within 30 days of obtaining the equipment and prior to use.
- Registrants must notify MDH when there is a change in ownership or when x-ray equipment is placed in storage or removed from the registrant’s physical location. This is to ensure the following:
  1. X-ray equipment is only possessed, used, or controlled by persons who have valid MDH facility registrations
  2. X-ray equipment is properly handled and secured
  3. Public health and safety are not compromised by the unauthorized use of x-ray equipment
- The Additional Registration application process may be found on our website.

FEES (4732.0210)

<table>
<thead>
<tr>
<th>Fee Type</th>
<th>Amount</th>
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<tbody>
<tr>
<td>Facility Base Fee: due initially and annually</td>
<td>$100</td>
</tr>
<tr>
<td>Veterinary X-ray Equipment Fee</td>
<td>$100</td>
</tr>
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<td>Dental X-ray Equipment Fee</td>
<td>$40</td>
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Submit your registration online at [www.health.state.mn.us/xray](http://www.health.state.mn.us/xray) or mail to:

Minnesota Department of Health
Radiation Control X-ray Unit
625 Robert Street North
PO Box 64497
St. Paul, Minnesota 55164-0497
INDIVIDUAL RESPONSIBLE FOR THE RADIATION SAFETY/QUALITY ASSURANCE PROGRAM (4732.0500)

You are responsible for x-ray equipment that is under your administrative control and must ensure your radiation safety/quality assurance program, your staff and the use of your x-ray equipment is in compliance with Minnesota Rules, Chapter 4732. To ensure adequate oversight is provided to your radiation safety/quality assurance program, a radiation safety officer must be designated and identified within your radiation safety/quality assurance program.

RADIATION SAFETY OFFICER (RSO) (4732.0500 - 4732.0505)

The Radiation Safety Officer (RSO) is responsible for the day-to-day operations of your radiation safety/quality assurance program. The RSO must receive RSO specific training and has additional responsibilities beyond his/her day to day job duties. The RSO must be provided sufficient time and commitment from your management to stop operations that he/she considers unsafe, ensure x-ray equipment is used safely, and maintain compliance with Minnesota Rules, Chapter 4732. These responsibilities, time and commitment must be delegated in writing from your management to the RSO.

Note: When the registrant is also the RSO, an RSO Delegation Agreement does not have to be completed. You must notify MDH in writing when there is a change to your RSO delegation.

Additional Radiation Safety Officer Information can be found in Attachment A and at the end of this guide:

- Typical duties and responsibilities
- Training and experience requirements
- Radiation Safety Officer Delegation Agreement
- [http://www.health.state.mn.us/divs/eh/radiation/xray/rso.html](http://www.health.state.mn.us/divs/eh/radiation/xray/rso.html)

Radiation Safety Officer Responsibilities

1. Ensuring the safe use of radiation
2. Identifying x-ray radiation protection problems
3. Recommending and providing corrective actions
4. Stopping unsafe activities
5. Managing the Radiation Protection Program.
6. Ensuring quality control tests are documented and completed
7. Verifying implementation of corrective actions
8. Ensuring compliance with state regulations
OPERATOR QUALIFICATIONS

Minnesota Rules, Chapter 4732 does not require operators of x-ray equipment for non-human use to meet minimum qualification requirements. However, the requirements of Minnesota Statue 156 for Veterinary Medicine must be met. Currently, these individuals include licensed veterinarians and any employee who is under the direction and supervision of a licensed veterinarian.

For specific questions regarding Minnesota Statute, Chapter 156 or the qualification requirements, you must contact the Minnesota Board of Veterinary Medicine.

- [https://www.revisor.mn.gov/statutes/?id=156.12](https://www.revisor.mn.gov/statutes/?id=156.12)
- [http://www.vetmed.state.mn.us](http://www.vetmed.state.mn.us)

OPERATOR TRAINING

Minnesota Rules, Chapter 4732.0510 requires that all individuals operating x-ray equipment must receive initial training in facility specific and system specific safe operating procedures, emergency procedures, quality control procedures, and proper protective shielding. Additional training must be conducted when there is a change to the radiation safety/quality assurance program, when existing x-ray equipment is upgraded, or when new x-ray equipment is added.

Additional training is required for staff that uses:

- Computed Tomography (CT), [Minnesota Rules, Chapter 4732.0860](https://www.revisor.mn.gov/statutes/?id=4732.0860)
- Fluoroscopic x-ray equipment, [Minnesota Rules, Chapter 4732.0825](https://www.revisor.mn.gov/statutes/?id=4732.0825)

Additional facility specific training requirement information can be found in Attachment B.

The exposure of an individual for training, instruction, demonstration, or maintenance is prohibited. The use of x-ray equipment for these purposes will result in enforcement action that may include an administrative penalty of up to $10,000.

OPERATION OF X-RAY EQUIPMENT

MDH identifies operation of x-ray equipment to include the use, manipulation, or adjustment of any components for the controlled production of x-rays that may affect the dose to the patient, occupational staff, public or the quality of the raw image. This includes minimally x-ray equipment and x-ray system that are defined in [Minnesota Rules, Chapter 4732.0110](https://www.revisor.mn.gov/statutes/?id=4732.0110). These components would include:

- X-ray high-voltage generator
- X-ray control
- X-ray tube housing assembly
- X-ray beam-limiting device
- The necessary supporting structures
The following items are examples of operating x-ray equipment:

- Positioning or repositioning of the patient table during activation of the x-ray beam
- Setting or the adjustment of the x-ray control
- Manipulating the image intensifier and x-ray tube
- Manipulating the collimators
- Manipulation or adjustment of any components that may affect the dose to the patient, occupational staff, public or the quality of the raw image.
- Making the exposure
- Evaluating quality control tests

The following items are examples of what is not considered “operation” of x-ray equipment and may be performed by unqualified individuals:

- Movement of the portable x-ray equipment into a room
- Make the necessary equipment connections
- Plug the x-ray unit in
- Turning on the “power” to the x-ray equipment
- Inputting the patient identification and examination information
- Movement of the x-ray equipment, x-ray table or image receptors prior to the patient being positioned for the examination
- Processing patient films
- Post-processing of digital patient images

**OPERATION OF FLUOROSCOPIC X-RAY EQUIPMENT**

Operation of fluoroscopic x-ray equipment in a non-human use veterinary facility must be performed only by individuals who have met the fluoroscopic training requirements of Minnesota Rules, Chapter 4732.0825 subpart 2, and only when an individual licensed by the Minnesota Board of Veterinary Medicine to practice veterinary medicine is physically present in the room in accordance with Minnesota Rules, Chapter 4732.0306.
RADIATION SAFETY/QUALITY ASSURANCE PROGRAM
(4732.0520)

You are required to have a radiation safety/quality assurance program in place prior to first use of your x-ray equipment. A radiation safety/quality assurance program includes administrative, radiation safety and quality control procedures to ensure:

- Occupational staff and the public are protected through the safe operation of x-ray equipment
- Consistent, high-quality images will be produced with a minimum exposure to occupational staff and the public
- Compliance with state regulations

Your radiation safety/quality assurance program must include:

- Radiation Safety procedures for the safe and proper use of the x-ray equipment
- Quality control procedures to include tests used for routine assessment of an x-ray imaging system specific to your x-ray equipment and processing systems
- Training of the operators of x-ray equipment including documentation
- Radiation program audits
- Equipment performance tests, including a listing of the required tests, can be found in Attachment D of this guidance document

RADIATION PROGRAM AUDIT (4732.0540)

The RSO and management are required to audit your radiation safety/quality assurance program to ensure the continued safe use of veterinary x-ray equipment at intervals not to exceed 12 months. An audit is a review of your established radiation safety/quality assurance policies, procedures, and their implementation to ensure the safe operation of the x-ray equipment, protection of staff and the public, and compliance with Minnesota Rules, Chapter 4732. It is essential that problems identified are addressed and corrected promptly and comprehensively. MDH will review your radiation safety/quality assurance program audit and determine if corrective actions are thorough, timely, and sufficient to prevent recurrence.

An audit program for a veterinary office should include, at minimum, a review of:

- Operating and emergency procedures
- Quality control procedures
- Required training
- Proper use of holding devices and personal protective garments
- Proper use of individual monitoring (if applicable)
- Proper use of technique charts
- Quarterly Retake/Reject analysis
- Performance evaluations of the x-ray equipment
- All required records and maintenance of these records
You are responsible for the content and implementation of your radiation safety and quality assurance program and for all actions of your employees. Each registrant must create a site specific audit and audit form. See Attachment C for questions to consider in an annual audit.

AS LOW AS REASONABLY ACHIEVABLE (ALARA) (4732.0530)

Every reasonable effort should be made to maintain radiation exposures as low as is reasonably achievable (ALARA). You are required to consider the ALARA philosophy in establishing your radiation safety/quality assurance program involving the use of your veterinary x-ray equipment.

A typical ALARA program in a veterinary setting may include:

- Commitment from management and staff
- Implement procedures for holding the patient or image receptor
- The manufacturer’s instructions on the proper use of image receptor holding devices
- The manufacturer’s instructions for proper image development
- Implementing site specific radiation safety procedures
- Implementing site specific quality control procedures

Operating and emergency procedures must be developed, implemented, and maintained to ensure that veterinary x-ray equipment is used only as designed, control and accountability are maintained, and radiation doses received by occupational workers and members of the public are ALARA and below the regulatory dose limits.

The success of an ALARA program depends on the cooperation of each person at your facility. Management should make a formal policy commitment to the ALARA philosophy and implement that commitment with adequate resources. A model ALARA management program is contained in Attachment E of this guide.
OPERATING AND EMERGENCY PROCEDURES
(4732.0500’S) (4732.0800’S)
Establishing and implementing operating and emergency procedures promotes good radiation safety practices to reduce the unnecessary radiation exposure received by occupational staff, individuals who must remain in the room during the x-ray examination, and the public. The registrant must implement site specific operating and emergency procedures including, but not limited to:

- The safe use of x-ray equipment
- Emergency operating procedures
- Holding of patient or image receptor and lead apron use
- Declaring pregnant staff
- Thyroid and eye protection
- Ordering of x-ray exams and equivalent procedure for identifying the individual ordering the examination
- Individual monitoring devices and assessing the need for individual monitoring

THE SAFE USE OF X-RAY EQUIPMENT
- Develop a technique chart to ensure a proper and consistent exposure for the anatomical region to be imaged
- Properly set the line voltage adjustment prior to the examination (if applicable)
- Proper use of mechanical holding devices
- Ensuring only trained individuals operate x-ray equipment
- Ensuring a physician is in the room during fluoroscopy procedures
- Ensuring all individuals are properly protected (see Holding of Patient or Image Receptor and Lead Apron Use)
- Ensuring there is an unobstructed view of the patient during x-ray exam
- All x-ray equipment must have the following warning label and it must be located on the control panel, be legible and accessible to view

WARNING
This x-ray unit may be dangerous to patient and operator unless safe exposure factors, operating instructions and maintenance schedules are observed

In addition, when dental intraoral imaging is performed, you must ensure:
- All individuals wear a 0.5 mm lead equivalent apron or must be at least 6 feet from the patient or x-ray tube during the exam
- The x-ray tube and arm assembly is stable
- There is an unobstructed view of the patient during x-ray exam
EMERGENCY OPERATING PROCEDURES (4732.0510)

You must have procedures in place in the event of an x-ray equipment malfunction, including:

- Switching off the power by turning off main power on the control panel or unplugging the x-ray equipment according to manufacturer’s specifications
- Remove the patient from the exam room
- Remove the x-ray equipment from service
- Contact the RSO
- Contact the service provider, when necessary
- Ensure x-ray equipment is functioning properly before next use

HOLDING OF PATIENT OR IMAGE RECEPTOR AND LEAD APRON USE (4732.0510)

Individuals must not hold patients or image receptors, unless protected by a 0.5 millimeter lead equivalent apron. Only individuals necessary for the exam may be allowed in the room during an x-ray exam.

- 0.5 millimeter lead equivalent apron use is required for any individual that must remain in the x-ray operatory
- Individuals must be positioned so that no part of the body, protected or unprotected, will be struck by the useful beam
- Collimate the primary x-ray beam to the area of interest
- Individuals may not routinely be used to hold the patient or imaging receptor
- Staff must be rotated
- Pregnant staff should not be used to hold the patient or image receptor

<table>
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<th>Equipment Type</th>
<th>Patient Apron Required?</th>
<th>0.5 mm Lead Apron Required for Others in the Room?</th>
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<tr>
<td>Dental</td>
<td>No</td>
<td>If within 6’ of patient or tube</td>
</tr>
<tr>
<td>Bone Densitometry</td>
<td>If Primary Beam within 2” of the gonads</td>
<td>If within 6’ of patient or tube</td>
</tr>
<tr>
<td>Portable Radiography</td>
<td>If Primary Beam within 2” of the gonads</td>
<td>If within 6’ of patient or tube</td>
</tr>
<tr>
<td>General Use Radiographic</td>
<td>If Primary Beam within 2” of the gonads</td>
<td>All personnel in room must wear an apron</td>
</tr>
<tr>
<td>Fluoroscopic</td>
<td>If Primary Beam within 2” of the gonads</td>
<td>All personnel in room must wear an apron</td>
</tr>
<tr>
<td>Computed Tomography</td>
<td>If Primary Beam within 2” of the gonads</td>
<td>All personnel in room must wear an apron</td>
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When dental imaging is performed, all individuals must wear a 0.5 mm lead equivalent apron, or must be at least 6 feet from the patient or tube during the exam.
THYROID PROTECTION AND EYE PROTECTION (4732.0410)

Eye and thyroid protection must be worn if the potential exposure to the individual would exceed the dose limits of Minnesota Rules, Chapter 4732.0410, including: dose to the lens of the eye exceeding 5 Rem, and dose to the thyroid exceeding 50 Rem.

Though exceeding these dose limits in diagnostic veterinary imaging is unlikely, you and your staff must be aware of the potential exposures to the eyes and thyroid and take precautions as needed.

DECLARED PREGNANT STAFF (4732.0415) (4732.0440)

You must have written procedures in place for declared pregnant staff and ensure the dose limits in are not exceeded when staff declares pregnancy in writing. Options are available to registrants for declared pregnant staff:

- Ensure the individual remains outside of the x-ray room/area when performing x-ray examinations and maintains safe operating procedures
- Remove individual from x-ray area during pregnancy
- Individual should not be in the x-ray room/area, hold the patient or image receptor during the x-ray exam
- Provide individual with fetal monitoring, when necessary, in accordance with Minnesota Rules, Chapter 4732.0415 and 4732.0440
- Additional information on declared pregnancy and prenatal dose may be found on our Topic Index, click on Prenatal Exposure.

ORDERING OF X-RAY EXAMS (4732.0560)

Minnesota Rules, Chapter 4732.0560 establishes minimum requirements on who must order x-ray examinations and what must be included in an x-ray order:

- X-rays must be ordered by a licensed veterinarian, per Minnesota Statute 156.12
- The operator may not carry out an examination unless ordered by a licensed veterinarian
- An order for an examination must be available to the operator of the x-ray equipment at the time of the examination
- The order must include:
  1. Identification of the patient to be radiographed
  2. Identification of the veterinarian ordering the examination through:
     - Signature
     - Electronic signature
     - Equivalent procedure
  3. Clearly stated clinical indications
  4. The exact anatomical part to be radiographed
  5. The examination to be performed
EQUIVALENT PROCEDURE FOR IDENTIFYING THE VETERINARIAN ORDERING THE EXAMINATION

You are responsible for ensuring that all examinations are ordered by a licensed veterinarian and the order is available to personnel at the time of the examination.

- The order must be available to the individual performing the examination when the order is not given directly to the operator of the x-ray equipment by the veterinarian
- The order may be in a hardcopy format to include the information required above
- The order may be in an electronic format to include the information required above
- The order may be verbally given to the operator of the x-ray equipment by the ordering veterinarian to include the information required above, including a signature on the order

INDIVIDUAL MONITORING DEVICES (4732.0400 - 4732.0440)

Note: Individual Monitoring device will be identified as “dosimetry badge” throughout this guide.

You must have written procedures in place for individual monitoring devices. Minnesota Rules, Chapter 4732.0440 requires you to provide occupational staff with appropriate dosimetry badges and require personnel to wear the monitoring devices if:

1. Personnel are likely to receive greater than 10% of the dose limits in Minnesota Rules, Chapter 4732.0410. See Assessing the Need for Individual Monitoring.

   Due to the nature of veterinary imaging, where x-ray operator and others assist in the x-ray imaging, veterinary occupational staff have the potential to receive greater than the 10% (500 millirem)

2. A declared pregnant women is likely to receive during the entire pregnancy a dose in excess of 0.1 rem (100 millirem)
   - Facilities may remove pregnant staff from performing or assisting in any x-ray imaging during the pregnancy. See Declared Pregnant Staff.
3. Each individual who enters a high radiation area or very high radiation area
   - Highly unlikely in diagnostic veterinary imaging
4. A minor likely to receive in one year a dose in excess of 0.1 rem (100 millirem).

When dosimetry badges are used, you are responsible, through the RSO, to ensure:

- Dosimetry badges are worn and cared for properly.
- Minnesota Rules, Chapter 4732.0440, subpart 2 and subpart 3, states that dosimetry badges are for each individual and must not to be shared.
- The specific locations for wearing dosimetry badges are as follows:
   1. When a protective apron is not worn, on the trunk of the body or at the unshielded location of the whole body likely to receive the highest exposure
2. When wearing a protective apron and a single dosimeter is worn:
   - Outside of the protective apron at the collar
   - Declared pregnant staff must wear the dosimeter at the abdomen and under the protective apron

3. When wearing a protective apron on and two dosimeters are worn:
   - Outside of the protective apron at the collar
   - At the waist under the protective apron
   - A control dosimetry badge should be included each time a new shipment of dosimetry badges is received. The control badge accompanies staff dosimetry badges and monitors any radiation received on the staff dosimetry badges during shipment.
     1. The control badge must be kept in an area of natural background radiation at the facility between shipments.
     2. It is good practice to maintain staff dosimetry badges with the control dosimetry badge when not in use.
     3. The control badge must be shipped back to the dosimetry provider each time staff dosimetry badges are sent in for reading.

- Individual monitoring reports are reviewed according to Minnesota Rules, Chapter 4732.0505, subpart E:
  1. The RSO is responsible to ensure individual monitors are returned to the dosimetry provider for evaluation at the established frequency, typically on a monthly or quarterly frequency
  2. Dosimetry badge reports are reviewed to ensure staff are maintaining individual exposure levels as low as possible

- Individual monitoring records are maintained according to Minnesota Rules, Chapter 4732.0440, subpart 13 and subpart 14:
  1. Records must be maintained for 30 years after termination or the lifetime of the individual
  2. Staff must receive a written notification of occupational dose received annually (not to exceed 12 months), as a hardcopy, or softcopy via email or the dosimetry provider’s website

- Terminated staff must receive a written report of their occupational dose received within 30 days of registrant’s receipt of dosimetry badge report from the dosimetry provider

- Attempt to obtain previous dose records for new employees
  1. Obtain information from the individual (termination exposure record)
  2. If you are unable to obtain pre-employment occupational dose records, you must reduce the individual’s annual limit by 1.25 rem (1250 millirem) for each quarter exposure records are unavailable

Example: An individual is hired in July and is unable to provide any previous occupational dose history from a previous employer where dosimetry badges were required:
  1. Assume the new employee received an occupational exposure of:
     - 1.25 rem (1250 millirem) for the first quarter (Jan-March)
     - 1.25 rem for the second quarter (April-June)
2. This total of 2.5 rem (2500 millirem) must be reduced from the annual limit of 5.0 rem (5000 millirem)
3. For the next two quarters employed by you, the employee must not receive an occupational dose of greater than 2.5 rem

Note: This is a conservative calculation and diagnostic veterinary occupational staff is unlikely to receive an exposure of 2.5 rem over two quarters. This should not affect the occupational duties of the employee regarding the use of x-ray equipment.

ASSESSING THE NEED FOR INDIVIDUAL MONITORING

Veterinary registrants must perform an evaluation to assess the need for individual monitoring of all occupational workers including veterinary technicians, veterinarians, and other individuals assisting in the x-ray examinations. Minnesota Rules, Chapter 4732 does not exempt a registrant, individual or x-ray equipment use from the individual monitoring requirements of Minnesota Rules, Chapter 4732.0440.

The evaluation process may be different for each registrant and must include a review of your entire radiation safety program and how it is implemented. This review may include:

1. An evaluation of previous dose history records.

Past dosimetry reports are the easiest way for Veterinary registrants to verify and document the likelihood of receiving greater than 10% of the dose limits of Minnesota Rules, Chapter 4732.0410.

2. Provide individual monitoring to staff for a designated time, from 3 to 6 months.
   - Dosimetry badges can be submitted to the dosimetry provider on a monthly or quarterly basis
   - Designated time must be representative of the typical volume and type of imaging performed
   - Maintain records according to Minnesota Rules, Chapter 4732.0330 and Minnesota Rules, Chapter 4732.0415

3. Review similar practices and procedures representative of the typical volume and type of imaging being performed at your facility.
   - Other similar facilities
   - Patient workload
   - General radiography
   - CT
   - Fluoroscopy
   - C-arm use

4. All x-ray equipment uses within your veterinary practice must be included in the overall evaluation process.
5. Individuals who perform specific duties that include more extensive use of the x-ray equipment than other staff should be evaluated independently to ensure they are not likely to exceed the 10% annual dose. Examples include:
   • Individuals hired exclusively to perform fluoroscopic x-ray examinations
   • Individuals hired exclusively to perform portable x-ray examinations
   • Registrant’s who have received a variance approval and must meet the individual monitoring requirements of the variance approval

6. You must retain records of the evaluation process used to determine that dosimetry badges are not necessary.
SHIELDING REQUIREMENTS

You are responsible for protecting your staff and the public from unnecessary radiation. MDH has requirements for the design of an x-ray room/area to ensure that you have met the minimum protection requirements.

If you are a registrant that has x-ray rooms/areas that were constructed, structurally remodeled, or placed into use prior to February 2008, you are responsible for compliance with Minnesota Rules, Chapter 4732.0220, subpart 3.

If you are a registrant that has purchased a facility with existing x-ray rooms/areas that were constructed, structurally remodeled, or placed into use prior to February 2008, you are responsible for obtaining shielding documentation from the previous owner to ensure compliance with Minnesota Rules, Chapter 4732.0220, subpart 3.

If you are a registrant that has x-ray rooms/areas that were constructed, structurally remodeled, or placed into use after February 2008, you are required to complete and submit a shielding plan in accordance with Minnesota Rules, Chapter 4732.0360. See Shielding Plans Section on pg. 17 of this guide.

The requirements for registrants that had x-ray rooms/areas that were constructed, structurally remodeled, or placed into use prior to February 2008 were in Minnesota Rules Chapter 4730.1670 Subpart 1 which was adopted in 1993 and required a registrant to perform a radiation survey at the time of initial installation of x-ray equipment and after any change in the facility or equipment which might cause a change in radiation hazard.

Minnesota Rules, Chapter 4730.1670, subpart 1 states that each registrant conducting diagnostic or therapeutic x-ray procedures must ensure that the radiation safety surveys specified in this part are site-specific and in compliance with this chapter. A survey must be performed at the time of initial installation and after any change in the facility or equipment which might cause a change in radiation hazard. A report of each survey must be prepared, maintained at the facility according to the record requirements in Minnesota Rules, Chapter 4730.1520, and made available to the commissioner on request. The safety survey must include the following:

1. An evaluation of the tube housing integrity
2. Calibrations
3. Equipment performance measurements
4. Maintenance and equipment modifications
5. Shielding plans or results from radiation shielding evaluations

Minnesota Rules, Chapter 4732.0220, subpart 3 requires all registrants to maintain documentation of the radiation shielding installed in their facility. The documentation must include:

1. A blue print or architectural drawing indicating installed shielding.
2. A shielding plan that was completed by a service provider or an appropriate radiological physicist.
• The actual structural composition and thickness or lead equivalent of all walls, doors, partitions, and, if occupied spaces above or below, the floor and ceiling of the rooms concerned

• Shielding plans must be completed by a service provider registered in Minnesota or by a radiological physicist

**Note:** Shielding plans completed by any other individual will be not be accepted by MDH and do not comply with the shielding plan submission requirements.

3. Calculation to ensure occupational staff and the public do not receive a dose in excess of the following dose limits:

• Occupational, [Minnesota Rules, Chapter 4732.0410](#)

• Embryo or Fetus, [Minnesota Rules, Chapter 4732.0415](#)

• Exposure to minors, [Minnesota Rules, Chapter 4732.0420](#)

• Members of the public, [Minnesota Rules, Chapter 4732.0430](#) OR

4. If shielding compliance can’t be verified by all the above, a detailed radiation survey covering the radiation levels at the operator position and at pertinent points outside the room during normal operation must be completed.

**As defined in Minnesota Rules, Chapter 4732.0110, “Survey” or “radiation survey” means an evaluation of the radiological conditions and potential hazards incident to the use of radiation-producing equipment. When appropriate, such evaluation includes, but is not limited to, tests, physical examinations, and measurements of levels of radiation.**

**Note:** Dental Intraoral x-ray equipment is exempt from shielding requirements and a shielding plan is not required to be completed or submitted to MDH.

[Minnesota Rules, Chapter 4732.0355](#) establishes general shielding and operator booth requirements of the x-ray equipment. Due to the nature of the imaging, veterinary facilities typically do not incorporate the use of an operator’s booth. Facilities that possess computed tomography (CT) equipment must have a control booth in the design of the room. [See Shielding Plan Requirements for CT](#).

**SHIELDING PLANS**

MDH does not develop or approve radiation shielding plans. MDH does verify the following at the time of inspection:

1. A post construction radiation evaluation has been completed to ensure the room/area has been constructed according to the submitted shielding plan and MDH shielding
requirements to including: Room/area design, operator’s booth design requirements, viewing conditions, and shielding calculations

2. Corrective actions are taken when a post construction radiation evaluation or subsequent analysis of operating conditions indicate:
   - The possibility of individuals receiving a dose in excess of the dose limits prescribed in *Minnesota Rules, Chapters 4732.0410 - 4732.0430*
   - Non-compliance with the design of the room/area

3. Permanent Placards are in place.

*Minnesota Rules, Chapter 4732.0360* establishes requirements for shielding plans to be completed and submitted to MDH after February 5, 2008 for the following:

1. New construction of an x-ray room/area.
   - This includes rooms/areas that were not originally designed for x-ray use and are now used routinely or permanently
2. Structural remodel of an existing x-ray room/area.
   - Removal or remodeling of existing exterior walls, ceiling, floor, or control booth of a room or area
3. Shielding plans must be submitted to MDH prior to the new construction, remodel or use in rooms/areas that were not originally designed for x-ray use.
4. Shielding plan forms and additional information may be found on the MDH X-ray website in the *Topic Index, under Shielding*.

If your x-ray room/area was constructed prior to February 5, 2008, you were not required to submit a shielding plan to MDH. You were required to perform a radiation or evaluation of your room/area to ensure that staff and the public are protected from unnecessary radiation and the dose received are not in excess of the limits identified in this guide.

*When a shielding plan has been completed and submitted, a permanent placard must be mounted in the room/area identifying the amount and type of shielding in the room/area.*

**SHIELDING REQUIREMENTS FOR CT**

*Minnesota Rules, Chapter 4732.0860*, subparts 2 through 5 establish design, viewing, and radiation survey requirements for rooms where computed tomography (CT) equipment is used.

- **Subpart 2**: Requires the operator to remain in a permanently protected area and must meet the design requirements of *Minnesota Rules, Chapter 4732.0355*, subpart 4.
- **Subpart 3**: Requires that the operator must have a means to continuously observe the patient from the control panel during the examination.
- **Subpart 4**: Is not applicable to veterinary facilities.
- **Subpart 5**: Requires that all CT systems installed after February 5, 2008 must have a radiation survey performed to identify the radiation levels, at the control panel where the operator is located, and exterior rooms and locations outside of the CT room or area.
The other subparts of Minnesota Rules, Chapter 4732.0860 will be discussed throughout the guidance document.

PERMANENT PLACARD REQUIREMENTS

A placard must be mounted in the room/area specifying the amount and type of shielding in all walls, doors, partitions, and, if occupied, spaces above or below the floor and ceiling.

• If mounting the shielding information is not practical, you may post a notice in the room/area that describes the document and states where it may be examined
• The placard is to remain in the room/area until the room/area is destroyed or remodeled at which time a new shielding plan must be competed and submitted if the room/area is to be used for x-ray purposes
• The placard must be mounted in a manner that the placard cannot easily be removed, defaced or destroyed
• In the event the placard is removed, defaced or destroyed, you must replace the placard

1. Example placard when shielding information is placed directly on the placard.

```
Radiation Shielding for (Room Name/Number)
Control Room Wall contains (x/xx) inch lead
  North Wall contains (x/xx) inch lead
  East Wall contains (x/xx) inch lead
  South Wall contains (x/xx) inch lead
  West Wall contains (x/xx) inch lead
Ceiling contains (x) inches concrete on metal deck
Floor is concrete on slab (unoccupied)
```

2. Example placard where it may be impractical to place the shielding information directly on the placard.

```
Radiation shielding information for (Room Name/Number) is maintained in the radiation safety manual located in the Radiation Safety office. This information is available for review by contacting (provide name of RSO and contact number).
```
QUALITY CONTROL PROCEDURES

Routinely evaluating and maintaining your x-ray equipment, imaging system, image processing, darkroom and other associated components is important for maintaining the quality and stability of your x-ray imaging. In an effort to assist in understanding of the quality control tests required by MDH, quality control instruction sheets are provided in Attachment F.

You are not exempt from implementing a site specific quality assurance program or implementing quality control procedures to monitor the x-ray system on a routine basis to ensure stable and reliable performance of the x-ray equipment, imaging system and the imaging conditions.

Quality Control Procedures include:

<table>
<thead>
<tr>
<th>Quality Control Procedure</th>
<th>Rule Part</th>
</tr>
</thead>
<tbody>
<tr>
<td>Retake/Reject Analysis Program</td>
<td>Minnesota Rules, Chapter 4732.0535</td>
</tr>
<tr>
<td>Processing Equipment</td>
<td>Minnesota Rules, Chapter 4732.0555, subpart 1</td>
</tr>
<tr>
<td>Darkroom or Glove Box Fog Tests</td>
<td>Minnesota Rules, Chapter 4732.0555, subpart 3</td>
</tr>
<tr>
<td>Outdated Film</td>
<td>Minnesota Rules, Chapter 4732.0555, subpart 4</td>
</tr>
<tr>
<td>X-ray Equipment Calibrations and Corrective Actions</td>
<td>Minnesota Rules, Chapter 4732.0700</td>
</tr>
<tr>
<td>Computed Tomography (CT)</td>
<td>Minnesota Rules, Chapter 4732.0860</td>
</tr>
<tr>
<td>Veterinary Medical Radiographic Systems</td>
<td>Minnesota Rules, Chapter 4732.1100, subpart 9</td>
</tr>
<tr>
<td>Radiographic X-ray Equipment</td>
<td>Minnesota Rules, Chapter 4732.0875</td>
</tr>
<tr>
<td>Fluoroscopic X-ray Equipment</td>
<td>Minnesota Rules, Chapter 4732.1100, subpart 5</td>
</tr>
<tr>
<td>Dental Intraoral X-ray Equipment</td>
<td>Minnesota Rules, Chapter 4732.1100, subpart 6</td>
</tr>
</tbody>
</table>

If you perform general and dental imaging, you must ensure that the required testing and frequency of testing listed above is performed for all x-ray equipment, processors, and processing conditions. Corrective actions must be taken and verified to have corrected the out-of-limit parameters prior to using the x-ray equipment or processing patient images.

Notes: Veterinary facilities are exempt from performing a daily processor quality control evaluation; Minnesota Rules, Chapter 4732.0555 subpart 2, Processing Quality Control. This is an evaluation of the processing chemistry and requires the use of a sensitometer and densitometer and/or the use of a dental crabtree device.

Minnesota Rules, Chapter 4732.0275 requires individuals that assemble, install, repair or replace components of your x-ray equipment, perform install calibrations, or equipment performance evaluations on your x-ray equipment that are not employed by you, must be registered as service providers with the state of Minnesota prior to providing these services.

FILM OR FILM/SCREEN COMBINATIONS

You must ensure that films are developed according to the film and chemistry manufacturer’s recommendations. Minnesota Rules, Chapter 4732.0555 has specific requirements depending on your processing method:

Manual Processing:
• Films must be developed according to the film and chemistry manufacturer’s time-temperature recommendations
• Films must be developed according to the time-temperature requirements listed in Minnesota Rules, Chapter 4732.0555 subpart 1 when the manufacturer’s recommendations are not available
• The temperature of the developer must be checked prior to developing each set of films

Automatic Processing:
• Films must be developed according to the film and chemistry manufacturers’ time-temperature recommendations
• The developer must be checked daily when the processor does not have a developer temperature readout or a “ready light” indicating the developer temperature is within range
• The developer must be checked weekly when a processor has a developer temperature readout or a “ready light” indicating the developer temperature is within range

DARKROOM OR GLOVE BOX FOG TEST
You must ensure films are processed under conditions that minimize unnecessary film fogging. See Attachment F for Procedures for Fog Testing.

A darkroom or glove box fog test must be performed:
1. To confirm the correct safelight filters and safelight filter placement, visible white light leaks, and any other conditions are corrected which may cause film fogging
2. Initially prior to first use of the darkroom or glove box, and at intervals not to exceed six months
3. For all processing conditions, including:
   • General veterinary images
   • Dental veterinary imaging (glove box)
   • Any time there may be a change in the processing conditions, such as: a new safelight, a new bulb, or moving glove box to a new location

   Note: Corrective actions must be taken and verified to have corrected the out-of-limit parameters prior to using the darkroom or processing patient images.

DIGITAL IMAGING
Minnesota Rules, Chapter 4732.0835 states that you must follow the quality control recommendations of the manufacturer for the image receptor(s) when you have a computed radiography (CR), direct radiography (DR) or a photostimulable storage phosphor (PSP) system.

1. Calibrations of the digital x-ray system must be performed if this digital system includes new or replacement x-ray equipment.
2. Performance evaluations of your x-ray equipment must be performed at the appropriate interval listed in Calibrations/Performance Evaluations or anytime maintenance is performed on the x-ray unit or system.

**CONE BEAM COMPUTED TOMOGRAPHY (CBCT)**

Facilities using Cone Beam Computed Tomography (CBCT) must follow the quality control recommendations of the manufacturer.

- Performance evaluations of your x-ray equipment must be performed at the appropriate interval listed in Calibrations/Performance Evaluations or anytime maintenance is performed on the x-ray unit or system.

**COMPUTED TOMOGRAPHY (CT)**

In addition to the general requirements, facilities using CT must comply with the specific requirements found in Minnesota Rules, Chapter 4732.0860.

- Performance evaluations of your x-ray equipment must be performed at the appropriate interval listed in Calibrations/Performance Evaluations or anytime maintenance is performed on the x-ray unit or system.

**CALIBRATIONS/PERFORMANCE EVALUATIONS**

(4732.0280) (4732.0700) (4732.1100)

Calibrations/Performance Evaluations must be performed prior to first use of your x-ray equipment and anytime maintenance is performed on the x-ray unit or system.

Performance evaluation of your x-ray equipment must be performed:

1. At intervals not to exceed 24 months for the following:
   - Radiographic x-ray equipment
   - CBCT x-ray equipment
   - Dental x-ray equipment
   - Screen contact (one or more cassettes)
   - Speed match testing with two or more cassettes and cassettes of the same speed/type

2. At intervals not to exceed 12 months for the following:
   - Fluoroscopic x-ray equipment
   - Computed Tomography x-ray equipment
   - Anytime maintenance is performed on the x-ray equipment or system

A listing of the required tests can be found in Minnesota Rules, Chapter 4732.1100.

*Note: Individuals that assemble, install, repair or replace components of your x-ray equipment, perform calibrations, or performance evaluations on your x-ray equipment*
must be registered as a service provider with the state of Minnesota prior to providing these services per Minnesota Rules, Chapter 4732.0275.

LEAD APRON/PROTECTIVE GARMENTS INTEGRITY EVALUATIONS (4732.0550)

Lead aprons must be monitored for integrity initially and at intervals not to exceed 24 months. This requirement is to ensure the radiation protection quality of the lead within the apron has not been compromised.

UTILIZATION LOG (4732.0545)

Veterinary registrants must maintain a daily utilization log for all radiographic procedures performed. The utilization log must contain the following information:

1. Patient identification, including name, patient id, and other means to identify the patient
2. Type of procedure(s) performed on the patient
3. The number of images taken (including retakes)
4. The date the procedure was performed
5. The name of the individual performing the procedure
6. The name(s) of individuals required in the room to assist in the procedure
7. Fluoroscopic on time for fluoroscopy procedures over 5 minutes

RETAKE/REJECT ANALYSIS (4732.0535)

You must perform a quarterly analysis of all radiographic images, including dental imaging, retaken or rejected. The quarterly results must be reviewed during the annual audit. A retake/reject analysis is a valuable tool for reducing unnecessary radiation dose to occupational staff, improving image quality and providing important information about existing and reoccurring imaging issues.

An analysis may indicate the need for:

- Corrective action for x-ray equipment failures
- Corrective action to processing issues
- Additional staff training
- Adjustment to procedures and or techniques
- The need for specialized devices to assist in imaging

HAND-HELD X-RAY EQUIPMENT

The use of hand-held x-ray equipment in the state of Minnesota is unauthorized, per Minnesota Rules, Chapter 4732.0306. You must submit a variance request and receive approval from MDH.
for hand-held x-ray equipment use prior to performing activities requiring the use of hand-held x-ray equipment.

A variance may be granted only to rule and will not be issued to a non-compliant activity after that activity has taken place. When applying for a variance, the request must be submitted by the veterinary registrant and include the following information.

1. The specific language in the rule or rules from which the variance is requested
2. The reasons why the rule cannot be met
3. Alternative measures that will be taken to assure a comparable degree of protection to health or the environment
4. The length of time for which the variance is requested
5. A statement that the party applying for the variance will comply with the terms of the variance, if granted
6. Other relevant information the commissioner determines necessary to properly evaluate the request for the variance

MDH may attach alternative measures or conditions to a variance approval based on the conditions of use. Examples are listed below, the specific device would replace “hand-held x-ray equipment”:

- Operators of the “hand-held x-ray equipment” must receive training provided by the manufacturer prior to use
- Operators of the “hand-held x-ray equipment” must receive specific radiation safety training for its safe use at each location used
- The “hand-held x-ray equipment” must be calibrated according to Minnesota Rules, Chapter 4732.1100
- Each operator of the “hand-held x-ray equipment” must be assigned an individual monitoring device, and it must be worn at the neck level (outside of the apron)
- The “hand-held x-ray equipment” must be secured and inaccessible to untrained personnel when not under direct supervision

Additional information on variance statutory and regulatory requirements is below.

<table>
<thead>
<tr>
<th>Rule/Statute</th>
<th>Rule/Statute Description</th>
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</thead>
<tbody>
<tr>
<td>14.055</td>
<td>Minnesota Statute, Rule Variances; Standards</td>
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<tr>
<td>4717.7000</td>
<td>Minnesota Rules, Variance Request</td>
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<tr>
<td>4717.7010</td>
<td>Minnesota Rules, Criteria for Decision</td>
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<td>4717.7020</td>
<td>Minnesota Rules, Notification of Decision</td>
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<td>4717.7030</td>
<td>Minnesota Rules, Effect of Alternative Measures</td>
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<td>4717.7040</td>
<td>Minnesota Rules, Renewal of Variance</td>
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<tr>
<td>4717.7050</td>
<td>Minnesota Rules, Denial, Revocation or Refusal</td>
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</tbody>
</table>
RECORD RETENTION (4732.0330)

MDH X-ray Unit record retention requirements are limited to the use of x-ray equipment, operator of x-ray equipment qualifications, training, registration, and those records associated with the establishment and implementation of a radiation safety and quality assurance program.

You must, at a minimum, maintain all records for review between MDH X-ray Unit inspections. This would include:

- Training of the operators of x-ray equipment since the last inspection including present, float, and temporary staff
- Calibration and performance evaluations
- Quality control testing
- Corrective action for any quality control tests which may have failed
- Procedures for the Radiation safety/quality assurance program audit
- Radiation safety/quality assurance program audit
- Utilization log
- Quarterly Retake and Reject Analysis
- Evaluation for assessing the need for individual monitoring
- Individual monitoring (where applicable)
- Radiation Safety Officer Delegation Agreement
- Registration information
- Shielding Plan information
**INSPECTIONS**

MDH X-ray Unit inspection staff is responsible to perform inspections of facility operations to ensure the safe use of your x-ray equipment and compliance with [Minnesota Rules, Chapter 4732](#). During an inspection, inspection staff may perform confirmatory testing of your x-ray equipment, interview you and your staff, and review your radiation safety procedures, quality assurance procedures, and records. At the completion of the inspection, the inspector will perform an exit interview with you, your administrator, radiation safety officer, and registrant designee to discuss potential findings or concerns. The exit interview is to ensure the inspector’s review of your program is complete and accurate, and that there are no misunderstandings with potential findings or violations you may receive.

Inspections may be conducted initially for new registrants to review your radiation safety and quality assurance procedures and at the below subsequent inspection intervals.

<table>
<thead>
<tr>
<th>Registrant</th>
<th>Inspection Interval</th>
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<tr>
<td>Chiropractic</td>
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<tr>
<td>Computed Tomography (CT)</td>
<td>Every 3 years</td>
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<tr>
<td>Computed Tomography (CT)/Fluoroscopic</td>
<td>Every 3 years</td>
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<tr>
<td>Dental</td>
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<tr>
<td>Industrial</td>
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<tr>
<td>Medical (general)</td>
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<tr>
<td>Radiation Therapy</td>
<td>Every 2 years</td>
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<tr>
<td>Veterinary</td>
<td>Every 4 years</td>
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</table>

Inspections may be performed at an increased frequency based on previous enforcement history, failure to respond to corrective actions, or if MDH receives a call of concern regarding a registrant’s x-ray operations.

Registrants must allow MDH X-ray Unit inspection staff, during reasonable hours of operation, the opportunity to inspect the premises, x-ray equipment and records.
Radiation Safety Officer Training Requirements (4732.0500)

The individual designated as a radiation safety officer must be either a licensed practitioner of the healing arts or an individual who has completed training in the following items:

- Fundamentals of radiation safety
- Familiarization with facility’s radiation-producing equipment
- Film processing, if applicable
- Digital imaging, if applicable
- Quality assurance program
- Audits of the quality assurance program
- Emergency procedures for radiation-producing equipment failures
- Proper use of personal dosimetry, if applicable
- Requirements of pertinent Minnesota Rules, Chapter 4732
- Veterinary registrant’s written operating and emergency procedures

Typical Duties and Responsibilities of the Radiation Safety Officer (4732.0505)

The RSO’s duties and responsibilities include ensuring radiological safety and compliance with Minnesota Rules, Chapter 4732 and the conditions of the radiation safety/quality assurance program. Typically, the RSO’s duties and responsibilities include and are not limited to:

- Establishing a radiation safety/quality assurance program
- Maintenance, and implementation of up-to-date operating and emergency procedures
- Ensure initial site specific training has been performed for safe operating procedures, emergency procedures and quality control procedures
- Radiation Safety/Quality Assurance program audits are performed at intervals not to exceed 12 months
- Identifying radiation protection problems and developing, implementing, and documenting timely corrective actions
- Stopping unsafe activities using x-ray equipment
- Ensuring compliance with regulations
- Ensuring that radiation exposures are ALARA
ATTACHMENT B
FACILITY SPECIFIC TRAINING
(4732.0510)

Facility Specific Training
Each operator must be instructed initially in site-specific and system specific procedures including:

- Safe operating and Emergency procedures
- Quality control procedures for all imaging receptors, film and digital
- The use of proper protective shielding for staff
- Additional training must be conducted at the time of any change to the quality assurance program or change in radiation output. Examples include, but are not limited to:
  - Changing from film/screen to Computed Radiography (CR) or Direct Radiography (DR)
  - Replacement of or addition of a new x-ray unit

If your facility is in possession and uses fluoroscopic, cone beam computed tomography (CBCT), or computed tomography (CT) x-ray equipment, your staff must receive system specific training:

1. Fluoroscopic training must include:
   - X-ray generation and control
   - X-ray dosimetry
   - Image formation
   - Image acquisition
   - Image processing and management
   - Radiation effects
   - Dose-management fundamentals
   - Staff radiation safety
   - Professional standards and regulatory requirements
   - Other miscellaneous items appropriate to site-specific use

2. CT/CBCT training must include:
   - Training by the manufacturer or equivalent
   - Training in appropriate CT positioning and anatomy for procedures performed at the facility

Training requirements for students, float staff, externs and temporary staff:

- Students, externs, and float staff are required to perform the initial training at only one location if they remain within a veterinary system that has an established radiation safety/quality assurance program for all sites.
- Students and externs who train within different practices must receive training at each location of practice.
- Temporary staff working within different practices must receive training at each location of practice.
Record Retention Training
Documentation of training must be available onsite at each registered location, either in electronic or hard copy.

- Training records must include site specific and modality specific: date of training, topics covered and names/signature of trained individuals
# ATTACHMENT C
## ANNUAL AUDIT FOR VETERINARY X-RAY REGISTRANTS
(4732.0540)

Only address those areas that apply to your activities and activities that have not occurred since the last audit.

<table>
<thead>
<tr>
<th>Audit History</th>
<th>4732.0540</th>
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<td>Is the written agreement in place for the RSO?</td>
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<tr>
<td>All x-ray equipment registered with the MDH?</td>
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<td>Technique charts completed and in place?</td>
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<td>Lead apron procedures in place, and in use?</td>
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Retake and Reject analysis performed quarterly and an evaluation included in the annual audit

Digital manufacturer’s quality control procedures followed?

Have records been maintained?

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Summary of findings:

Corrective and preventive actions:

Comments:

Audit conducted by: ____________________________ Date: ____________________________
ATTTACHMENT D
CALIBRATIONS AND PERFORMANCE EVALUATION TESTS
(4732.1100)

Calibrations and performance evaluations are required in order for you to ensure your x-ray equipment is functioning in accordance with the specifications of the manufacturer, Minnesota Rules, Chapter 4732 or the Code of Federal Regulations, title 21, sections 1020.30 and 1020.31. If the manufacturer’s specifications are unknown the x-ray equipment must meet the specification of Minnesota Rules, Chapter 4732 or the Code of Federal Regulations, title 21, sections 1020.30 and 1020.31.

Calibrations and performance evaluations must be performed by a service provider who is currently registered with the MDH and the following information must be included on a calibration or performance evaluation report:

- The name and registration information of the service provider
- The numerical results for each test where applicable, and any test images
- Written recommendations necessary to bring x-ray equipment failures into compliance
- The date the equipment performance tests were completed
- The serial number of the equipment, room number, or name, if applicable

*Note: If the service provider is using the manufacturer’s specifications for compliance, the manufacturer’s specifications must be available onsite for review.*

The following tests are required at initial installation of the x-ray equipment, at the following required intervals and any time there is a change or replacement to the x-ray equipment where applicable. Performance evaluations are to be performed at all clinically used settings.

- **Beam Size (Dental x-ray equipment):** The area exposed to radiation must be limited to the area of interest and image receptor. It is very important to obtain the proper placement of the tube and image receptor to reduce the risk of clipping the area of interest and or exposing areas of the patient that is not to be imaged.
  1. Intraoral X-ray units:
     - If you are using a long cone veterinary unit the x-ray field size can be no larger than 2.75 inches (7cm) at the end of the cone.
     - If you are using a short cone veterinary unit the x-ray field size can be no larger than 2.36 inches (6cm) at the end of the cone.

- **Filtration or Half Value Layer (HVL):** This refers to “hardening” of the x-ray beam by filtering out the lower energy x-rays to allow for only the higher energy x-rays which are able to penetrate the bony structure and soft tissue to reach the patient and produce an image. Lower energy x-rays which cannot penetrate the bony structure or soft tissue add to the patient overall exposure but does not affect the x-ray image.

- **Radiation Exposure at the End of Cone (Dental Human Use):** MDH limits the maximum dose at the end of the cone on intraoral x-rays to reduce the patient dose. These maximums are based on the measured kVp of your x-ray unit and the speed of the film or imaging system you use. With
proper development of the image, optimal image quality can be obtained using less than the maximums established by your service provider.

- **Timer Accuracy:** Ensures the radiation exposure is consistent with the time that is set on the x-ray control. Timer setting sets the duration of the radiation exposure.

- **Timer Reproducibility:** This verifies the x-ray unit timer settings are accurate and will provide a reproducible exposure (density) of your image.

- **kVp accuracy:** Ensures the Kilovoltage Peak (kVp) set is consistent with the kVp that is set on the x-ray control. KVP controls the speed and energy of the x-rays and is the factor which allows the x-rays to penetrate the area of interest and provide the contrast to your images.

- **Reproducibility:** This verifies that the technique factors set (kVp, mA and timer setting) provide a reproducible exposure of your image.

- **Linearity:** This test is required only if your x-ray unit has multiple milliamperage (mA) settings that are used clinically. Milliamperage controls the quantity of electrons and with the timer setting the quantity of x-rays.

- **Dead man exposure switch:** When the x-ray exposure button is released the x-ray unit must stop producing an x-ray.

- **Audible and visible indication of an x-ray exposure:** There must be an audible and visible indication during the x-ray exposure.

- **Tube head stability:** The x-ray tube must remain stable during an x-ray exposure without the assistance of an individual or holding device.

- **Multiple tubes with one control:** A veterinary unit which operates more than one tube must have an indication on the x-ray control and on or near the tube housing assembly which has been selected.

- **SID Indicator Accuracy:** Ensures the actual distance at which the examination is performed is consistent with the distance noted on the technique chart.

- **X-ray to Light Field Alignment:** Ensures the area to be exposed to the x-ray field is congruent and the same size of the light field that is used to collimate to the patient area of interest.

- **X-ray to Image receptor Alignment:** Ensures the x-ray field is centered to the image receptor.
You may include the text as it appears here or if you prefer or you may develop your own ALARA (As Low as Reasonably Achievable) program for MDH review at the time of an inspection.

Management Commitment

- We, the management of this facility, are committed to the program described herein for keeping individual and collective doses As Low as Reasonably Achievable (ALARA). In accord with this commitment, we hereby describe an administrative organization for radiation safety and will develop the necessary written policy, procedures, and instructions to foster the ALARA concept within our institution. The organization will include a radiation safety officer.

- We will perform a formal annual review of the radiation safety program, including ALARA considerations. This will include reviews of operating procedures, past dose records (if applicable), inspections, etc., and any modifications to operating and maintenance procedures or to x-ray equipment and facilities will be reviewed and include consultations with the radiation safety staff or outside consultants.

- In addition to maintaining doses to individuals as far below the limits as is reasonably achievable, the sum of the doses received by all exposed individuals will also be maintained at the lowest practicable level.

Registrant’s Responsibility to Supervised Individuals

- Veterinary registrants will explain the ALARA concept and the need to maintain exposures as low as reasonably achievable to all staff.

- Veterinary registrants will ensure that the supervised individuals who are subject to occupational radiation exposure are trained and educated in safe radiation practices involving time, distance, shielding, and appropriate techniques in maintaining exposures.

Veterinary registrants will ensure staffs that are subject to occupational radiation exposure are trained and educated in practices involving time, distance, shielding and appropriate techniques in maintaining exposures ALARA.
In this Attachment you will find suggested methods for performing the applicable quality control testing required in a veterinary facility using x-ray equipment. Included in these attachments are procedures and guidance documents for the following quality control tests:

PROCEDURES FOR REPEAT/RETAKE ANALYSIS
PROCEDURES FOR FOG TESTING
PROCEDURES FOR INTRAORAL FOG TESTING
PROCEDURES FOR EXTRAORAL SCREEN CONTACT TEST
PROCEDURES FOR SCREEN SPEED MATCH TESTING
PROCEDURES FOR ADDING OR CONVERTING TO PSP OR DIGITAL IMAGING SYSTEM IN AN OFFICE
PROCEDURES FOR LEAD APRON INTEGRITY EVALUATION

INITIAL REGISTRATION
ADDITIONAL REGISTRATION
SAMPLE RADIATION SAFETY OFFICER DELEGATION AGREEMENT
SAMPLE UTILIZATION LOG
SAMPLE REPEAT/REJECT ANALYSIS WORKSHEET
PROCEDURES FOR REPEAT AND REJECT ANALYSIS

What is it?
Repeat and Reject Analysis is a mechanism for tracking and evaluating images that have to be repeated or are rejected.

Why is it important?
It is a tool for improving image quality and reducing unnecessary occupational dose by tracking and evaluating poor images that may be caused, for example; malfunctioning x-ray equipment, changes in procedure and staff that require additional raining, or the need for specialized devices to assist in imaging.

When is it performed?
Repeat and Reject Analysis is an ongoing process where the repeated or rejected images are tracked on your daily utilization log with an analysis performed quarterly. An annual review of your repeat and reject analysis must also be reviewed by the RSO during the annual audit.

What is the requirement?
Minnesota Rules, Chapter 4732.0535 requires the repeat and reject analysis be:
- Performed at least quarterly
- Include the overall retake or reject rate and a summary of the causes for the retakes or rejects
- A review of your repeats and rejects during the annual audit
- Develop written facility specific procedures for the retake and reject analysis
- Maintain records according to Minnesota Rules, Chapter 4732.0330

Items needed
- Utilization log from the previous quarter
- Repeat and Reject analysis form

Procedure
1. Review your daily utilization log for the previous quarter.
2. Document on your Repeat and Reject Analysis Form the number of repeated or rejected films corresponding to the reason for the repeat or reject.
3. Calculate the total number of repeated or rejected images.
4. Calculate the total number of images taken in the quarter.
5. Use the one of the following formulas for calculating the repeat or reject rate:
   - Repeat Rate = Total Number Repeats for the Quarter / Total Number of Films Taken
   - Repeat Rate = Total Number Repeats for a specific exam in the Quarter / Total Number of Films Taken

Corrective Actions
MDH does not have a maximum percentage rate for the repeat and reject analysis. It is the RSO’s responsibility to review these rates and determine what, if any, corrective action should be taken. All corrective actions must be documented and retained in your Radiation Safety/Quality Assurance Manual for future reference. Changes in staffing, x-ray equipment or from film to digital imaging as examples where the repeat and reject rate may be higher than what is typically seen at your facility.
PROCEDURES FOR FOG TESTING

What is it?
The darkroom fog test is meant to determine and minimize the amount of unwanted light within the darkroom. This test is not required for CR, DR, or PSP imaging systems.

Why is it important?
Improper safelights and unwanted light in the darkroom can compromise the quality of your radiographs by reducing contrast and darkening the image. This can jeopardize image quality to the point of repeating the image or misdiagnosis.

When is it performed?
Fog testing must be performed initially, at intervals not to exceed 6 months, and any time there is a change in the darkroom conditions that may have the potential for unnecessary light to affect the quality of your images.

What is the requirement?
Minnesota Rules, Chapter 4732.0555 requires the darkroom/glove box test be performed:
- Initially and at intervals not to exceed six (6) months
- Anytime fog is suspected
- Anytime there is a filter or bulb change
- Any other change in darkroom conditions

The amount of fog for a two-minute test must not allow visualization of a density difference between the covered and uncovered side of the fog test film. Minnesota Rules, Chapter 4732.0330 requires records be maintained for review by the X-ray unit.

Items needed
- Loaded imaging cassette (preferably the smallest cassette used in your practice)
- Your x-ray unit
- Established technique factors for the darkroom fog test
- Timer set at 2 minutes

Procedure
1. Load the cassette with unexposed film under your normal darkroom conditions.
2. Take the loaded cassette into your x-ray room and place on the patient table.
3. Set a distance typically used for imaging (40 inches to table top or the cassette tray).
4. Collimate to about a 6” by 8” field size on the cassette.
5. Set a technique that would be used for a small animal extremity:
   - For example 50 kvp at 1 mAs.
   - You will have to adjust your techniques according to you specific film system.
• The processed film should have a density that is light enough to visualize the writing on a newspaper, and yet dark enough to make the writing difficult to read. 1.0 optical density if a densitometer was to be used.

6. Take the cassette into the darkroom and under the same conditions that would be used for processing patient films.
7. Place the cassette in the location most used for unloading and loading during processing.
8. Remove the film from the cassette and cover half of the film lengthwise with something that is light opaque. A cassette works well.
9. Etch a line in the film along the edge of the light opaque object.
10. Start the 2 minute timer.
11. Stand back from the film to ensure your body is not shadowing the fog test film. Take time to look around the darkroom for any potential light leaks or sources of unwanted light.
12. When the timer goes off, process the film as usual.
13. Take the processed film to a view box and review the density on the side of the film that was covered with the density on the side of the film that was uncovered.
14. If the densities between the covered and uncovered side are significant your fog test fails. Greater than a 0.08 optical density if an densitometer was to be used for measurement.
15. Corrective action must be taken and another fog test must be performed to verify the corrective action was acceptable.
16. Record the date, the results of the test as pass/fail, and save the film for state inspection.

Helpful Hints
Below are some common conditions as to why the fog test may fail.

Safelight/filter:
• Not compatible with the film being used
• Bulb in the safelight is too high a wattage
• Cracks in the filter
• Filter emulsion flaking off

Electronic equipment indicator lights:
• Radio
• Internet modems
• Phone

Additional conditions:
• Flames from boilers, water heaters or furnaces
• Ceiling tiles that are not installed correctly
• Light leaks around ceiling fixtures or doors

Corrective Actions
Repeating a fog test without the safelight on and the fog is removed, the safelight may need to be replaced or moved further away from the processor. Any light other than what is from the safelight can potentially fog your patient films. Remove or completely cover any of these sources of unwanted light:
• Close cupboards or place items behind a curtain
• Place a curtain covering the entire darkroom door entrance and use a curtain rod or hooks to move the curtain out of the way when film processing is not being performed
• Tape around light leaks in the ceiling
• Attach weather stripping around the darkroom door
PROCEDURES FOR INTRAORAL DENTAL FOG TEST

What is it?
The glove box (daylight processor) fog test is meant to determine and minimize the amount of unwanted light within the glove box.

Why is it important?
Unwanted light in the glove box can compromise the quality of your radiographs by reducing contrast and darkening the image. This can jeopardize image quality to the point of repeating the image or misdiagnosis.

When is it performed?
Fog testing must be performed initially, at intervals not to exceed 6 months, and any time there is a change in the glove box conditions that may introduce the potential for unnecessary light to affect the quality of your images.

Note: If you use the same darkroom conditions for processing intraoral film as you do for general radiography films, you do not need to perform this test. Your general radiography film is your most sensitive film.

What is the requirement?
Minnesota Rules, Chapter 4732.0555 requires the glove box test be performed:

- Initially and at intervals not to exceed six (6) months
- Anytime fog is suspected
- Anytime there is a filter or bulb change
- Any other change in darkroom conditions

The amount of fog for a two-minute test must not allow visualization of the outline of a coin on the intraoral film.

Minnesota Rules, Chapter 4732.0330 requires records be maintained for review by the X-ray unit.

Items Needed

- Timer Set for two minutes
- Coin
- Unexposed Intraoral Dental Film Packet

Procedure

1. Place all of the items needed for this test in the glove box.
2. Ensure fog test is performed using the same processing conditions that are used for processing patient films.
3. Under your normal processing conditions unwrap the film from the film packet and place the film in glove box. You may need to place the film on a covered developer, fixer or water cup.
4. Place the coin on top of the film.
5. Start the 2 minute timer.
6. Keep your hands in the cuffs and lean to ensure your body is not shadowing the viewing window.

7. Evaluate the condition of the cuffs and any seals for potential light leaks.

8. When the timer goes off, process the film as usual.

9. Review the film to ensure it passes.
   - Figure 1 shows a passing fog test.
   - Figure 2 shows a failing fog test. If any difference between the covered and uncovered portions is seen, it may indicate the presence of darkroom fog. See corrective actions below.

10. If the fog test fails, corrective action must be taken and another fog test must performed to verify the corrective action was acceptable.

11. Record the date, the results of the test as pass/fail, and save the film for state inspection.

Helpful Hints
### Common conditions why the fog test may fail:
- Glove box is placed under direct fluorescent lights
- Glove box cuffs are worn and fit loosely around the wrists
- Filter cover may be damaged or is not compatible with film used

Corrective Actions
- If a daylight processor glove box fog test fails, try covering the viewing window filter when redoing the fog test to see if this is the source of fog. If that is the case, moving the processor to a different location or changing surrounding lighting conditions may help to remove the fogging conditions.
- Replace cuffs that are loose fitting.
PROCEDURES FOR FILM/SCREEN CONTACT TEST

The film/screen contact test is not required for digital imaging systems

What is it?
The screen contact test is used to confirm there is good contact between the screens and the film inside of the x-ray cassette and must be performed on all x-ray cassettes used clinically. Repeated expose to x-rays does not cause x-ray screens to wear out. Typically the cause for poor contact requiring replacement of the screen(s) is due to improper maintenance and handling. Be sure to follow the manufacturer’s recommendations for cleaning and care.

Why is it important?
Poor contact between the screen and the film inside of an x-ray cassette can cause an x-ray image to look blurred, density fluctuations throughout the film, and artifacts which may reduce the diagnostic quality of your patient films and add unnecessary radiation dose to your patients if the films must be repeated.

When is it performed?
The screen contact test must be performed initially prior to patient use, at intervals not to exceed twenty four (24) months and any time there has been a change to the cassette that may affect the film/screen contact including new hinges, felt padding or screen(s). This is the same frequency as the calibration/performance evaluations of your x-ray equipment and it may be of value to have the service provider perform this test for you.

What is the requirement?
Minnesota Rules, Chapter 4732.1100 requires the screen contact test to be performed:
- Initially (new) and at intervals not to exceed twenty four (24) months
- Anytime screen damage is suspected
- Minnesota Rules, Chapter 4732.0330 requires records be maintained for review by the X-ray unit

Items Needed
- 8 wire/inch mesh test tool or 7 holes per inch test tool
- All imaging cassettes. Each cassette must be identified along with the respective test film.
- View box

Procedure
1. Load with film each imaging cassette under your normal darkroom/glove box conditions allowing them to sit for at least 15 minutes after loading. This will give any air trapped in the cassettes time to dissipate.
2. Take the loaded imaging cassette into the x-ray room.
3. Rotate the x-ray tube so that the x-ray beam is pointed towards the floor or x-ray table.
4. Place the cassette on the floor or x-ray table underneath the x-ray tube.
5. Place the screen contact test tool on top of the cassette.
6. The cassette should be placed on the floor or x-ray table with the tube at a distance of at least 40”. This will provide enough distance from the tube to the cassette to allow the x-ray field to cover the entire cassette.
7. Expose the test tool and cassette using approximately the same setting you used for your fog test evaluation, Optical Density (O.D.) of approximately 1.0.
8. Take each cassette into the darkroom and process the film under your normal processing conditions.

MDH Veterinary X-ray Regulatory Guide
9. View each processed film on a view box in a dimly lit room from a distance of approximately six feet or more.
10. Look for areas that are darker and/or more blurry than the rest of the film. This indicates poor contact.
11. If there is an area of poor contact located in an area of interest on a film, remove the cassette from service.

**Helpful Hints**
Some common causes of poor screen-film contact:
- Worn felt behind the screen(s)
- Loose, bent or broken hinges or latches
- Warped screens or cassettes
- Sprung or cracked cassette frame
- Foreign matter under the screen

X-ray cassettes and screen will last indefinitely when they are properly handled and maintained by following the manufacturer’s recommendations.

*What the test tool looks like...*  
*What your films may look like...*
PROCEDURES FOR SCREEN SPEED MATCH TESTING

What is it?
The screen speed match test is performed to confirm there is a consistent image density from one cassette to another and must be performed on all x-ray cassettes used clinically.

Why is it important?
The screen speed match test is used to ensure that the effective film density remains consistent from one cassette to another at a given technique. If you use a number of cassettes interchangeably, you need to be aware of the density for the combination of film and screen that you're using and you need to perform a proper speed match test to make sure that each of the interchangeable cassettes produces the same effective density. An adjustment in the technique may be required for cassettes that do not provide a similar density.

When is it performed?
The screen speed match test must be performed initially prior to patient use, at intervals not to exceed twenty four (24) months and any time there has been a change to the cassette that may affect the screen speed. This is the same frequency as the calibration/performance evaluations of your x-ray equipment and it may be of value to have the service provider perform this test for you.

What is the requirement?
Minnesota Rules, Chapter 4732.1100 requires the screen speed match test to be performed:

- Initially and at intervals not to exceed twenty four months
- Anytime screen damage is suspected

Minnesota Rules, Chapter 4732.0330 requires records be maintained for review by the X-ray unit.

Items needed
- All cassettes used clinically
- Identifier for each cassette and processed film, such as paperclips or lead markers
- Technique used for your darkroom fog test evaluation
- View box

Procedure

1. Select a small cassette to be used as a “master” cassette. Make a note of the cassette number or some other identifier for the master cassette, and record it. 
   *Ex: Cassette #40 is the master.*
2. Load the master cassette and all other cassettes with film from the same box of film. Place the film in the corner that will be exposed on larger cassettes. Remember what corner of the large cassette the film is in.
3. Set a distance of 40” to the table top.
4. Place three cassettes including your master cassette on your x-ray table so that one corner of each cassette touches each other making sure the film is in the corner you select on the larger cassettes.
5. Center the x-ray field to the center where the four cassette corners meet and come down so that you are exposing an area of approximately 4 inches by 4 inches on each cassette. *See arrows on the picture.*

6. Make an exposure using a similar technique that you use for your darkroom fog test evaluation. A paw or carpus technique on a small animal.

7. Take the cassettes into the darkroom and process the films under your normal processing conditions.

8. Place the processed films on a viewbox similar to how the cassettes were placed on the table top.

9. Visually inspect the films for significant density differences between the films.

10. If a film shows a significant density difference, remove the associated cassette from service. If you are reading the difference with a densitometer, the difference must be less than 0.10 Optical Density.

11. If you have more than four cassettes, you must repeat steps 2 through 9 using the same master cassette with the additional cassettes. *Ex: Cassette #40.*

12. Save all associated films and documentation until the next inspection by the state.

**Helpful Hints**

Some common causes of screen speed match failures:

- Tested cassette screens are not of the same speed
- Film and screens are not compatible
- Incorrect cleaner used on screens

X-ray cassettes and screens can have a long service life as long as they are properly handled and maintained following the manufacturer’s recommendations.
PROCEDURES FOR ADDING OR CONVERTING TO A DIGITAL IMAGING SYSTEM

The exposure of an individual for training, instruction, demonstration, or maintenance is prohibited. The use of x-ray equipment for these purposes will result in enforcement action that may include an administrative penalty of up to $10,000.

What is it?
Computed Radiography (CR), Direct Radiography (DR) and Photostimulable Storage Phosphor (PSP) imaging are the manner in which the x-rays are received and processed to provide for a diagnostic image.

• Many registrants converting x-ray units to a CR or PSP imaging system may only replace the film imaging cassettes with CR or PSP imaging cassettes.
• Regardless of the imaging system an x-ray tube is necessary and the patient must receive radiation in order to generate the image.
• CR and PSP imaging systems require the imaging cassette to be placed in an image reader to obtain the x-ray image.
• For digital imaging systems, the x-ray image is obtained directly from the sensor and received on the computer monitor without the need for an image reader.

Why is it important?
1. Digital imaging does not require processing of the image in the same manner as film.
2. Dose reduction is not an “automatic” when converting to digital imaging.
   • Film/screen imaging is not forgiving, what you see is what you get
   • Digital imaging allows for manipulation of the raw image to enhance the final image
3. Doses may be higher during the transitional stages from film to digital imaging.
4. The “dose creep” phenomenon or the theory that “this technique works for everyone” is the tendency to overexpose a patient in order to maintain the ability to manipulate the raw image.
   • Overexposing allows for the ability to enhance the images through adjustment of the window and level post exposure
   • Underexposing does not allow for the ability to enhance the images and increases the noise of the images
   • Proper exposure allows for the enhancement of the images while maintain the dose as low as reasonable
   • Exposure is reduced with the use of Automatic Exposure Control (AEC)

What you must do?
• Submit an email to the X-ray Unit health.xray@state.mn.us stating that you have gone digital.
• Retain the email for your records.
• When installing new x-ray equipment in your digital conversion, the service provider must complete an installation calibration.
• Work closely with the service provider to give you the best image quality and maintain the patient dose as low as possible and adjust your technique charts accordingly.
• The service provider or you must adjust the preprogrammed techniques if they are to be used.
• Review your digital technical manuals very carefully, including:
  1. Manufacturer’s specifications for maintenance and quality control of the digital imaging system must be maintained onsite

MDH Veterinary X-ray Regulatory Guide 48
2. Maintenance and quality control testing of the digital imaging system must be performed according to manufacturer’s specifications

- Training must be done at the time of conversion and documented for all those who operate the digital system to ensure staff is aware of new exposure techniques, proper equipment usage, including use of holders, and equipment maintenance and quality control requirements.
- Update your Radiation Safety/Quality Assurance Manual to include procedures for the use digital imaging.
- Ensure you have updated your techniques for all imaging receptors in use.
PROcedures for Lead Apron Integrity Evaluation

What is it?
The Lead Apron Integrity Evaluation is an evaluation of the lead inside of the lead aprons, half-aprons, gloves and thyroid collars has maintained its protective characteristics.

Why is it important?
Lead aprons, half-aprons, gloves, and thyroid collars protect the staff and the public (when needed) from unnecessary radiation exposure during radiology procedures. The Lead Apron Integrity Evaluation ensures that lead aprons, half-aprons, gloves, and thyroid collars provide an ideal level of protection against radiation exposure.

When is it performed?
The Lead Apron Integrity Evaluation must be performed initially prior to patient use, at intervals not to exceed twenty four (24) months

What is the requirement?
Minnesota Rules, Chapter 4732.0550 requires lead apron integrity evaluations to be performed
• Initially and at intervals not to exceed twenty four months
• Anytime damage is suspected
• Minnesota Rules, Chapter 4732.0330 requires records be maintained for review by the X-ray unit

Note: Lead apron integrity evaluation must be performed for all personal protective garments regardless of the shielding material they are made of. The following procedures specify personal protective garments made with lead, but may be used for the different types of shielding material.

Items Needed
• All lead aprons, half-aprons, gloves and thyroid collars in use.
• Identifier for each personal protective garment in use
• Technique of ~ 70 kVp and 10 mAs.

Procedure
Film and CR Imaging

1. Lay the lead apron as flat as possible on the table top lengthwise. (Remove any wrinkles)
2. Place a loaded 14” X17” cassette crosswise in the table bucky (cassette tray) and position the cassette and x-ray field over the chest area of the lead apron
3. Collimate to the cassette
4. Setting a technique of approximately 70 kVp and 10 mAs, make an exposure
5. Using a second loaded cassette place it lengthwise and position the cassette and x-ray field over the abdomen and pelvis area of the lead apron
6. Make another exposure
7. Process the images as normal
MDH X-ray Unit is always striving to better the information that we provide to registrants. This may include additions to the information presented in this guide. There may be occasion for revisions to this guide. **THESE REVISIONS ARE NOT CHANGES TO MINNESOTA RULES, CHAPTER 4732** and are intended to clarify or supplement what is already within the guide. Any revisions to this guide will be documented in the **SUMMARY OF REVISIONS**.

<table>
<thead>
<tr>
<th>REVISION</th>
<th>SECTION</th>
<th>DESCRIPTION</th>
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<tbody>
<tr>
<td>03/2016</td>
<td>QC Procedures</td>
<td>Updated procedures for adding or converting to a digital imaging system.</td>
</tr>
<tr>
<td>03/2016</td>
<td>QC Procedures</td>
<td>Updated procedure for film/screen contact test.</td>
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<tr>
<td>03/2016</td>
<td>Throughout guidance</td>
<td>Revised grammatical and formatting errors.</td>
</tr>
</tbody>
</table>
| 03/2016  | Shielding Requirements    | Clarified shielding and shielding documentation requirements for registrants that have or have purchased facilities that were constructed before February of 2008.
Delegation of Authority for a Radiation Safety Officer for an X-ray Facility

(Please retain for your records)

Facility Name: __________________________________________________________

Facility Registration Number: _______________________________________________

Memo To: Radiation Safety Officer

From: Chief Executive Officer

Subject: Delegation of Authority

You, ________________________, have been appointed Radiation Safety Officer for our x-ray department. You are responsible for ensuring the safe use of radiation. Your responsibilities include managing the radiation protection program, identifying x-ray radiation protection problems, ensuring quality control tests are completed and documented, recommending, or providing corrective actions; verifying implementation of corrective actions; stopping unsafe activities; and ensuring compliance with state regulations.

You are hereby delegated the time and authority necessary to meet those responsibilities, including prohibiting the use of radiation-producing equipment by employees who do not meet the necessary requirements and shutting down operations where radiation safety is compromised. You are required to notify management if staff do not cooperate and do not address radiation safety issues. In addition, you are free to raise issues with the Minnesota Department of Health at any time.

It is estimated that you will spend _______ hours per week conducting radiation protection activities.

Your signature below indicates acceptance of the above responsibilities.

Name of Radiation Safety Officer  Name of Management Representative

______________________________  _________________________________
Signature of Radiation Safety Officer  Signature of Management Representative

______________________________  _________________________________
Date  Date

cc: Department Heads
## Patient Utilization Log

<table>
<thead>
<tr>
<th>Date</th>
<th>Patient ID</th>
<th>Exam</th>
<th># of Films</th>
<th># of Retakes</th>
<th>Retake Reason</th>
<th>Operators</th>
<th>Holding Assistants</th>
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</table>

### Retake Reason

1. Positioning
2. Technique
3. Too light
4. Too dark
5. Motion
6. Jewelry
7. Artifact
8. Mechanical
9. Static
10. Fog
11. Other

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Radiation Control, X-ray Unit
625 North Robert Street
PO Box 64497
St. Paul, Minnesota  55164-0497
651-201-4545
[www.health.state.mn.us/xray](http://www.health.state.mn.us/xray)
health.xray@state.mn.us
Repeat/Reject Analysis Worksheet

Time Period: ____________ to ____________
Total # Images Used: ____________

<table>
<thead>
<tr>
<th>Cause</th>
<th>Number of Images</th>
<th>Total Number</th>
<th>Total Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image - Black</td>
<td></td>
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<tr>
<td>Image - Dark</td>
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<td>Image - Good</td>
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<tr>
<td>Image - Light</td>
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<tr>
<td>Fog - Cassette</td>
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<tr>
<td>Fog - Darkroom</td>
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<tr>
<td>Mechanical</td>
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<tr>
<td>Other</td>
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<tr>
<td>Patient Motion</td>
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</tr>
<tr>
<td>Positioning</td>
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<tr>
<td>Static</td>
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</tbody>
</table>

The analysis must include at a minimum the overall retake or reject rate, and a summary of causes for the retakes. Include corrective actions if needed.

Repeat Rate = \( \frac{\text{Total Number Repeats}}{\text{Total Images Taken}} \) \times 100

Quarterly Repeat Rate _________

Optional: Individual Cause Repeat Rate = \( \frac{\text{Total Number Repeats for That Cause}}{\text{Total Number of Repeats}} \) \times 100