The American College of Radiology, with more than 30,000 members, is the principal organization of radiologists, radiation oncologists, and clinical medical physicists in the United States. The College is a nonprofit professional society whose primary purposes are to advance the science of radiology, improve radiologic services to the patient, study the socioeconomic aspects of the practice of radiology, and encourage continuing education for radiologists, radiation oncologists, medical physicists, and persons practicing in allied professional fields.

The American College of Radiology will periodically define new practice parameters and technical standards for radiologic practice to help advance the science of radiology and to improve the quality of service to patients throughout the United States. Existing practice parameters and technical standards will be reviewed for revision or renewal, as appropriate, on their fifth anniversary or sooner, if indicated.

Each practice parameter and technical standard, representing a policy statement by the College, has undergone a thorough consensus process in which it has been subjected to extensive review and approval. The practice parameters and technical standards recognize that the safe and effective use of diagnostic and therapeutic radiology requires specific training, skills, and techniques, as described in each document. Reproduction or modification of the published practice parameter and technical standard by those entities not providing these services is not authorized.

Revised 2019 (CSC/BOC)*

ACR–AAPM TECHNICAL STANDARD FOR MEDICAL PHYSICS PERFORMANCE MONITORING OF IMAGE-GUIDED RADIATION THERAPY (IGRT)

PREAMBLE

This document is an educational tool designed to assist practitioners in providing appropriate radiologic care for patients. Practice Parameters and Technical Standards are not inflexible rules or requirements of practice and are not intended, nor should they be used, to establish a legal standard of care. For these reasons and those set forth below, the American College of Radiology and our collaborating medical specialty societies caution against the use of these documents in litigation in which the clinical decisions of a practitioner are called into question. The ultimate judgment regarding the propriety of any specific procedure or course of action must be made by the practitioner considering all the circumstances presented. Thus, an approach that differs from the guidance in this document, standing alone, does not necessarily imply that the approach was below the standard of care. To the contrary, a conscientious practitioner may responsibly adopt a course of action different from that set forth in this document when, in the reasonable judgment of the practitioner, such course of action is indicated by variables such as the condition of the patient, limitations of available resources, or advances in knowledge or technology after publication of this document. However, a practitioner who employs an approach substantially different from the guidance in this document may consider documenting in the patient record information sufficient to explain the approach taken.

The practice of medicine involves the science, and the art of dealing with the prevention, diagnosis, alleviation, and treatment of disease. The variety and complexity of human conditions make it impossible to always reach the most appropriate diagnosis or to predict with certainty a particular response to treatment. Therefore, it should be recognized that adherence to the guidance in this document will not assure an accurate diagnosis or a successful outcome. All that should be expected is that the practitioner will follow a reasonable course of action based on current knowledge, available resources, and the needs of the patient to deliver effective and safe medical care. The purpose of this document is to assist practitioners in achieving this objective.

1 Iowa Medical Society and Iowa Society of Anesthesiologists v. Iowa Board of Nursing 831 N.W.2d 826 (Iowa 2013) Iowa Supreme Court refuses to find that the ACR Technical Standard for Management of the Use of Radiation in Fluoroscopic Procedures (Revised 2008) sets a national standard for who may perform fluoroscopic procedures in light of the standard’s stated purpose that ACR standards are educational tools and not intended to establish a legal standard of care. See also, Stanley v. McCarver, 63 P.3d 1076 (Ariz. App. 2003) where in a concurring opinion the Court stated that “published standards or guidelines of specialty medical organizations are useful in determining the duty owed or the standard of care applicable in a given situation” even though ACR standards themselves do not establish the standard of care.
I. INTRODUCTION

This technical standard was revised collaboratively by the American College of Radiology (ACR) and the American Association of Physicists in Medicine (AAPM).

This technical standard is intended to provide guidance for quality assurance (QA) of systems used for external-beam image-guided radiation therapy (IGRT) (or image guidance) and adaptive radiation therapy (ART). IGRT is a general term addressing the imaging application in the entire process of radiation therapy. However, in this document, IGRT discussion is limited to only imaging application in the treatment room.

The goal of radiation therapy treatment is to deliver radiation to the planned target volumes as precisely and accurately as possible while minimizing dose to critical normal tissues. IGRT is the process of using various imaging techniques to locate target and critical tissues and, if needed, reposition the patient for the delivery of radiotherapy. Image registration techniques are used to determine the amount of adjustment of the patient and therefore the target relative to the treatment beam. The basic imaging tools employed in IGRT have remained essentially the same since the publication of the ACR–AAPM Technical Standard on IGRT in 2014. However, advancements in IGRT techniques have seen a paradigm shift toward ART in which image guidance is coupled with image registration and replanning to generate an “updated” treatment plan that is “adapted” to the current patient anatomy if it is sufficiently different than that at the time of the initial planning.

II. QUALIFICATIONS AND RESPONSIBILITIES OF QUALIFIED MEDICAL PHYSICIST

A Qualified Medical Physicist is an individual who is competent to practice independently one or more of the subfields in medical physics. The American College of Radiology (ACR) considers certification, continuing education, and experience in the appropriate subfield(s) to demonstrate that an individual is competent to practice one or more of the subfields in medical physics and to be a Qualified Medical Physicist. The ACR strongly recommends that the individual be certified in the appropriate subfield(s) by the American Board of Radiology (ABR), the Canadian College of Physics in Medicine, the American Board of Science in Nuclear Medicine (ABSNM), or the American Board of Medical Physics (ABMP).

A Qualified Medical Physicist should meet the ACR Practice Parameter for Continuing Medical Education (CME). [1]

The appropriate subfield of medical physics for this technical standard is Therapeutic Medical Physics (previous medical physics certification categories that include Radiological Physics and Therapeutic Radiological Physics are also acceptable). (ACR Resolution 17, adopted in 1996 – revised in 2008, 2012, 2022, Resolution 41f)

With the advancement of imaging technologies and new techniques being developed on IGRT, a Qualified Medical Physicist may not have the necessary experience on a particular IGRT technique and must be trained accordingly before implementing or performing the new IGRT technique in the clinic.

The Qualified Medical Physicist is responsible for the technical aspects of IGRT. Those responsibilities should be clearly defined and should include the following:

1. Acceptance testing and commissioning of the IGRT system, thereby ensuring its mechanical, software, and geometric precision and accuracy as well as its image quality verification and documentation. These generally include, but are not limited to:
   a. Communication with the treatment planning system
   b. Communication with the treatment delivery system
   c. Evaluation of adequate image quality and imaging dose
   d. Testing of image registration software and application of results to patient shift coordinates
   e. Communication of patient data between storage, retrieval, and display devices
2. Implementation and management of a QA program for the IGRT system to monitor and ensure each of the following:
   a. The geometric relationship between the image guidance system and the treatment delivery system.
   b. The proper functioning of the registration software that compares planning image data sets to IGRT data sets.

3. Development, implementation, and documentation of standard operating procedures together with the radiation oncologists for the use of IGRT (including how, when, and who is to perform the IGRT procedures for each patient treatment protocol).

III. IGRT METHODS

Imaging has been used to verify patient position since the earliest days of external-beam radiation therapy (EBRT). The first method of imaging internal anatomy to verify the patient’s position on the treatment couch used the treatment beam to expose radiographic film. These images are called port films and are acquired during the treatment course with a frequency that varies but is often at the start of treatment and weekly thereafter. This may be considered as the earliest form of IGRT. Portal imaging remains an important part of patient treatment management, even though radiographic films have been replaced by electronic imaging detectors. Several imaging techniques have been developed over the years to improve the accuracy in the patient positioning/target verification process.

Ultrasound and in-room computed tomography (CT) were two early methods for routine volumetric imaging at the time of treatment delivery. Digital 2-D techniques were developed along a similar timeline and have evolved into 3-D and/or 4-D methodologies [2,3]. One of these 2-D methods uses the treatment beam to generate port images on an electronic imager (or flat panel detector) that can be compared with treatment planning images. An important addition to all of these techniques is the implementation of computer software that manually or automatically registers the current image data set with the planning or reference image set to determine the treatment couch shifts (and/or rotations), which are then executed to correctly align the target to the treatment field. Another development of the in-room 2-D imaging approach is the use of dual electronic imager(s) and kilovoltage (kV) x-ray sources mounted in a fixed geometry in the treatment room to provide stereoscopic information to identify the spatial location of target’s anatomy. The use of kV energies produce better image quality as compared with portal images acquired with the megavoltage (MV) treatment source. Some manufacturers mount the kV x-ray source and electronic imager on the treatment unit so that orthogonal images can be sequentially obtained with a simple rotation of the gantry.

Other innovations implemented by various linear accelerator manufacturers include integrated spiral data acquisition using a narrow MV beam at lower energy than the treatment beam to produce CT images of useful quality. Another technique that has become a standard feature of modern linear accelerators is cone beam CT (CBCT). CBCT is acquired using an integrated rotating large-field x-ray source and an opposing flat panel detector. After a full or partial rotation of the device around the patient, standard CT reconstruction algorithms are used to generate volumetric representation of the anatomy. A diagnostic quality 3-D volumetric data set is generated for IGRT purposes.

Systems using the treatment beam are typically referred to as mega-voltage (MV) imaging systems, whereas those using a diagnostic x-ray source are kilo-voltage (kV) imaging systems. For some MV imaging systems, beam tuning and/or the target are changed to improve image quality.

Magnetic resonance imaging (MRI) has been introduced for IGRT in the last 10 years (ie, MR-guided radiation therapy (MRgRT)). The major advantage of MRI is its superior soft-tissue contrast, allowing more accurate delineation of target and organs-at-risk (OARs) and robust image registration with the planning CT. The initial design of the commercially integrated MRgRT unit combined a MR scanner with a specially designed teletherapy machine. Newer MRgRT units are LINAC-based with a more powerful magnet for the MR scanner. Regardless of the design, the ultimate goal in MRgRT is to address intrafraction motion.
In recent years, there has been a paradigm shift in IGRT toward ART. There are three timescales for ART: offline replanning between treatments, online replanning immediately prior to a treatment, and real-time replanning during treatment [4]. Offline ART is carried out for tumors that are known to respond to radiation during the course of treatment. In offline ART, a patient may still have IGRT for each fraction. However, a new CT scan of the patient will be performed after 2 to 3 weeks of radiation therapy or if significant changes in patient anatomy are noticed at any time. The new CT data set is compared with the initial planning CT via image registration. If a tumor change is observed, new contour sets will be outlined and a new treatment plan generated for subsequent treatments. Typically, there would not be more than two replanning instances in offline ART.

In online ART, replanning is performed either on a CBCT or an MRI system, the evaluation of the new plan is done by a Qualified Medical Physicist in real time, and the patient is treated with the new plan. Online ART is still an evolving practice as issues with CBCT artifacts are active research subjects [5,6], whereas real-time replanning with MRgRT is currently building on the experience of a limited number of institutions. It is beyond the scope of this Technical Standard to discuss the essentials of both online and real-time ART.

Image registration is a crucial step in ART whether it is offline or online. Implicit in image registration is the transfer of contours from the original plan to the new image data set. Advanced image registration algorithms allow automatic morphing of the target and OAR structures to the new image data set (deformable image registration [DIR]).

The various technologies currently available for in-room patient setup and target localization are summarized below. The differences inherent in the design of each IGRT system dictate the appropriate QA procedure that is needed for its safe clinical implementation. They are fundamentally described as either 2-D (planar or surface) approaches or 3-D volumetric approaches.

1. 2-D plane
   - kV x-ray head with opposed electronic imager panels mounted on treatment unit for obtaining two or more fixed views of the patient’s anatomy.
   - Dual kV x-ray heads with opposed imaging electronic imager panel mounted at fixed positions in the treatment room.
   - Electronic imager panel mounted on the accelerator unit opposed to the treatment beam for obtaining two or more fixed views of the patient’s anatomy.

2. 3-D surface
   - Systems that use cameras and/or surface-mapping systems for external localization.

3. 3-D volumetric
   - Ultrasound unit.
   - In-room diagnostic CT.
   - kV x-ray head with opposed electronic imaging panel mounted on treatment unit for CBCT.
   - Electronic imaging panel mounted on the accelerator unit opposed to the treatment beam for obtaining MV CBCT images, with one view using a kV electronic imager and another view using MV electronic imager.
   - A narrow MV beam with opposed detector for spiral CT data acquisition.
   - MR systems.

For 2-D IGRT, the process begins with the construction of a reference image set, such as 2-D digitally reconstructed radiographs or surface rendering from the treatment planning CT data set. A 3-D IGRT utilizes the entire treatment-planning CT image set as the reference data set. These reference images are compared with the online IGRT images obtained before and/or during the treatment delivery process. The patient is repositioned based on the congruence of these image data sets such that the images are aligned to within some predetermined localization criteria. In this
manner, the treatment will be delivered precisely and accurately according to the treatment plan approved by the radiation oncologist. Implicit in this IGRT approach is the assumption of rigid target geometry.

For the purpose of this technical standard, the software components that facilitate image registration and provide the shift coordinates for the patient support system are considered to be an essential part of the IGRT system. Additionally, improvements in motion management during IGRT, while applying separate technologies and methods, will also be considered as an essential component of IGRT in this technical standard.

Given this wide spectrum of IGRT methodologies, it is difficult to devise a single generic test procedure that is appropriate for guaranteeing the safe use of this integral technology. However, this technical standard will describe the features of tests that should be included so that the potential for error is minimized. This information will allow individual institutions to compare a particular QA procedure against this list of essential features.

The record of the IGRT procedure registered in the radiation oncology information system should be reviewed to confirm accurate reporting on the session in terms of applied displacements and timeline.

IV. IGRT QA

IGRT offers the possibility of improving the accuracy and precision of dose delivery in radiation therapy and is an important advance in terms of margin reduction to better limit the dose to critical structures. However, similar to the changes in the QA procedures that occur with the adoption of any new emerging technology, the introduction of this in-room imaging step in the radiation therapy process also leads to additional QA procedures for the dose delivery system. It is critical that a QA program addresses the equipment, procedures, and safety of image guidance.

One of the most important IGRT QA components is to guarantee accurate geometric alignment of the imaging system with the treatment delivery system [7,8]. This QA procedure must be designed to include tests to ensure that the image registration part of the IGRT process performs within the stated tolerances. QA methods must also be incorporated to evaluate the accuracy of motion-management techniques in order to ensure optimal image registration. Moreover, QA tests to evaluate image quality must be performed periodically when IGRT is used. Given the recent availability of special patient support systems that allow translational and rotational position corrections, the tests must also include some verification of the performance of this part of the IGRT chain.

The test frequency has been recommended according to TG-142 and TG-226 (MPPG #2.a) [8,9]. However, it should be performed at least each month when standard fractionation treatments are scheduled. It should be performed each morning when stereotactic radiosurgery, (SRS) stereotactic radiation therapy (SRT), or stereotactic body radiation therapy (SBRT) treatment is scheduled.

A. Equipment Performance and Integration

An image-guidance system consists of components to acquire images (radiation sources, detectors, and their mechanical assemblies), measure position (image alignment tools), and perform adjustments (interfaces and equipment for position adjustment). Each of these components requires validation prior to implementation as well as routine checks to ensure safe and effective utilization.

1. Image quality

Image quality is typically characterized by physical measurements, such as contrast, resolution, and signal-to-noise ratio (SNR). It can also be evaluated by its impact on the performance of a person or alignment system that uses the images (eg, via a receiver operating characteristic curve). It is important that consistent measurement methods and phantoms are used for image-quality evaluation and that the methods are ultimately tied to the ability to use these images in practices. AAPM TG-142, TG-179, and TG-226 (MPPG #2.a) provide recommendations for the comprehensive and essential QA practices, respectively [8-10].
2. Mechanical integrity
   Whether room-mounted and stationary or gantry-mounted and moving, imaging equipment must be able to maintain a known relationship to the treatment coordinate system. The configuration and its stability should be established and monitored (e.g., checking of flex maps for centering projections as a function of angle in a gantry-mounted system). AAPM TG-142 and TG-226 (MPPG #2.a) provide recommendations for the comprehensive and essential QA practices, respectively [8,9].

3. Registration software
   IGRT equipment has both manual (e.g., visual alignment) and automated image registration tools. These tools have advantages and limitations and should be understood by evaluation prior to patient imaging. Accuracy and reproducibility of alignment results should be tested using images similar to those acquired in a clinic. AAPM TG-142 and TG-226 (MPPG #2.a) provide recommendations for the comprehensive and essential QA practices, respectively [8,9].

In clinical practice, ART requires QA of the software performing deformable image registration (DIR) to ensure the integrity of the DIR functionality [11]. The TG-132 provides useful information and guidelines on general techniques, commissioning, and QA procedures for image registration in the context of clinical utilization [12].

4. Motion-management system
   Use of motion management in IGRT may be done by multiple methods. Each of these methods requires its own evaluation for accuracy and effectiveness. Appropriate QA testing of each methodology should be done prior to its incorporation into the IGRT process. AAPM TG-76 and AAPM TG-142 contain recommended guidelines for QA and implementation of respiratory motion management [8,13].

5. Imaging dose
   Imaging parameters and associated doses for different IGRT applications should also be carefully assessed as defined by AAPM TG-75 and its complementary AAPM TG-180 [14,15]. It is important to clearly understand the imaging dose to the whole imaging volume for each IGRT procedure, especially when it applies to motion imaging. Note that the imaging volume is much larger than the treatment volume [14].

6. System integration
   An appropriate phantom is essential for IGRT implementation. There are many commercially available QA phantoms for testing integration of IGRT systems. These phantoms and various test devices often come with a description of the recommended test procedure. It is essential that users verify the appropriateness of the test equipment and procedures to ensure the accuracy and precision of the different IGRT systems in their clinic, such as, for example, the congruence of the imaging and treatment isocenter in both 2-D and 3-D imaging systems and the proper functioning of the image registration software in terms of accuracy in positioning and repositioning (couch shifts) [9]. The results of the end-to-end testing must be documented in detail as they form the baseline values for all subsequent tests of the IGRT systems.

7. User- and technology-dependent issues
   Ultrasound imaging localizes the interfaces of tissues that have different acoustic impedances. Whether ultrasound images are aligned to reference ultrasound or CT images, issues, such as interobserver variability, difficulty standardizing scanning techniques, and organ motion that is due to probe pressure, all affect accuracy in verification of patient positioning and should be appropriately considered in safe and effective use of ultrasound for positioning [16].

The use of cameras and/or surface imaging is not generally considered stand-alone image guidance for patient positioning because of the lack of internal anatomical information. Its utility for initial patient positioning and motion management must typically be supplemented with another form of internal imaging for verification. It is most useful for initial positioning and as a real-time surface monitoring aid to therapists.
as well as for cases requiring beam gating. AAPM TG 147 provides guidelines and recommendations on clinical implementation and QA procedures for nonradiographic localization systems [17].

Most Qualified Medical Physicists in radiation therapy are neither sufficiently trained in MR imaging nor qualified to perform QA of the MR scanner, and, more importantly, identify artifacts and differentiate them from actual pathology. It is prudent that a Qualified Medical Physicist with sufficient working knowledge on MRI be present at least in the first fraction of MRgRT and be available in subsequent fractions.

8. Information technology
The introduction of IGRT creates a substantial amount of image data and associated information requiring storage and management. Information systems to manage the patient data, image data, treatment data, clinical trials data, etc, can be quite challenging (eg, data must be in DICOM format, HIPAA compliant, and accessible from multiple systems) and expensive, despite the advancement in computer technology. The efficiency of storage, retrieval, and display may have significant impact on the clinical operation and information accuracy. The information flow from storage to retrieval should be tested for its accuracy, efficiency, and integrity [18]. Often, a number of information systems are involved in a single radiation oncology facility; effective and accurate communication between these systems should be ensured when implementing IGRT processes [19].

B. Correction Strategies
Use of image guidance involves determining a strategy for selecting when to image, which method to use (eg, ultrasound, x-ray, CT, or MRI), and how to utilize the acquired images. Appropriate staff qualifications and training must be considered. It is critical that implementation and maintenance of IGRT be supported by a rigorous program of documentation and training [3]. It is also important to maintain adequate staffing (a backup team for example) for each step in the IGRT process.

There exist three classes of correction strategies. The most common is online measurement and adjustment of position. For online adjustment, decisions should be made about the tolerance for a correctable action, taking into account both the accuracy with which measurement and correction can be realistically applied and the sensitivity of plan objectives to these actions. Offline corrections based on retrospective analysis of images from prior fractions need appropriate testing mechanisms to correctly evaluate and implement the measured adjustments [20]. The last method, adaptive adjustment and plan modification, will require a plan-dependent decision process to be put in place [6,21].

The QA team, consisting of physicians, physicists, dosimetrists, and therapists, should work as a group to define image-guidance and correction strategies [22]. Dry runs of a given strategy should be performed to ensure that the processes and documentation are sufficient. Of significant importance is a practical understanding of the limits of information available for alignment. A physician’s specific knowledge may be needed for image evaluation at the treatment unit. The availability of a Qualified Medical Physicist during the IGRT process is necessary to help solve issues that may arise on the imaging device, image registration, couch movements, etc. The practical tradeoff between treatment margins and the effort required to correct for errors needs to be evaluated. Another contribution to error that IGRT does not address is target delineation uncertainty, which can be potentially significant [23]. The issue is even more critical in DIR, but its broad scope precludes its inclusion in this technical standard.

C. Patient Dose
Imaging dose assessment is an important component for IGRT QA as recommended in AAPM TG-75 and AAPM TG-180 [14,15]. AAPM TG-75, TG-180, and TG-226 (MPPG #2.a) provide a useful overview of methods for measuring imaging dose from various IGRT modalities. These methods should aid the quality management program in the process of dose management [9,14,15]. IGRT methods using ionizing radiation will deliver an absorbed radiation dose to the patient that can, in some situations, be a significant fraction of the prescribed dose for treatment. Furthermore, x-ray imaging irradiates a significantly larger region than the treatment volume, and therefore, doses
to critical structures may be larger than intended. Management of the IGRT doses requires radiation physics expertise because (1) the method of measuring dose depends on the imaging geometry (eg, 2-D or 3-D, fan beam or cone beam) and (2) comparing generalized diagnostic imaging metrics, such as air kerma or CT dose index, with individualized therapeutic absorbed doses is nontrivial. On the other hand, MR does not result in radiation dose to a patient. MR-guided radiation therapy is an established though continually evolving IGRT technique that has seen increasing usage in the clinics. It is expected to play an increasingly significant role in image-guided adaptive radiation therapy [24].

AAPM TG-75 [14] and TG-226 [9] provide a useful overview of methods for measuring imaging dose from various IGRT modalities. These tools should aid the quality management program in the process of dose management. Image quality is to be judged based on the critical endpoint of the IGRT process: targeting. This endpoint provides a distinction in IGRT image quality that is different from that associated with diagnostic image quality. Thus, to control IGRT patient dose, it is recommended that dose management techniques that decrease the ionizing radiation imaging dose without affecting targeting be implemented whenever possible.

ACKNOWLEDGEMENTS

This technical standard was revised according to the process described under the heading The Process for Developing ACR Practice Parameters and Technical Standards on the ACR website (https://www.acr.org/Clinical-Resources/Practice-Parameters-and-Technical-Standards) by the Committee on Practice Parameters and Technical Standards – Medical Physics of the ACR Commission on Medical Physics in collaboration with the AAPM.

Collaborative Committee – members represent their societies in the initial and final revision of this technical standard

<table>
<thead>
<tr>
<th>ACR</th>
<th>AAPM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chee-Wai Cheng, PhD, FAAPM, Chair</td>
<td>Brent C. Parker, PhD</td>
</tr>
<tr>
<td>Join Y. Luh, MD</td>
<td>Jonas D. Fontenot, PhD</td>
</tr>
<tr>
<td>Victoria Yu, PhD</td>
<td>Arthur J. Olch, PhD</td>
</tr>
<tr>
<td>Tianyu Zhao, PhD</td>
<td></td>
</tr>
</tbody>
</table>

Committee on Practice Parameters and Technical Standards – Medical Physics
(ACR Committee responsible for sponsoring the draft through the process)

Maxwell R. Amurao, PhD, MBA, Chair
Mary Ann Keenan, DMP, Vice Chair
Priscilla F. Butler, MS, FACR
Chee-Wai Cheng, PhD, FAAPM
William R. Geiser, MS
Per H. Halvorsen, MS, FACR
Loretta M. Johnson, PhD
Lijun Ma, PhD, FAAPM
Tariq A. Mian, PhD, FACP
Jonathon A. Nye, PhD
Matthew A. Pacella, MS, FACR
Anshuman Panda, PhD
Douglas E. Pfeiffer, MS, FACR
Premavathy Rassiah, PhD
Christopher J. Watchman, PhD

Mahadevappa Mahesh, MS, PhD, FACP, Chair, Commission on Medical Physics
Jacqueline Anne Bello, MD, FACP, Chair, Commission on Quality and Safety
Matthew S. Pollack, MD, FACP, Chair, Committee on Practice Parameters and Technical Standards
Mary S. Newell, MD, FACR, Vice Chair, Committee on Practice Parameters and Technical Standards

Comments Reconciliation Committee
Ralph P. Lieto, MS, FACP, Chair
K. Elizabeth Hawk, MD, MS, PhD, FACP, Co-Chair
Maxwell R. Amurao, PhD, MBA
Jacqueline A. Bello, MD, FACP
Richard Benson, PhD
Join Y. Luh, MD
Mahadevappa Mahesh, MS, PhD, FACP
Jean M. Moran, PhD
Mary S. Newell, MD, FACP
Arthur J. Olch, PhD
REFERENCES


**GLOSSARY**

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>AAPM</td>
<td>American Association of Physicists in Medicine</td>
</tr>
<tr>
<td>ACR</td>
<td>American College of Radiology</td>
</tr>
<tr>
<td>ART</td>
<td>Adaptive radiation therapy</td>
</tr>
<tr>
<td>CBCT</td>
<td>Cone-beam CT</td>
</tr>
<tr>
<td>CT</td>
<td>Computed tomography</td>
</tr>
<tr>
<td>DDR</td>
<td>Digitally reconstructed radiographs</td>
</tr>
<tr>
<td>DIR</td>
<td>Deformable image registration</td>
</tr>
<tr>
<td>EBRT</td>
<td>External beam radiation therapy</td>
</tr>
<tr>
<td>IGRT</td>
<td>Image-guided radiotherapy</td>
</tr>
<tr>
<td>IMRT</td>
<td>Intensity modulated radiation therapy</td>
</tr>
<tr>
<td>kV</td>
<td>kilovoltage</td>
</tr>
<tr>
<td>MRgRT</td>
<td>Magnetic resonance-guided radiation therapy</td>
</tr>
<tr>
<td>MV</td>
<td>Megavoltage</td>
</tr>
<tr>
<td>QA</td>
<td>Quality Assurance</td>
</tr>
<tr>
<td>SRS</td>
<td>Stereotactic radiosurgery</td>
</tr>
<tr>
<td>SRT</td>
<td>Stereotactic radiation therapy</td>
</tr>
<tr>
<td>SBRT</td>
<td>Stereotactic body radiation therapy</td>
</tr>
</tbody>
</table>

*As of May 2015, all practice parameters and technical standards that are collaborative with only the American Association of Physics in Medicine are approved by the ACR Council Steering Committee and the ACR Board of Chancellors and will not go through the ACR Council (ACR Resolution 54, 2015). The effective date is the first day of the month following a 60-day period that begins on the date the document was approved.*
Development Chronology for this Technical Standard
2009 (Resolution 5)
Revised 2014 (Resolution 36)
Revised 2019 (CSC/BOC)
Amended 2022 (Resolution 41f)
Amended 2023 (Resolution 2c)