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 guidelines 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 guidelines and technical standards will be reviewed for revision or renewal, as appropriate, on their fifth anniversary or sooner, if indicated.

Each practice guideline 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 guidelines 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 guideline and technical standard by those entities not providing these services is not authorized.

**ACR–ASTRO PRACTICE GUIDELINE FOR IMAGE-GUIDED RADIATION THERAPY (IGRT)**

**PREAMBLE**

These guidelines are an educational tool designed to assist practitioners in providing appropriate radiation oncology care for patients. They 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 cautions against the use of these guidelines 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 physician or medical physicist in light of all the circumstances presented. Thus, an approach that differs from the guidelines, 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 the guidelines when, in the reasonable judgment of the practitioner, such course of action is indicated by the condition of the patient, limitations of available resources, or advances in knowledge or technology subsequent to publication of the guidelines. However, a practitioner who employs an approach substantially different from these guidelines is advised to document in the patient record information sufficient to explain the approach taken.

The practice of medicine involves not only the science, but also 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

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1Iowa Medical Society and Iowa Society of Anesthesiologists v. Iowa Board of Nursing, ___ N.W.2d ___ (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.
certainty a particular response to treatment. Therefore, it should be recognized that adherence to
these guidelines 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 sole purpose of these guidelines is to assist practitioners in achieving this objective.

I. INTRODUCTION

This guideline was developed revised collaboratively by the American College of Radiology
(ACR) and the American Society for Radiation Oncology (ASTRO).

Image-guided radiation therapy (IGRT) is radiation therapy that employs imaging to
maximize accuracy and precision throughout its entire process. This process includes target
and normal tissue delineation, radiation delivery, and adaptation of therapy to anatomic
and biological changes over time in individual patients. This guideline focuses on image-
guidance at the time of radiation delivery to ensure its adherence to the planned treatment,
referred to as in-room IGRT.

Radiation therapy has long been image-guided, but rapidly evolving imaging technologies
led to substantially greater accuracy and precision of radiation delivery. The need for this
improved accuracy and precision has been amplified by ongoing advances in radiation
planning and delivery that permit much more conformal dose distributions with sharper
dose gradients. Thus IGRT is particularly applicable to highly conformal treatment
modalities such as 3-D conformal radiation therapy (3-D CRT), intensity-modulated
radiation therapy (IMRT), or proton/hadron therapy [1,2]. In the very specialized case of
stereotactic body radiation therapy (SBRT) or stereotactic ablative radiotherapy (SABR)
[3]. IGRT is considered a necessary and integral component of the entire procedure.
Nevertheless, accurate radiation therapy is important even for simple treatments. A broad
range of IGRT modalities is now available, and adoption of some form of IGRT is now
widespread [4].

For many years the only means of verifying the proper orientation of treatment beams during
radiation therapy was the use of Historically, megavoltage “port films” were used as an early
form of IGRT, but lacked 3-D visualization of soft tissue targets and often was not applied
to every fraction, periodically obtained during the course of treatment. Such images can indicate
the location of a beam isocenter and field outlines reasonably well relative to bony landmarks.
However the tumor being treated is often a mobile soft tissue mass within the body and patient
repositioning based on bony landmarks alone is subject to Addressing these uncertainties by
error. One solution to address this error would be to expanding the radiation field sizes
sufficiently adequately to cover the entire range of potential tumor positions within the body. This
approach by default incorporates to ensure target coverage inevitably irradiates a large volume
of normal tissue that might receive unnecessary radiation unnecessarily in the process. With
improving soft tissue localization and increasing frequency of imaging and correction,
uncertainty is mitigated allowing correspondingly reduced margins and safer
administration of curative radiation doses. Therefore it would be preferable to limit the
radiation field size if possible

In its current state of evolution, in-room IGRT (hereafter referred to simply as IGRT) is the
use of imaging at the time of treatment delivery to ensure that the location of the target
relative to the treatment beams based on a pre-determined plan is reproduced. In most
cases, this spatial relationship is determined from a 3-D image, commonly X-ray computed tomography (CT), acquired at the initial simulation.

As techniques of radiation therapy administration have evolved in recent years, methods of imaging a tumor or target volume within a patient have been coupled with treatment delivery technology that allows near-simultaneous localization of the tumor and repositioning of the patient. The goal is to direct the radiation beam toward the true location of the tumor volume within the patient, allowing for more tightly focused treatment fields. In this manner, the images are used to guide the radiation therapy; hence the term image-guided radiation therapy (IGRT).

The IGRT process of care begins with the construction of digitally reconstructed radiographs (DRRs) from the approved computer treatment plan computed tomography (CT) data set. These images will be compared to the images obtained before and/or during the treatment delivery process. At the time of treatment delivery, an IGRT modality is employed to determine the location of the target (and often the surrounding normal organs) at some frequency, most often at the beginning, to as often as nearly continuously throughout delivery. The target location may be determined by a range of methods from soft tissue volumetric imaging (e.g., kV or MV CT, ultrasound, magnetic resonance imaging) to localization of surrogates such as implanted fiducial markers or external surface markers or features (e.g., by planar imaging or fluoroscopy, electromagnetic localization, or optical surface imaging). The match or discrepancy between the simulated location and the “live” IGRT measurement at the time of treatment may be determined manually, or in some cases using automated image analysis software. If a discrepancy is found, a correction is applied.

Corrections may include repositioning the patient, either through rigid corrections (shift and/or rotation) or readjustment of anatomic relationship (e.g., neck and shoulder manipulations for head/neck treatments), or movement or reshaping of the radiation beam to match the target position, or holding the beam until the target falls in the correct location (e.g., respiratory gating). Images obtained before and/or during the treatment delivery process. The patient will be moved based on the congruence of these images data sets such that the images align 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.

IGRT can be understood as a procedure that refines the delivery of therapeutic radiation by applying image-based target relocalization to allow proper patient repositioning for the purpose of ensuring accurate treatment and minimizing the volume of normal tissue exposed to ionizing radiation. It is a procedure that is separate from the actual delivery of radiation therapy. There is a technical component that involves the acquisition and registration of images and a professional component that involves physician work to review the images and approve the extent of patient repositioning required.

IGRT can be performed to enhance either 3-dimensional conformal radiation therapy (3DCRT) or intensity-modulated radiation therapy (IMRT). In the very specialized case of stereotactic body radiation therapy (SBRT), IGRT is considered a necessary, integral component of the entire procedure. There are numerous types of imaging modalities that can be incorporated into an IGRT system. Examples include ultrasound images, low energy (kV) CT scan images, high energy (MV) CT scan images, and kV images that reveal internal landmarks or fiducials. Typically, an IGRT system includes software with the capacity to accomplish an automated fusion of the acquired images with the expected image appearance. The software then calculates the vector displacement in 3D space of the actual target location from the expected location. In some cases, rotational distortion
is calculated in addition to linear misalignment. The $x$, $y$, and $z$ axis displacement (and sometimes rotational error) are then corrected by moving the couch on which the patient is immobilized.

This guideline addresses qualifications and responsibilities of personnel, clinical IGRT implementation, of IGRT including personnel qualifications, quality assurance standards and suggested documentation, and quality control and improvement, safety, and patient education.

A literature search was performed and reviewed to identify published articles regarding guidelines and standards in IGRT. Selected articles are found in the suggested additional reading section.

II. QUALIFICATIONS AND RESPONSIBILITIES OF PERSONNEL

See the ACR Practice Guideline for Radiation Oncology [5] where qualifications, credentialing, professional relationships and development are outlined.

A. Radiation Oncologist


1. The qualifications of the radiation oncologist shall be clearly defined and should include the following:

   a. Certification in radiology by the American Board of Radiology of a physician who confines his/her professional practice to radiation oncology or certification in radiation oncology or therapeutic radiology by the American Board of Radiology, the American Osteopathic Board of Radiology, the Royal College of Physicians and Surgeons of Canada, or the Collège des Médecins du Québec may be considered proof of adequate physician qualifications. If this certification did not include IGRT, then specific training in IGRT should be obtained before performing any stereotactic procedures.

   or

   b. Satisfactory completion of a residency program in radiation oncology approved by the Accreditation Council for Graduate Medical Education (ACGME), the Royal College of Physicians and Surgeons of Canada (RCPSC), the Collège des Médecins du Québec, or the American Osteopathic Association (AOA). If this training did not include IGRT, then specific training in IGRT should be obtained before performing any stereotactic procedures.

2. The responsibilities of the radiation oncologist shall be clearly defined and should include the following:

   a. The radiation oncologist will manage the overall disease-specific treatment regimen, including: careful evaluation of disease stage; assessment of comorbidities and previous treatments; thorough exploration of various treatment options; ample and understandable discussion with patients regarding the impact of treatment, including benefits and potential harm; knowledgeable conduct of IGRT as outlined below; and prudent follow-up after treatment.

   b. The radiation oncologist will determine and recommend and approve: a proper patient positioning method with attention to disease-specific targeting concerns; patient-specific capabilities (eg, arm position in arthritic patients, degree of recumbency recumbences) in patients with severe chronic obstructive pulmonary disease); patient comfort; stability of setup; and accommodation of devices
accounting for organ motion (eg, gating equipment) required for targeting through the IGRT approach.

c. The radiation oncologist will determine and recommend and approve a procedure to account for the intra-treatment motion/variation, and the potential residuals from onboard image registration, localization and correction procedures inherent organ motion (eg, breathing movement) for targets that are significantly influenced by such motion (eg, lung and liver tumors) as they relate to and integrate with the IGRT approach chosen. This activity may include execution of a variety of methods, including: respiratory gating; tumor tracking; organ motion dampening; or patient-directed methods (eg, active breath holding).

d. It is the radiation oncologist’s responsibility to supervise the patient’s IGRT simulation using appropriate imaging methods (eg, 4-D CT for the case of thorax lesions). The radiation oncologist needs to be aware of the spatial accuracy and precision of the simulation modality and the IGRT delivery. Steps must be taken to ensure that all aspects of simulation, including positioning, immobilization, and accounting for inherent organ motions, are properly carried out using IGRT in a consistent fashion.

e. After the planning images have been acquired, they will be transferred to the treatment-planning computer, and the radiation oncologist will contour the outline of the IGRT targets of interest. Normal organ structures may be contoured by the physicist or dosimetrist and reviewed by the radiation oncologist. Specific structures that may be used to facilitate IGRT may also be contoured. Various imaging platforms known to be useful for the specific disease treated should be fused into the planning dataset for targeting. via useful mutual information Subsequently the radiation oncologist will coordinate the design for the proper planning target volume (PTV) beyond the tumor targets. In addition to these tumor targets, the radiation oncologist will confirm that relevant normal tissues adjacent to and near the targets are contoured such that dose volume limits are considered. Locating and specifying the target volumes and relevant critical normal tissues will be carried out after consideration of all relevant imaging studies.

f. The radiation oncologist will convey case-specific expectations for prescribing the radiation dose to the target volume and set limits on dose to adjacent normal tissue. It may be required that certain normal tissues be tracked with the IGRT process just as with the tumor target(s). Participating in the iterative process of plan development, the radiation oncologist will approve the final treatment plan in collaboration with a medical physicist and dosimetrist.

g. After obtaining informed consent for the IGRT procedures and treatment, the radiation oncologist will supervise oversee the actual treatment process. The conduct of all members of the treatment team will be under the supervision of the radiation oncologist. The radiation oncologist will be responsible for deciding what are the acceptable or unacceptable day-to-day variations in the treatment setup or provide the acceptable limit on movements.

h. The radiation oncologist will participate in the quality assurance (QA) processes, such as approval of IGRT assessments, in order to insure that the intended treatment is being delivered in the prescribed fashion.
B. Qualified Medical Physicist

For the qualifications of the Qualified Medical Physicist, see the ACR–AAPM Technical Standard for Medical Physics Performance Monitoring of Image-Guided Radiation Therapy (IGRT).

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, or by the American Board of Medical Physics (ABMP).

A Qualified Medical Physicist should meet the ACR Practice Guideline for Continuing Medical Education (CME). (ACR Resolution 17, 1996—revised in 2012, Resolution 42)

The appropriate subfield of medical physics for this guideline is Therapeutic Medical Physics. (Previous medical physics certification categories including Radiological Physics and Therapeutic Radiological Physics are also acceptable.)

If the above training did not include IGRT, then specific training in IGRT should be obtained prior to performing any IGRT procedures.

The medical physicist is responsible for the technical aspects of IGRT. Those responsibilities shall be clearly defined and should include the following:

1. Acceptance testing and commissioning of the IGRT system, thereby assuring its mechanical, software, and geometric precision and accuracy, as well as image quality verification and documentation. This includes
   a. Communication with the treatment planning system
   b. Communication with the treatment delivery system
   c. Testing of image registration software and translation to patient shift coordinates.
   d. Ensuring the storage and retrieval of patient data

2. Implementing and managing a QA program for the IGRT system to monitor and assure 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. Together with the radiation oncologist, developing and implementing standard operating procedures (SOPs) for the use of IGRT

C. Medical Dosimetrist

The responsibilities of the medical dosimetrist or otherwise designated treatment planner shall be clearly defined and should include the following:

1. Contouring clearly discernible critical normal structures
2. Ensuring proper orientation of volumetric patient image data on the radiation treatment planning (RTP) system (from CT and other fused image data sets)
3. Designing and generating the treatment plan under the direction of the radiation oncologist and medical physicist is required
4. Generating all technical documentation required to implement the IGRT treatment plan
5. Being available for the first treatment and assisting with verification for subsequent treatments as necessary

D. Radiation Therapist

The responsibilities of the radiation therapist shall be clearly defined and should include the following:

1. Understanding the proper use of the patient immobilization/repositioning system and fabricating and understanding the proper use of devices for IGRT
2. Under the supervision of the radiation oncologist and medical physicist, performing initial (planning) simulation of the patient and generating the medical imaging data appropriate for the RTP system
3. Implementing the IGRT treatment plan under the supervision of the radiation oncologist and the medical physicist or of the medical dosimetrist under the direction of the medical physicist
4. Acquiring periodic verification images for review by the radiation oncologist
5. Performing periodic evaluation of the stability and ongoing reproducibility of the immobilization/repositioning system and reporting inconsistencies immediately to the radiation oncologist and the medical physicist

E. Continuing Medical Education

Continuing medical education programs should include radiation oncologists, medical physicists, medical dosimetrists and radiation therapists.

The continuing education of the physician and medical physicist should be in accordance with the ACR Practice Guideline for Continuing Medical Education (CME) [6].

F. Staffing Levels

It is the responsibility of an institution to ensure appropriate staffing levels for the support of clinical practice. Staffing levels will be dependent on, among other things, the complexity of treatment and number of new technologies introduced in the clinic and supported for clinical use. Institutions should review their staffing levels before and after new technologies are introduced to ensure quality and safety in their standard of care.

III. IGRT IMPLEMENTATION

Introducing IGRT in clinical application includes comprehensive device operation evaluation, acceptance/commissioning, establishment of routine QA procedures, identification of appropriate disease sites, and creation of disease site and/or technique specific policies/procedures. Enforcement of sufficient initial and ongoing staff training is essential for a safe and efficient IGRT program for targeting and reduction in margin.

As IGRT technology evolves, it is the responsibility of all staff to keep an up-to-date knowledge on the technology and operational details of newly introduced and updated
IGRT devices, eg, MRI guidance, more sophisticated fiducial markers with electromagnetic localization and dose tracking, and better imaging techniques with CT, ultrasound and/or camera-based systems [7].

The commissioning/acceptance for these IGRT systems should follow technical recommendations from national profession organizations. IGRT has been routinely implemented for various disease sites, such as: brain; head and neck; lung/thorax; breast; liver; prostate/pelvis; pelvis/gynecologic tumors; spine; and for techniques such as IMRT and SBRT/SRS. The frequency of IGRT usage should be carefully balanced between the needs of the disease/technique, imaging dose and resource requirements.

A. Patient Dose

One of the undeniable benefits of IGRT is the minimization of irradiation of surrounding tissues (organs at risk and other nontarget tissues). However this generally comes at the cost of increased dose due to increased imaging, eg, fluoroscopic imaging or MV cine imaging.

As discussed in the ACR–AAPM Technical Standard for Medical Physics Performance Monitoring of Image-Guided Radiation Therapy (IGRT), imaging parameters and associated doses for different IGRT applications should also be carefully assessed as defined by AAPM TG-75. It is important to have a clear picture about the imaging dose to the whole imaging volume (much larger than target volume) for each IGRT procedure, especially when applies to motion imaging. Note that the imaging volume is much larger than the treatment volumes [8]. IGRT offers the possibility of significantly enhancing the accuracy and precision of radiation therapy methods and is an important advance in terms of margin reduction to better limit the dose to critical structures.

B. Fiducial Markers

When the target is not clearly visible and bony anatomy is not sufficient for adequate target alignment, fiducial markers may be needed.

With either simple megavoltage port films to helical in-room kilovoltage CT scanners, fiducial markers (either already present or specifically implanted) can be used as surrogates to target areas. With the use of MV or kV X-rays, fiducial markers are needed when bony alignments or soft tissue imaging quality are inadequate. The use of implanted fiducial markers in small lung and liver lesions has also enabled real-time tracking using in-room X-rays, particularly improving the accuracy of radiosurgical types of approaches [9,10]. Use of other marker based techniques such as electromagnetic tracking without the acquisition of images is an extension of the use of implanted fiducial markers where only a 3D coordinate is generated to perform the guidance [8,11]. Helical or cone beam CT scans or planar X-ray alignments are also efficient and reliable image guidance methods.

C. Moving Targets

Although the patient may be immobilized relative to an external reference system, the reproducibility of target position will vary due to the motion of internal organs during a given treatment fraction, and also due to the displacement, deformation, or alteration of targets and other tissues between fractions. Both of these factors – intrafraction motion and interfraction
motion – must be taken into account when determining the margins around the clinical target value (CTV) that will define the PTV for a given course of treatment.

Several methods may be employed using IGRT to help assess and account for such target motion, thereby leading to better coverage of the target volume and less exposure of nontarget surrounding normal tissues. Which specific method will be chosen in a given clinical situation will depend on a number of factors, such as: the technologies available; the appropriate imaging technology for the target tissue in question; the relative levels of potential intrafraction and interfraction organ motion; and the deployment of fiducial markers or other tracking devices.

At the time of patient simulation for IGRT, the radiation oncologist must decide how, if at all, both intrafraction and interfraction target motion will be taken into account.

1. Intrafraction organ motion

   Several methods may be helpful in determining the extent of potential intra-fraction organ motion, including, eg: slow or multiple acquisition of CT images; 4-D CT imaging; 4-D positron emission tomography (PET) imaging; and dynamic fluoroscopic imaging of targets or fiducial markers. Assessment of the extent of internal organ motion, including organ excursion, deformation, speed, frequency, and the presence of phase shifts, may be useful in determining which techniques, if any, would be most appropriate to compensate for or control organ motion.

   Several validated forms of motion monitoring and control exist, such as respiratory gating, abdominal compression, tumor tracking, or active breath control. A QA program for the methodology should exist for the procedure, and the clinical tolerances should be predetermined.

2. Interfraction organ motion

   Displacement of internal organs may occur, lessening the accuracy and reproducibility of the external reference system at the time of treatment delivery. Methods for compensating for this problem include those that directly image the internal target in question, or those that indirectly image the target through the use of fiducial markers or other tracking devices.

   Direct imaging may include, eg: MV radiographs acquired with either radiographic film or an electronic portal imaging device (EPID); planar kV imaging for better differentiation of bony anatomy; ultrasound images; MRI; or CT imaging at the time of treatment delivery for better delineation of soft tissues, either offline, such as the adaptive-radiation-therapy (ART) strategy, or online, such as the in-room cone-beam CT (CBCT) approach.

3. **Planning target volume** definition

   Definition of PTV, in terms of the margins used to expand the CTV, must take into account the interfraction and intrafraction motion characteristics of the target, the mechanical tolerances of the imaging modalities and treatment unit, the associated uncertainties of imaging methods used at the time of simulation and treatment delivery, and the position uncertainties of fiducials or other tracking devices relative to the target in question, as well as any residual immobilization and setup uncertainties.
D. Soft Tissue

Fiducial markers do not provide information about changes in the size and shape of tumors that may occur during a several-week course of radiation therapy [12-16]. Various modalities, including orthogonal imaging, ultrasonography, or MRI/CT, can be used for real-time imaging of the tumor and surrounding tissues during radiation therapy. Using the tumor itself or the surrounding bony anatomy as a surrogate for the target, these techniques incorporate specialized software to determine positional deviations relative to pretreatment CT simulation images and to adjust patient positioning. Applications of various IGRT systems may be tumor-specific and/or site-specific depending not only on the properties of the imaging modality, but also on the type of tumor and its anatomical relation with the surrounding healthy tissues [13].

Conventional CT or CBCT cone beam CT using kilovoltage X-rays (kVCT) can be used to identify most superficial or deep-seated tumors [14,15]. Ultrasonography depicts echogenicity differences between tumors and the surrounding tissues and has been used for several years, mainly for localizing the prostate and other superficial tumors [12]. Ultrasonography may also be used to localize tumors that are found to be isodense or hypodense by unenhanced CT, such as certain tumors in the liver, to obviate the need for contrast-enhanced CT. Recently, megavoltage X-ray CT (MVCT) and in-room MRI have become available for IGRT. Even though MVCT provides images of inferior resolution compared to kVCT, this modality results in better soft tissue imaging in anatomic regions adjacent to metallic prostheses, such as dental fillings in patients with head and neck cancers, and hip prostheses in patients with pelvic tumors [16].

E. Patient Positioning

Patient immobilization could improve accuracy and reproducibility in patient positioning relative to the IGRT device and treatment unit. This can be achieved with the use of immobilization devices and is especially important when the IGRT unit is separate from the treatment unit.

F. Image Acquisition and Imaging Dose

The IGRT system needs to be calibrated to ensure high quality of imaging. The calibration ensures system performance characteristics such as slice thickness uniformity, image contrast, and spatial resolution. The IGRT system must also be accurately aligned to the isocenter of the linear accelerator and registered with the treatment planning system. The software used to identify and correct couch misalignments needs to be assessed for accuracy. Orthogonal images should be obtained and compared to digitally reconstructed radiographs (DRRs) for coincidence when applicable. Each facility needs to develop QA procedures to ensure reliability and reproducibility of the IGRT process.

Imaging dose needs to be carefully evaluated for imaging protocols used. Extensive efforts have been engaged to reduce imaging dose while maintaining image quality when radiation-based IGRT systems are used. At the time of this report, the imaging dose per-image ranges from 0.1 to 0.6 mGy for planar kV imaging, 1 to 3 mGy for MV planar imaging, and 10 to 50 mGy for 3D X-ray imaging. For 4-D image acquisition or tracking with radiation-based systems, accumulated dose from these imaging should be evaluated, eg, imaging dose from fluoroscopy can reach over 1,000 mGy/hour [7].
G. Treatment Verification

IGRT images need to be reviewed by the physician initially and then periodically to ensure treatment accuracy and reproducibility. Each facility, under the direction of the radiation oncologist, should consider establishing a threshold of couch positioning changes above which the physician is required to review the patient setup and images before treatment is delivered.

IV. DOCUMENTATION

Reporting should be in accordance with the ACR Practice Guideline for Communication: Radiation Oncology [17].

Successful IGRT implementation includes specification of the type of imaging modality used, its frequency, and the anatomical or fiducial targets employed. As noted above, various verification methodologies of IGRT implementation are in current use, and one or more appropriate methodologies should be incorporated into the patient’s record, as part of documentation of treatment parameters.

V. QUALITY CONTROL AND IMPROVEMENT, SAFETY, AND PATIENT EDUCATION

Policies and procedures related to quality, patient education, infection control, and safety should be developed and implemented in accordance with the ACR Policy on Quality Control and Improvement, Safety, Infection Control, and Patient Education appearing under the heading Position Statement on QC & Improvement, Safety, Infection Control, and Patient Education on the ACR website (http://www.acr.org/guidelines).

In-room IGRT is different than older techniques that use imaging for patient setup. It uses specialized computer techniques to register the image data set (planning to periodic images obtained during or very near the time of treatment in the accelerator room) images acquired at time of radiation delivery to ensure its adherence to the planned treatment. A more precisely monitor and/or correct setup deviations. The use of IGRT requires additional QA procedures to demonstrate that the treatment and image guidance systems are geometrically related to fall within a stated tolerance. IGRT requires a modification to traditional QA procedures. An additional step must be inserted to demonstrate that the treatment and image guidance systems are geometrically related to fall within a stated tolerance. The complete image guidance process includes the treatment planning component that assigns field positions and creates the images that are used to verify the match that is achieved through the image registration step.

IGRT is a complete process; a complex interaction of different systems must be monitored through a comprehensive QA procedure or set of procedures. This beginning end-to-end QA test must start with the CT simulation procedure and go through all steps that are required until reaching the final step treatment. and extend through treatment planning, IGRT positioning and finally the treatment step. It may be set up as a 2 step process. The first step is to position test markers in space using the IGRT system, and the second step must irradiate and image these markers with the treatment beam. The connection of the 2 steps is through the treatment planning system.

There are different end-to-end tests that have been developed. ways of accomplishing the test described above. One simple example. However, one of the simplest that can be applied to most
IGRT systems involves uses a plastic block phantom with a few embedded radio-opaque fiducial markers. The exact position of these markers is determined during CT simulation process. The treatment planning step places a series of small fields that hit each of these markers from at least 2 orthogonal directions, and creates digitally reconstructed radiographs (DRRs) showing the expected position of the markers in the treatment fields. The phantom is then placed on the treatment couch with intentional setup errors. After IGRT correction of the position of the phantom position, the treatment beam is used to irradiate and image the markers. Any detected difference in the position of the markers quantifies the overall error in the system.

A. Fiducial Markers

To serve as adequate surrogates, implanted fiducial markers need to be stable. The stability of intraprostatic fiducials has been well documented [18,19]. However, for lung and liver lesions treated with small margins, fiducial migration needs to be ruled out [9,20]. If more than 1 fiducial marker is implanted, intermarker distances are simple measures of migration. However, if migration is suspected, a CT scan should be obtained to document and reestablish the correlation between the target areas and the implanted fiducial marker. This is particularly important in situations where image interpretation as part of the image registration process during the guidance is minimal, or when an image is not obtained at all (eg, electromagnetic tracking). It is also important to avoid the mistake of interpreting bone versus fiducial location variations or deformation of target areas rather than migration. The larger the magnitude of intermarker distances or the magnitude of bone versus fiducial distances, the more migration is the likely explanation of positional variations.

B. Moving Targets

At the time of patient simulation for IGRT, the radiation oncologist must decide how, if at all, both intrafraction and interfraction target motion will be taken into account.

1. Intrafraction organ motion

Several methods may be helpful in determining the extent of potential intrafraction organ motion, including, eg, 4-D CT imaging, 4-D PET imaging, and dynamic fluoroscopic imaging of targets or fiducial markers in or proximal to the target volume or other visible surrogate. Assessment of the extent of internal organ motion – including organ excursion, deformation, speed, frequency, and the presence of phase shifts – may be useful in determining which techniques, if any, would be most appropriate to compensate for or control organ motion.

Several validated forms of motion monitoring and control management exist, such as respiratory gating, abdominal compression, tumor tracking, or active breath control. The QA procedures being used should be relevant to the motion management system that is clinically implemented in their institution. A QA program should exist for these procedures, and the clinical tolerances should be explicitly determined.

2. Interfraction organ motion

Displacement, deformation, or growth or shrinkage of targets or other organs may occur over time, lessening the accuracy and reproducibility of the external reference system over a course of treatment. at the time of treatment delivery. Methods for compensating for this problem include those that directly image the internal target in question, or those
that indirectly image the target through the use of fiducial markers or other tracking devices.

Direct imaging at time of treatment may include, for example, MV radiographs acquired with either radiographic film or EPID, planar kV imaging for better differentiation of bony anatomy or CT imaging at the time of treatment delivery for better delineation of soft tissues and bony anatomy, either offline such as the ART strategy or online such as the CBCT approach.

Imaging of fiducial markers or other tracking devices may be used to provide indirect information regarding the position of the target at the time of treatment delivery, particularly when direct imaging technologies provide lower resolution or greater uncertainty about the position of the tissues in question.

Deployment of a fiducial marker system, however, requires assessment of potential uncertainties of the position of the fiducials relative to the position of the target, including the potential motion of the fiducial markers relative to the target volume over the entire time period of the treatment course.

Any imaging system, whether direct or indirect, should have an associated QA program with its clinical tolerances explicitly determined and parameter limits clearly defined.

3. Target definition

Definition of PTV, in terms of the margins used to expand the CTV, must take into account several constraints including, but not limited to: the interfraction and intrafraction motion characteristics of the target; the mechanical tolerances of the imaging modalities and treatment unit; the associated uncertainties of imaging methods used at the time of simulation and treatment delivery; and the position uncertainties of fiducial markers or other tracking devices relative to the target in question, as well as any residual immobilization and setup uncertainties.

C. Soft Tissues

Since IGRT is used in conjunction with highly conformal radiotherapy techniques, its accurate implementation is of IGRT technologies and workflow are extremely important in ensuring adequate tumor coverage and avoidance of organs at risk. Moreover, IGRT may detect changes in the shape and/or size of tumors, which may necessitate modifications of the initial radiation dose distribution. The radiation oncologist must review the IGRT images and may need to revise the initial plan according to the individual patient’s clinical situation.

Verification of accuracy in treatment delivery requires understanding of the individual IGRT system and the ability to interpret the IGRT images in relation to those acquired at treatment planning. This process should include verification of patient positioning and documentation of the required couch shifts. It should result in congruence between portals, CT, or ultrasonographic images and DRRs created from the planning CT or other initial imaging. IGRT images should be reviewed and approved by the radiation oncologist to ensure that the radiation doses will be delivered to the designated clinical volumes as planned.

Each facility should develop its own clinical guidelines for the initial and ongoing implementation and documentation of IGRT throughout a course of radiation treatment. In
particular, consideration should be given in establishing a threshold of couch positioning changes that requires the radiation oncologist’s involvement before the treatment is delivered, to verify the patient/tumor positioning and assess whether any couch adjustments are warranted. Confirmation of treatment positioning should otherwise be performed at least weekly throughout the course of radiation therapy. IGRT should be used in combination with other QA processes such as those employed to ensure proper gantry, jaw, and multileaf collimator settings or those used to verify IMRT plans.

The Medical Director of Radiation Oncology is responsible for ensuring that there is an appropriate continuing quality improvement (CQI) program as described in the ACR Practice Guideline for Radiation Oncology and the ACR Practice Guideline for the Performance of Radiation Oncology Physics for External Beam Therapy [5,21]. It is the director’s responsibility to respond to identified problems, see that the actions are taken, and evaluate the effectiveness of the actions.

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REFERENCES


17. American College of Radiology. ACR practice guideline for communication: radiation oncology. Available at:


**Suggested Reading** (Additional articles that are not cited in the document but that the committee recommends for further reading on the topic)


*Guidelines and standards are published annually with an effective date of October 1 in the year in which amended, revised, or approved by the ACR Council. For guidelines and standards published before 1999, the effective date was January 1 following the year in which the guideline or standard was amended, revised, or approved by the ACR Council.

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