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–ASTRO PRACTICE PARAMETER FOR THE PERFORMANCE OF STEREOTACTIC BODY RADIATION THERAPY

PREAMBLE

This document is an educational tool designed to assist practitioners in providing appropriate radiation oncology 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 in light of 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 the condition of the patient, limitations of available resources, or advances in knowledge or technology subsequent to publication of this document. However, a practitioner who employs an approach substantially different from the guidance in this document 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 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 sole 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, ___ 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.
I. INTRODUCTION

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

Advancements in target delineation, motion management, treatment planning, and image-guided radiation therapy (IGRT) have served as cornerstones to the development of stereotactic body radiation therapy (SBRT) in radiation oncology. SBRT is also referred to as stereotactic ablative radiation therapy (SABR) in some publications; however, for the purposes of this practice parameter, the term SBRT will exclusively be used. Despite the presence of consensus guidelines recommending SBRT as standard of care [1], there exists variability in the practice of this modality [2-5]. The central aim of this practice parameter is to highlight critical components that are recommended for safe and appropriate implementation of this complex special procedure in radiation oncology.

SBRT is a strategy employed to precisely deliver dose to a well-defined extracranial target in 5 fractions or less [6] and commonly employs substantially higher doses per treatment than in conventional fractionation. SBRT thus demands a significantly greater degree of precision and accuracy and requires a coordinated team effort. The clinical experience reported in lung SBRT has significantly helped contribute toward the continued advancement of SBRT across other disease sites [3,5]. Optimal SBRT schedules vary by cancer type and proximity to specific organs at risk. The biological equivalence relationships of these higher doses per fraction are not completely understood, require continued study, and are outside the scope of this parameter. Given the added complexity of this special procedure, patients under consideration for SBRT benefit from multidisciplinary review, and enrollment on open clinical trials is encouraged to better understand the relative safety and efficacy of this complex treatment modality over time.

SBRT is a continuously evolving therapy, including a variety of techniques to address the challenges posted by the motion of the target and surrounding organs. Although treatment of intracranial sites may be understood conceptually as a form of SBRT, for the purpose of this document, SBRT is strictly defined as radiation therapy delivered via stereotactic guidance with high levels of targeting accuracy to extracranial targets. For information regarding intracranial target treatments, refer to the ACR Practice Parameter for the Performance of Brain Stereotactic Radiosurgery [7].

The purpose of this practice parameter is to provide guidance to physicians and medical physicists to define quality criteria in view of the high technical demands of SBRT. Megavoltage photons and protons may be used for SBRT. During irradiation with photons, multiple options are available for the radiation beams. These options include static fields with or without intensity-modulated radiation therapy (IMRT) techniques, rotational fields with or without/volumetric modulated arc therapy (VMAT), helical TomoTherapy, and multiple robotically directed beams.

SBRT requires stereotactic target localization and improved delivery precision over those required for conventional 3-D treatment delivery. For example, a conventional multileaf collimator (MLC) leaf width of 1 cm is inadequate for treating small targets. Higher confidence in targeting accuracy facilitated by imaging and positioning techniques is necessary to reduce uncertainties and corresponding target margins. Maneuvers to either limit or compensate for movement of the patient (as in breathing) or of the tumor during treatment planning and delivery are vital components to this technique [6]. Motion assessment techniques include 4-D computed tomography (CT), breath-hold techniques, fluoroscopy, and magnetic resonances (MR) guidance to assess tumor motion during simulation and treatment delivery [8,9]. Likewise, similar maneuvers are used on the treatment machine to compensate for tumor motion. Patient maneuvers on the treatment machine include breath-hold techniques and motion dampening with abdominal compression. Equipment maneuvers include techniques such as beam gating and tumor tracking. All delivery methods must ensure treatment of the entire tumor pathway during inhalation and exhalation using either patient- and/or equipment-based techniques.

SBRT, like conventional radiation therapy, uses target definitions defined within Reports 50 and 62 published by the International Commission on Radiation Units and Measurements [10,11]. Definitions of gross target volume (GTV), clinical target volume (CTV), planning target volume (PTV), and organs at risk (OAR) are employed. The
CTV may fluctuate in size and position because of respiratory motion or organ dynamics in certain sites. This is generally accounted for by adding an internal motion margin to the CTV, resulting in an internal target volume (ITV). As the delineation of an ITV implicitly includes the CTV under the assumption that microscopic disease extension is incorporated within the region of internal motion, under these circumstances the CTV often is left undefined. Stable sites may not require an ITV.

The use of multiple nonoverlapping beams or arc(s) spanning a large angular range is the principle means to achieve a rapid dose fall-off with SBRT. To ensure accurate radiation beam placement, stereotactic localization of the target is accomplished for each treatment with imaging and/or placement of fiducial marker(s). Whether delivery systems have room-mounted or on-board x-ray or CT-based localization for image guidance, the expected results are similar. For either localization system, visible tumors, fiducial markers, or appropriate anatomic landmarks are used. Finally, localization or preparatory filling/emptying (ie, full bladder protocols) of critical normal structures may be confirmed by imaging as well, as needed.

Strict protocols for quality assurance (QA) must be followed. SBRT requires levels of precision and accuracy that surpass the requirements of conventionally fractionated radiation therapy or intensity-modulated delivery because of the high radiation doses used per fraction. A variety of task groups and reports are available that provide guidance toward commissioning and QA of SBRT delivery devices [6,12] as well as imaging and treatment planning systems [13–15]. The SBRT process requires a coordinated team effort between the radiation oncologist, medical physicist, medical dosimetrist, nurse, and radiation therapist. Refer to ACR-AAPM Technical Standard for Medical Physics Performance Monitoring of Stereotactic Body Radiation Therapy (SBRT).

Figure 1 demonstrates the typical SBRT workflow.

This practice parameter addresses qualifications and responsibilities of personnel, clinical implementation of SBRT, quality control and improvement, safety, and patient education. Despite the known variance in clinical practice on the use of SBRT, we believe that this practice parameter will help set the minimum requirements to ensure consistent clinical outcomes with the evolving greater scope of SBRT in clinical practice.
### Fig 1. SBRT Practice Parameters and Working Flow

<table>
<thead>
<tr>
<th>RADIATION ONCOLOGY DEPARTMENT SBRT TEAM</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Equipment: linear accelerator with in-room image guidance, CT, and/or MR simulator</td>
</tr>
<tr>
<td>• Written department processes and protocols</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>SPECIAL PHYSICS CONSULTATION</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>SIMULATION AND TREATMENT PLANNING</th>
</tr>
</thead>
<tbody>
<tr>
<td>• 4-D CT</td>
</tr>
<tr>
<td>• ITV-based motion management strategy</td>
</tr>
<tr>
<td>• Immobilization</td>
</tr>
<tr>
<td>• Protocolled patient preparatory maneuvers (ie, bladder filling)</td>
</tr>
<tr>
<td>• Additional imaging (additional CT, MRI, PET, etc) for volume delineation</td>
</tr>
<tr>
<td>• Peer review</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PRE-TREATMENT ASSESSMENT FOR VITALS, PAIN, INTERVAL HISTORY</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Consideration of medication for patient comfort.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>TREATMENT DELIVERY</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Daily pretreatment image guidance</td>
</tr>
<tr>
<td>• Supervision by radiation oncologist and medical physicist according to institutional guidelines</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>FOLLOW-UP</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Structured follow-up for clinical outcome assessment</td>
</tr>
</tbody>
</table>
II. QUALIFICATIONS AND RESPONSIBILITIES OF PERSONNEL

See the ACR–ASTRO Practice Parameter for Radiation Oncology [16] for outlined qualifications, credentialing, professional relationships, and treatment practice parameters.

The following are minimal recommendations for staffing levels and staff responsibilities while participating in an SBRT procedure. Specific duties may be reassigned as appropriate.

A. Radiation Oncologist

Certification or satisfactory completion of training in radiation oncology should be documented for the radiation oncologist as detailed in the ACR–ASTRO Practice Parameter for Radiation Oncology [16]. As in any procedure, physicians should do either that which they feel comfortable they can deliver safely or seek additional educational and practical experience under guidance of those more experienced.

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

1. The radiation oncologist will manage the overall disease-specific treatment regimen, including careful evaluation of disease stage, assessment of comorbidity and previous treatments, thorough exploration of various treatment options (including multidisciplinary conferences and consultation where appropriate), ample and understandable discussion of treatment impact, including its benefits and potential harms, knowledgeable design and conduct of treatment as outlined below, and prudent follow-up after treatment.

2. The radiation oncologist, in collaboration with the qualified medical physicist, will determine and recommend proper patient positioning and immobilization methods with attention to site-specific targeting concerns, patient-specific positioning, patient comfort for typically long treatment sessions, stability of setup, and accommodation of devices accounting for organ motion.

3. The radiation oncologist, in collaboration with the qualified medical physicist, will determine and recommend a procedure to account for inherent organ motion for targets that are significantly influenced by such motion. This activity may include execution of a variety of methods, including respiratory gating, tumor tracking, organ motion dampening, or patient-directed methods if necessary, depending on treatment site. Other devices may be used to stabilize or pull away organs-at-risk (OARs) from inadvertently moving into the field of treatment. Examples of these include rectal balloons and injectable gel spacers.

4. The radiation oncologist will be responsible for supervision of patient simulation using appropriate imaging methods. The radiation oncologist needs to be aware of the spatial accuracy and precision of the imaging modality, including assessment of motion [17]. Steps must be taken to ensure that all aspects of simulation, including positioning, immobilization, and methods to account for inherent organ motions, are properly incorporated.

5. After the planning images have been acquired, the images will be transferred to the treatment-planning system or contouring software. The radiation oncologist will contour the outline of the GTV. Generally, only visible tumor will be targeted, but in certain circumstances, the radiation oncologist will use his/her knowledge of the pattern of microscopic spread and knowledge of normal tissue tolerance to enlarge the GTV to constitute the CTV. In some instances of using tumor motion information obtained at the time of simulation, the concept of a “motion-corrected” CTV may be used to create an ITV. Subsequently, with knowledge of the mechanical uncertainty of the treatment apparatus, the extent of setup uncertainty, inherent and residual organ motion, and other patient or system-specific uncertainties, the radiation oncologist will coordinate the design for the proper PTV. In addition to these tumor targets, the radiation oncologist will see that relevant normal tissue OARs are contoured so that dose volume limits may be considered [18]. Locating and specifying the target volumes and relevant critical normal tissues will be carried out after consideration of all relevant imaging studies. The importance of careful contouring and review is emphasized given tight CTV and PTV margins inherent in SBRT. Methods to assess specific uncertainty levels in contouring site specific targets should be assessed by physicians in collaboration with the physicist to derive the optimal margin.

6. The radiation oncologist will generate a case-specific prescription for the radiation dose to the target volume and for dose/volume limits to normal tissues. After participating in the iterative process of plan...
development, the radiation oncologist will approve the final treatment plan in collaboration with medical physicists or dosimetrists.
7. The radiation oncologist will attend and direct the actual treatment process. Premedications, sedation, and/or pain medicine may be prescribed by the radiation oncologist as appropriate. Patients will be positioned according to the simulation and treatment plan. Treatment devices used for stereotactic targeting and methods that account for inherent organ motion will be employed. The conduct of all members of the treatment team will be under the supervision of the radiation oncologist.

B. Qualified Medical Physicist

For further information, see the ACR–AAPM Technical Standard for Medical Physics the Performance of Stereotactic Body Radiation Therapy (SBRT).

C. Radiation Therapist

A radiation therapist must fulfill state licensing requirements and should have American Registry of Radiologic Technologists (ARRT) certification in radiation therapy. The responsibilities of the radiation therapist should be clearly defined and may include the following:

1. Preparing the treatment room for the SBRT procedure
2. Assisting the treatment team with patient positioning/immobilization
3. Performing appropriate timeouts to confirm proper setup and accuracy
4. Performing IGRT and assisting with review
5. Operating the treatment unit after the radiation oncologist and medical physicist have approved the clinical and technical aspects for beam delivery
6. Monitoring the treatment to ensure treatment is completed within tolerance.
7. Specialized training is encouraged as many of the technologies used in SBRT may not be typically used for standard radiation treatment [19].

D. Other Participants

The radiation oncologist, as the primary physician involved in the assessment of patient suitability and supervision of the delivery of SBRT, may choose to obtain consultation from other specialists as necessary.

III. SPECIFICATIONS OF THE PROCEDURE

The accuracy and precision of SBRT treatment planning and delivery are critical. The treatment-delivery unit requires the implementation of, and adherence to, an ongoing QA program [12]. The mechanical tolerance of the radiation delivery apparatus should be appropriate for the clinical task. Additional tolerances to account for setup error and variation of target localization may be applied, and these are detailed in section VI. Precision and accuracy should be validated by a reliable QA process. It is recognized that various test procedures may be used with equal validity to ascertain that the treatment delivery unit is functioning properly and safely. The test results should be documented, signed by the person doing the testing, and archived.

Substantive maneuvers will be used to accurately deliver the dose plan, which was developed on the patient model and approved by the physician, to the patient on the treatment day with consideration to all patient specific changes from the simulation time. In many cases, this will require reliable immobilization and reproducible positioning maneuvers, accurate targeting with image guidance, and motion management. Efforts need to be made to account for inherent organ motion that might influence target precision. Dose distributions surrounding the target with rapid falloff to normal tissue may be achieved with the use of numerous beams or large-arced beams of radiation with carefully controlled aperture(s) as well as with IMRT or VMAT in some cases.
IV. DOCUMENTATION

Reporting should be in accordance with the ACR–ASTRO Practice Parameter for Communication: Radiation Oncology [18].

Documentation of delivered doses to volumes of target and nontarget tissues, in the forms of dose-volume histograms and representative cross-sectional isodose treatment diagrams, should be maintained in the patient’s written or electronic record. Successful SBRT implementation requires specification of treatment unit parameters, patient positioning details, and type and frequency of imaging modalities used. As noted elsewhere in this document, various verification methodologies of SBRT implementation are in current use, and documentation of the methodology used should be incorporated into the patient radiation oncology record.

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

Relative to conventional radiation therapy treatments, SBRT is a process that requires increased precision in the positioning of the treatment beams because of the large doses of radiation administered for each treatment fraction. Thus, QA of an SBRT system must guarantee that both the image-guided system and the treatment delivery system are functioning within acceptable limits for SBRT as outlined in the TG-142 report [12]. It is essential that the QA process ensures that these two systems communicate such that the information gathered by the imaging system properly directs the selected beams to the position within the patient determined by the treatment planning process. It is important to understand that it is not acceptable to test the two systems separately. The testing must be tied together.

This procedure must be a two-step process: The first step must be designed to use the image-guided system to position 1 or more test points, eg, fiducials, in space at known coordinates. The second step must work through the treatment planning system for irradiation of these test points with the actual treatment beam, using an appropriate imaging technique that verifies acceptable target localization. Some technologies employ two independent imaging systems. For example, the image-guided system might be a cone-beam device mounted on the linear accelerator, whereas the verification imaging is performed with the megavoltage beam impinging on an electronic portal imaging device (EPID) [20].

Frame-based systems may incorporate a head and neck mask or body molds (with polyurethane foam cast or vacuum bags) to immobilize patients in a stable treatment position. They might include external reference fiducials for targeting, but relying solely on body frame localization is strongly not recommended (TG-101) [6]. Frameless stereotactic methods include metallic seed implantation within a tumor; use of surrogate anatomy, such as bone, whose position is well established in relation to the target; use of the target itself as a fiducial; and optical surface guidance. Frameless systems primarily rely on image guidance of the direct target position or of a surrogate marker (implanted fiducial markers or an anatomical structure—if the relation between the surrogate and target remain stable or fixed). Immobilization with appropriate noninvasive immobilization tools, such as masks, body cushions, etc, may still be considered for frameless systems.

1. Quality control of images

SBRT is an image guidance–based treatment modality. All salient anatomical features of the SBRT patient, both normal and abnormal, are defined with CT, MRI, positron emission tomography (PET), or angiography. Image fusion with a planning CT data set may be useful in defining the target volume(s). Both high 3-D spatial accuracy and tissue contrast definition are important imaging features for using SBRT to its fullest positional accuracy. This includes considering thin slice thickness (1 to 2 mm) for simulation CT and additional planning scans (eg, MRI) if required to maximize spatial accuracy. The correct patient’s imaging is transferred into contouring and/or planning computer, including correct name of patient and date of imaging.
The images used in SBRT are critical to the entire process. The quality of patient care and treatment delivery is predicated on the ability to define the target and normal tissue boundaries as well as to generate target coordinates at which the treatment beams are to be aimed. They are used for creating an anatomical patient model (virtual patient) for treatment planning, and they contain the morphology required for the treatment plan evaluation and dose calculation.

General consideration should be given to the following issues:

Target/critical structure definition must be accurately fused to the planning CT data set to ensure spatial linearity and precise anatomical localization. Depending on the treatment site, contrast might be recommended during the planning CT. For instance, contrast may have utility in patients with soft-tissue disease to help distinguish from normal tissue but may complicate tumor delineation in bony tumors where contrast can washout the gross tumor volume. Motion at the time of simulation may affect ideal tumor delineation, thus immobilization and/or other motion management techniques may be used at the time of simulation and subsequently used during treatment to minimize this.

CT is the most useful, spatially undistorted, and practical imaging modality for SBRT. It permits the creation of the 3-D anatomical patient model and electron density distribution that are used in the treatment planning and dose calculation processes. Some CT considerations include partial volume averaging, pixel size, slice thickness, distance between slices (ideally small for SBRT), and timing of CT with respect to contrast injection, contrast washout, and image reformatting for the treatment planning system, as well as potential intrascan organ movement.

In some cases, target tissues and normal tissue structures may be better visualized by MRI. The considerations enumerated for CT also apply to the use of MRI. Additional caution is warranted in MRI because of magnetic susceptibility artifacts and image distortion. As such, MRI must be verified with CT images or through a rigorous QA program. Techniques such as combining MRI with CT images via image fusion can be used to minimize geometrical distortions inherent in MR images. Although auto-registration tools in commercial software can be used to aid fusion of MRI or other advanced imaging to CT simulation image set, the physician should ensure the fusion is adequate for the target delineation specifically at hand. Furthermore, care must be taken when contouring directly on these fused images, which assumes accurate image registration and lack of image distortion between MRI and CT.

2. Quality control for the treatment planning system

For further information, see the ACR-AAPM Technical Standard for Medical Physics Performance Monitoring of Stereotactic Body Radiation Therapy (SBRT).

VI. SIMULATION AND TREATMENT

The tolerance for radiation targeting accuracy, which includes accounting for systematic and random errors associated with setup and target motion, needs to be determined for each organ system in each department performing the SBRT by actual measurement of organ motion and setup uncertainty.

A. Positioning and Immobilization

For frame-based stereotaxy, fiducials are rigidly attached to a stereotactic index board and registered to the target or patient position. Daily image guidance is used for treatment setup.

Frameless stereotaxy uses the fiducials that are registered immediately before or during the targeting procedure. Examples of frameless stereotaxy include image capture of one or more metallic seeds (each constituting a single fiducial) placed within or near a tumor; using surrogate anatomy, such as bone (constituting a volumetric fiducial), whose position is well established in relation to the target; or using the target itself (eg, identified on the image guidance system). External fiducials or optical surface imaging may be used for initial setup, as well as tracking
during treatment delivery. For accurate SBRT treatment delivery the target position should be confirmed with a radiographic imaging method when using optical surface imaging.

The patient is positioned appropriately with respect to the stereotactic coordinate system used, ensuring that the target is within physically attainable fiducial space. The treatment position should be comfortable enough for the patient to hold still for the entire duration of the SBRT procedure. Immobilization may involve the use of devices, such as a thermoplastic mold or mask, a vacuum mold, a vacuum pillow, or immobilization cushions.

B. Respiratory Motion Assessment and Control Techniques

Following the choice of positioning and immobilization, methods to manage tumor motion during simulation and treatment are undertaken. This activity is divided into two distinct phases: (1) motion assessment and (2) motion control.

Motion assessment is the process whereby the actual time-dependent 3-D displacements of the tumor target or a reliable surrogate is quantified. Motion assessment is typically performed with 4-D CT but can also be performed with real-time fluoroscopy or other time-dependent imaging platforms. The quantified time-dependent motion trajectory for the specific patient’s tumor is considered in the context of the planning processes, techniques, and constraints with specific emphasis on the method of motion control utilized. For example, treatment planning using an ITV/PTV expansion approach requires the motion envelop to be very similar in size to the target volume (ie, very little motion) to avoid unacceptable toxicity after delivery of ablative dose. In contrast, tumors that can be effectively tracked or gated may be allowed to have a considerably larger motion envelope. In either case, the target displacements over time should be quantified in 3-D planes to create custom motion control.

Motion control techniques include abdominal compression, breath-hold, gating, and tracking. Breath-hold and gating are associated with a duty cycle where the beam is engaged and disengaged based on an ongoing understanding of the tumor’s position. Tracking requires an accurate motion model to “predict” the location of the tumor in the next moment. Abdominal compression techniques attempt to constrain the patient to perform relatively more chest wall breathing as opposed to diaphragmatic breathing (the former is associated with less motion). In any case, the method of motion control utilized must be consistently applied throughout the simulation, planning, and treatment process.

The utilization of 4-D CT scans are particularly useful for creating more accurate representations of motion effects and facilitating treatment planning. Reconstructed data sets, such as the maximum and minimum intensity projections (MIP and MinIP), can be useful for determining the motion envelopes for lung and liver tumors during the treatment planning process, respectively. In turn, the average intensity projection is used by many centers as the planning data set for dosimetry, especially when used in conjunction with abdominal compression. Simulation is performed with the respiratory control systems activated. MRI simulation or fusion of MRI and CT images may be necessary as well.

There should be a quality control (QC) program for the method of respiratory motion assessment and control used, and the clinical tolerances should be explicitly determined [8].

C. Treatment Planning

Treatment planning involves contouring of GTV and the normal structures, review of iterations of treatment plans for PTV adequate dose coverage, review of proper falloff gradients, and review of dose/volume statistics by the radiation oncologist. Every effort should be made to minimize the volume of surrounding normal tissues exposed to high and medium dose levels. This requires minimizing the consequential high dose (ie, dose levels on the order of the prescription dose) and intermediate dose (such as ITV 50% of the prescription dose) resulting from entrance of beams, exit of beams, beam overlap, scatter radiation, and enlargement of beam apertures required to allow for target position uncertainties. The target dose distribution conforms to the shape of the target in a 3-D plan, thereby avoiding unnecessary prescription dose levels occurring within surrounding normal tissues. In intensity-modulated treatment planning, the entire target volume may not be included in a beam aperture or segment. Interplay effect of tumor and MLC motions should be evaluated with appropriate methods for targets in movable anatomy for correct
dose delivery. Quantification of the dose/volume statistics for the surrounding tissues and organs is needed so that volume-based tolerances are not exceeded. It should be understood that reduction of high dose levels within normal tissue volume may require additional exposure of normal tissues to low dose levels (ie, increased integral dose).

The planning system must support an advanced type b or c (model based) dose calculation algorithm that can accurately predict the dose deposition within heterogeneous media.

D. Treatment Delivery and Verification

Precision should be validated by the QC process and maintained throughout the entire treatment process. At the onset of the SBRT program, an end-to-end test should be performed and periodically repeated when components of the SBRT equipment and procedure change.

The radiation oncologist is responsible for ensuring that patient positioning and field placement are accurate for each fraction. The image-guided stereotactic procedure is used to verify or correct the patient’s position relative to the planning image data set. Given how critical patient positioning is for this procedure, positioning prior to each treatment must be verified. The team (radiation oncologist, physicist, and therapist) must be aware of the limitations of the setup, and the tolerance or threshold for stopping, repositioning, and reimagining. To accomplish this, treatment site–specific protocols may be developed and standardized for each department based on available equipment and its limitations. The team must communicate and update each other when errors or malfunctions are suspected so that they can be readily corrected. Patient treatment should be paused if it is felt that the patient is out of position, beyond any thresholds previously specified by the radiation oncologist. Intrafractional systems may be used to track patient positioning during treatment. The Qualified Medical Physicist should be present for the setup, image guidance, and motion review for the entirety of the first fraction. The radiation oncologist should approve the image guidance and motion review and be present at the start of each treatment fraction. The Qualified Medical Physicist and radiation oncologist must be readily available should issues arise during treatment delivery [21].

Surrogate or secondary systems may be used if the primary patient positioning system fails, but this must undergo proper QA as well.

E. Correction Strategies

Three classes of correction strategies exist. 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 test mechanisms for correctly evaluating and implementing measured adjustments. The last method, adaptive adjustment and plan modification, will require a plan-dependent decision process to be in place. At present, adaptive therapy and a related technology deformable image registration are still too early in the developmental stage for appropriate guidelines to be given.

VII. FOLLOW-UP

There should be follow-up of all patients treated, and appropriate records should be maintained to determine local control, survival, and normal tissue injury. It is recommended that the treating radiation oncologist be involved in the clinical follow-up and review the post-SBRT diagnostic imaging as some abnormal radiographic changes, often mistaken as tumor progression, may actually be posttreatment effects. As with any form of radiation therapy, there is a potential risk of subacute or late toxicities that may occur months or even years after treatment. The data should be collected in a manner that complies with statutory and regulatory peer-review procedures to protect the confidentiality of the peer-review data.
VIII. CONTINUOUS QUALITY IMPROVEMENT

Continuous quality improvement (CQI) is crucial for SBRT given the potential risks associated with errors or misadministration. The requirements are outlined in ACR–ASTRO Practice Parameter for Radiation Oncology [16]. Proactive, even prospective (prior to treatment delivery), peer review is strongly suggested given the limited number of fractions which limits the time to institute any changes suggested by peer review.

IX. SUMMARY

The quality of a SBRT program depends on the coordinated interactions of a team of skilled health care professionals. A high degree of spatial accuracy is necessary in the treatment planning and delivery process. Because SBRT uses either single-fraction treatment or a hypofractionated regimen, there is little chance for adjustment once treatment has been initiated. This demands considerable time for planning and treatment verification by the radiation oncologist and medical physicist.

ACKNOWLEDGEMENTS

This practice parameter 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 – Radiation Oncology of the ACR Commission on Radiation Oncology in collaboration with the ASTRO.

Collaborative Committee – members represent their societies in the initial and final revision of this practice parameter

ACR
Samuel T. Chao, MD, Chair
Neil B. Desai, MD
Matthew A. Pacella, MS, FACP
Ying Xiao, PhD

ASTRO
Luqman K. Dad, MD
Laura A. Dawson, MD
Ramesh Rengan, MD, PhD
Kamil Yenice, PhD

Committee on Practice Parameters – Radiation Oncology
(ACR Committee responsible for sponsoring the draft through the process)

Alan C. Hartford, MD, PhD, FACR, Chair
Naomi R. Schechter, MD, Vice Chair
Nathan H. J. Bittner, MD
Samuel T. Chao, MD
Chee-Wai Cheng, PhD, FAAPM
Neil B. Desai, MD
Nancy A. Ellerbroek, MD, FACR
Mark Hurwitz, MD
Lesley A. Jarvis, MD, PhD

Seth A. Rosenthal, MD, FACR, Chair

Comments Reconciliation Committee
William Small, Jr., MD, FACR, Chair
K. Elizabeth Hawk, MD, MS, PhD, FACR (Co-Chair)
Samuel T. Chao, MD
Luqman K. Dad, MD
Laura A. Dawson, MD
Neil B. Desai, MD

Ramesh Rengan, MD, PhD
Seth A. Rosenthal, MD, FACR
Naomi R. Schechter, MD
Charles B. Simone, II, MD
Naomi R. Schechter, MD
Timothy L. Swan, MD, FACR

PRACTICE PARAMETER 11 SBRT
REFERENCES


*Practice parameters and technical 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 practice parameters and technical standards published before 1999, the effective date was January 1 following the year in which the practice parameter or technical standard was amended, revised, or approved by the ACR Council.

**Development Chronology for this Practice Parameter**

- 2004 (Resolution 20)
- Amended 2006 (Resolution 16g, 36)
- Revised 2009 (Resolution 4)
- Revised 2014 (CSC/BOC) – Effective June 25, 2014
- Revised 2019 (CSC/BOC)