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 2022 (BOC/CSC)*

ACR–ARS PRACTICE PARAMETER ON INFORMED CONSENT RADIATION ONCOLOGY

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

The process of informed consent is designed to protect a patient’s autonomy in their medical decision-making in the context of an asymmetrical relationship with the health care system. A proper consent process assures a patient’s individual autonomy, reduces the opportunity for abusive conduct or conflicts of interest, and raises trust levels among participants.

Consent is a communications process between the patient and a health care provider in which both parties have the opportunity to ask questions and exchange information relevant to the patient’s diagnosis and treatment. For this process to be effective, both parties must actively participate in the process, and both parties share the responsibility for the accurate exchange of information. Patients have a right to self-determination and to consent to any medical treatment before it is given. Physicians have a legal and ethical duty to obtain informed consent from the patient. For informed consent to be appropriately obtained, the physician is obligated to provide accurate medical information regarding any recommended treatment or procedure such that the patient or the individual responsible for the patient’s care (ie, legal representative) has sufficient understanding to make an informed decision [1-4]. In addition, the patient must be provided with adequate opportunity to ask and have satisfactorily addressed all and concerns regarding the recommended treatment or procedure.

The physician has the duty to help the patient make decisions consistent with the patient’s best interests from among the therapeutic alternatives consistent with good medical practice. The process should also reasonably guarantee freedom from coercion. Thus, informed consent must be obtained in a manner that is free from fear of the practitioner’s subsequent actions, honors both ethical and legal principles, and minimizes the question of undue inducement. The American Medical Association Code of Medical Ethics outlines its perspective on informed consent in Opinion 8.08 [4]. If medical treatment is given without informed consent having been appropriately obtained, a claim of battery or malpractice may be made against the physician or health care professional performing the procedure [1,2,5,6]. For information on exceptions to the informed consent rule, see section II.

Informed consent is a process that must not be mistaken as the simple act of signing a formal document. The informed consent document does, however, provide important documentation of the complex shared decision-making process undertaken by the patient and physician. By providing their signature, a patient indicates that they understand and consent to the treatments and procedures that will be performed. Informed consent with appropriate documentation must follow institutional policies and procedures and comply with applicable state and federal law.

II. INDICATIONS

Informed Consent Rule: Informed consent must be obtained and appropriately documented prior to the initiation of any complex medical treatment, including, but not limited to, the following procedures:

1. Imaging for simulation of radiotherapy treatment setup including field placement and/or treatment planning and any associated procedures, such as use of contrast agents and tattoo placement
2. Radiotherapy treatment including external beam radiation therapy, radiopharmaceutical therapy, high-dose rate or low-dose rate brachytherapy, etc
3. Administration of conscious sedation

Exceptions to this informed consent rule include:

1. Emergency Situation: Cases in which the patient is unable to provide consent, the patient’s legal representative is unavailable, no significant evidence exists to indicate that the patient (or the patient’s legal representative) would refuse the treatment, and immediate treatment is required to alleviate a condition causing severe pain or to prevent serious disability or death. In these cases, treatment only for the emergency condition can be given without informed consent. Should the patient regain ability to consent, or if a legal representative becomes available, the process of informed consent must be obtained for the treatment to continue [1,2].
2. Patient Declination: If a patient specifically asks not to be informed of certain facts of their diagnosis
or risks of treatment, treatment may be given with less than full disclosure. The request of the patient must be documented, including details regarding what was discussed and what was omitted. If the patient gives permission for their next of kin or legal representative to receive information relevant to informed consent, the nature of the information given should be documented. At a minimum a patient must be informed:

a. That the procedure involves some known risks, which may be serious
b. That there are some potential benefits, which should be outlined
c. That following the treatment there may be some late effects, which should be outlined
d. About how to cooperate with treatment [2]

III. QUALIFICATIONS AND RESPONSIBILITIES OF PERSONNEL

A. Physician

Informed consent for radiation oncology procedures must be obtained by or under the supervision of a licensed physician qualified to perform the procedure. The supervising physician must be familiar with the procedure being performed. For procedures in which multiple physicians will be responsible, multiple physicians may participate in the same consent process.

B. Other Health Care Professionals

Other health care professionals may provide information and standardized information materials. Members of the treatment team may also serve to verify that the patient understands the procedure. Ultimately, the physician performing the procedure is responsible for the accuracy of the information presented during the informed consent process as well as for clarifying that this information was understood by the patient or, as appropriate, by their legal representative [1,2,8,9].

C. Witnesses

It is recommended, but not required (unless by state law), that a witness be present at the time at which the patient signs the consent form. Each designated witness must be at least 18 years of age or of legal majority in the state in which the consent is given. Failure to have the patient’s signed consent witnessed does not invalidate the consent. In cases in which a mark is made by a patient to document informed consent, however, because they are unable to provide a signature, the mark must be witnessed. The witness signature serves to verify that the patient signed the consent form.
D. Interpreters

Patients who do not feel competent to understand the language spoken by the physician should be provided with an interpreter who can accurately convey what the physician is saying and can interpret any questions or concerns that the patient may have.

Patients with hearing, vision, speaking, or other communication impairments should be provided with reasonable assistance when necessary to afford such persons the ability to give informed consent.

Although less ideal, after being informed of the availability of an interpreter and if allowed by the institution, a patient may choose to use a family member or friend to serve as their interpreter. This preference should be documented, and the use of minors should be avoided.

When interpreters are used, documentation should be placed in the patient’s medical record indicating the name of the person who acted as the interpreter and that person’s title or their relationship to the patient.

E. Patient’s Legal Representative

Patients who are unable to consent for themselves, such as minors or incompetent adults, and those who choose to have another person consent for them, have the right to be represented by someone who will protect their interests and preserve their basic rights. State law also may designate who may act for a patient when he or she is incapable of giving consent. In such situations, the physician or other qualified person performing the procedure must obtain informed consent from the patient’s legal representative and arrange for the legal representative’s signature on a consent form [2]. In certain rare situations, there may be no identified surrogate decision maker. In such cases, state law and institutional policy should guide the process of consent and, to the best extent possible, follow any wishes that the patient may have previously made known.

IV. SPECIFICATIONS FOR OBTAINING INFORMED CONSENT

A. Standard Procedure

The process to obtain informed consent is designed to facilitate the disclosure of all relevant information according to the individual needs of the patient. This can be tailored to one of several possible standards: the professional standard, the level of detail that is conventional for a professional in the field to mention; the reasonable person standard, a level that a typical person may desire; or an individual standard, a level of detail that the participant determines is appropriate for them. The appropriate standard to apply is primarily dependent upon the patient requirements.

To obtain informed consent, the physician informs the patient or legal representative of:

1. The nature of the patient’s diagnosis, what is known of the extent of the disease, the goals of care, and intent of treatment (curative or palliative intent)
2. The nature of the proposed treatment, the parts of the body to be treated (including laterality when relevant), and the method by which the treatment will be given
3. The expected side effects and/or complications that a prudent practitioner would find meaningful [10] including:
   a. Complications or side effects that occur commonly and are likely to occur
   b. Complications or side effects that may be rare but serious if they do occur and that occur with enough frequency that a reasonable patient would want to be informed before deciding to accept treatment. If there is doubt about the likelihood of a complication, that doubt may be communicated as appropriate.
4. Reasonable treatment alternatives
5. The potential benefits of treatment
6. The potential consequences of refusal of treatment [2]
And as relevant:

7. For patients who retain radioactive sources or receive therapeutic radionuclides, the potential risks to family members or to the general public and restrictions for patient and public safety as applicable and/or required by federal, state, or institutional policy

8. The teratogenic effects of radiation therapy for individuals who are of childbearing potential

9. The potential impact of prior radiation therapy on the treatment options and the potential of early- and late-occurring side effects or complications of the current treatment course

10. The potential impact of radiotherapy on electronic medical devices such as pacemakers, defibrillators, etc, that may require special monitoring

These must be explained in a way in which the patient or legal representative can understand. It is generally recognized that no one method is appropriate for all physicians and all patients [3,9,11,12]. The general underlying principle is that the communication should be sincere and focused on the patient, taking into account cultural and language barriers, and individualized to the type of treatment offered [1-3,8,9,12].

Particular challenges in obtaining consent for radiation therapy include the negative connotations associated with radiation and lack of understanding of therapeutic application of radiation by the lay public [13-15]. Many patients have limited understanding of radiation therapy, with approximately half having read or heard negative stories about radiation therapy [13-15]. Preconceptions of radiation and lack of understanding may result in increased anxiety and distorted perceptions of the associated risks and benefits of treatment, making the process of informed consent more challenging [13-15].

It is recommended to include in the patient’s medical record reference to any written or other educational materials provided [1,3].

The process of informed consent can take place at the time of consultation or, preferably, over a period of time including 1 or more follow-up appointments. Informed consent is an iterative process that continues throughout a course of treatment, with provision of new relevant information should any become available. Patients have the right to ask further questions and always retain the right to withdraw consent at any time.

The consent must not be obtained in any coercive way, and the consent form must not contain any coercive statements [1,2].

**B. Special Circumstances**

1. **Underage or incompetent patients**
   For patients younger than 18 years of age, minors as determined by state law, or incompetent adults, the patient’s parent, legal guardian, or person with a medical power of attorney must give informed consent and sign the form [1,2]. In certain rare situations, there may be no identified surrogate decision maker. Informed consent with appropriate documentation must follow institutional policies and procedures and comply with applicable state law. Institutional and departmental policies and local community practice should guide the process of consent and, to the best extent possible, follow any wishes that the patient may have previously made known.

2. **Patient consent via telehealth or telephone**
   Telehealth, or telemedicine, is the use of electronic information and telecommunication technologies to provide care when the patient and the doctor are unable to be in the same place at the same time. Increased use of telehealth consultations present added challenges to the consent process. Although most patients will be seen and treated in a radiation oncology department affording the opportunity for in-person interaction, occasions arise (such as with radionuclide therapies) in which treatment is not administered in the department. Additional considerations for communication limited to telephone consent include two factor identification of the patient and inclusion of a second member of the health care team to verbally verify the consent of the patient which should be formally documented on the consent form. In addition, the patient should be provided the opportunity to meet with the physician in person to ensure all questions
and/or concerns are satisfactorily addressed prior to treatment.

3. Second party consent by telehealth, telephone, or facsimile
When a patient’s legal representative is available to give informed consent but is not physically present to sign the form, consent by telehealth using audio and visual communication, telephone or facsimile may be obtained. The responsible physician must, to the extent possible, provide the patient’s legal representative with the information the physician would disclose in a face-to-face discussion. Documentation should include the following information:
   a. Name and position of the person who spoke to the patient’s relative or legal representative
   b. Name and relationship of the legal representative
   c. A summary of the information conveyed to that legal representative
   d. A brief account of questions posed to the legal representative along with a summary of the answers to these questions
   e. Date and time of entry in the patient record as well as the writer’s signature [2]
   f. Date, time, and telephone number called

4. Clinical research
If a patient is participating in a clinical research study, he or she must not only give the standard facility informed consent but must also sign a study-specific informed consent document. The research study and the consent form must be approved by the Institutional Review Board (IRB) with jurisdiction over research at the treating facility prior to the initiation of treatment per the research protocol. If the treating physician is uncertain whether any study involving patients constitutes research requiring IRB approval, this information can be obtained by contacting the Office of Human Subjects Research in the Office of the Deputy Director for Intramural Research, National Institutes of Health.

V. DOCUMENTATION

A. The informed consent document should contain, at minimum, the following information:
   1. The patient’s name and identification number, date of birth, or other second identifier
   2. The name of the person(s) or practice group performing the procedure
   3. A statement in the first person with the patient’s name or the word “myself” authorizing administration of the proposed treatment
   4. A statement in the first person that the nature of the treatment, alternatives, side effects, and risks of injury despite precautions have been explained to the patient or person signing the form for the patient
   5. A statement in the first-person authorizing tattoos if applicable
   6. A statement in the first-person authorizing photographs for documentation
   7. A statement, when applicable, regarding the risks of radiation related to pregnancy
   8. A statement in the first person acknowledging that others may participate in the care of the patient

B. The informed consent document should have a place for:
   1. The signature of the patient or patient’s legal representative
   2. Relationship of signer if other than the patient
   3. The date
   4. Reason patient did not sign, if applicable
   5. Signature of witness, if applicable
   6. Identification of translator, if applicable

C. Additional Information

There must be a place in the permanent medical record, often on a separate document, in which the informing physician:
   1. States that he or she has informed the patient of the nature of the procedure or treatment; the risks, complications, and expected benefits or effects of such treatment or refusal; and the alternatives and their risks and benefits.
   2. States that the patient and/or patient’s legal representative have had the opportunity to have their questions answered to their satisfaction.
3. Specifies the type of radiation therapy recommended (e.g., external beam versus brachytherapy). The type of brachytherapy procedure should be specified, if applicable.

4. Documents the need for immediate treatment for patients treated on an emergency basis. Documentation should include information establishing the nature, immediacy, and magnitude of the problem and the difficulty of obtaining consent under the circumstances. All notes should include the date and time [1,2].

5. Provides special documentation for the rare situations when the patient specifically asks not to be fully informed, or where therapeutic privilege is invoked (see also section II). Communication to specified next of kin or legal representatives of the information withheld from the patient should be documented in the medical record to the extent legally allowed [1,2].

6. States when tissue or, as applicable, radiographic diagnosis has not been obtained and the implications of diagnostic uncertainty in such circumstances.

Some states require that special forms be used in the informed consent process in certain circumstances. The informed consent process must comply with state and federal law and with institutional policies and procedures.

A copy of all pertinent consent documentation should be kept in the patient’s medical record.

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REFERENCES

5. Anscher MS, Anscher BM. Medical malpractice in the age of technology: how specialty societies can make a difference. Brachytherapy 2006;5:131-4.

*As of May 2015, all practice parameters and technical standards that are collaborative with Radiation Oncology societies 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.

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