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, requiring the approval of the Commission on Quality and Safety as well as the ACR Board of Chancellors, the ACR Council Steering Committee, and the ACR Council. 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.

Revised 2012 (Resolution 6)*

ACR–ASNR–ASSR–SIR–SNIS PRACTICE GUIDELINE FOR THE PERFORMANCE OF VERTEBRAL AUGMENTATION

PREAMBLE

These guidelines are an educational tool designed to assist practitioners in providing appropriate radiologic and 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 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 revised collaboratively by the American College of Radiology (ACR), the American Society of Neuroradiology (ASNR), the American Society of Spine Radiology (ASSR), the Society of Interventional Radiology (SIR), and the Society of NeuroInterventional Surgery (SNIS).

This document addresses vertebral augmentation which includes all percutaneous techniques used to achieve internal vertebral body stabilization. Vertebral augmentation encompasses a variety of procedures for treating pathologically weakened vertebral bodies. The more common ones are vertebroplasty and acrylic vertebroplasty, which involve injecting surgical bone cement; balloon kyphoplasty (also called balloon-assisted vertebroplasty), which involves inflation of a balloon in the weakened vertebral body to attempt fracture reduction before cement is injected; and radiofrequency ablation (RFA) and coblation techniques. Other less common procedures include mechanical void creation (also called mechanical cavitation) with an osteotome, injection of bone graft material or bone substitutes, and insertion of materials in an attempt to restore the patient’s height. The field is evolving rapidly and this document also applies to
any new methods for achieving the same end: vertebral augmentation.

A thorough review of the literature was performed. When published data were felt to be inadequate, data from the expert panel members’ own quality assurance programs were used to supplement. Thresholds for quality assurance have been updated in accordance with available data in the literature.

Introduced by Galibert and Deramond et al in France in 1987 [1], vertebroplasty entails injection of material into the weakened vertebra. Radiologic imaging has been a critical part of vertebroplasty from its inception. Most procedures are performed using fluoroscopic guidance for needle placement and material injection or placement. The use of computed tomography (CT) has also been described for these purposes [2-3].

Vertebral augmentation is an established and safe procedure [1-2,4-21]. Two blinded, randomized controlled trials failed to demonstrate an advantage in their study populations for vertebroplasty over a control intervention for either pain reduction or disability improvement [22-23]. However, larger, non-blinded, prospective randomized controlled studies and other studies of vertebral augmentation have shown its efficacy [24-36]. As with any invasive procedure, the patient is most likely to benefit when the procedure is performed in an appropriate environment by qualified physicians.

These guidelines are intended to be used in quality improvement programs to assess vertebral augmentation procedures. The most important processes of care are 1) patient selection, 2) performing the procedure, and 3) monitoring the patient. The outcome measures or indicators for these processes are indications, success rates, and complication rates. Outcome measures are assigned threshold levels.

Use of other technologies to treat patients for the same indications should yield similar or better success rates and complication profiles.

II. DEFINITIONS

Vertebral augmentation includes all percutaneous techniques used to achieve internal vertebral body stabilization.

Vertebroplasty is a minimally invasive surgical or interventional procedure, performed by percutaneously injecting radiopaque bone cement into a painful osteoporotic or neoplastic compression fracture or a painful vertebral body weakened by any other etiology.

Kyphoplasty is an image-guided percutaneous procedure that creates a cavity within the bone that is then filled with material.

Failure of medical therapy is defined as:

1. For a patient rendered nonambulatory due to pain from weakened or fractured vertebral body, pain persisting at a level that prevents ambulation despite 24 hours of analgesic therapy.

or

2. For a patient with sufficient pain from weakened or fractured vertebral body that physical therapy is intolerable, pain persisting at that level despite 24 hours of analgesic therapy.

or

3. For any patient with weakened or fractured vertebral body, unacceptable side effects such as excessive sedation, confusion, or constipation due to the analgesic therapy necessary to reduce pain to a tolerable level.

III. OVERVIEW

Vertebral compression fractures are a common and often debilitating complication of osteoporosis [37-41]. Although most fractures heal within a few weeks or months, a minority of patients continue to suffer pain that does not respond to conservative therapy [42-44]. Vertebral compression fractures are a leading cause of nursing home admission. Open surgical fixation is rarely used to treat these fractures. The poor quality of bone at the adjacent unfractured levels does not provide a good anchor for surgical hardware, and the advanced age of most affected patients increases the morbidity and mortality risks of major surgery.

Initial success with vertebroplasty for treating aggressive hemangiomas [1,12] and osteolytic neoplasms [10,21] led to extension of the indications to include osteoporotic compression fractures refractory to medical therapy [2,4-9,11,13-19]. Vertebral augmentation is currently being used to treat a wide variety of osteolytic metastases and multiple myelomas.

Perioperative imaging that identifies the painful vertebral body in concordance with the clinical examination is considered essential for the safe and effective performance of vertebral augmentation.

IV. INDICATIONS AND CONTRAINDICATIONS

The major indication for vertebral augmentation is the treatment of symptomatic osteoporotic vertebral body fracture(s) refractory to medical therapy or vertebral bodies weakened due to neoplasia. Currently, there is no indication for the use of vertebral augmentation for
prophylaxis against future fracture. The indications and contraindications for vertebral augmentation may change in the future as more research and information become available.

A. Indication Threshold 95%
   1. Painful osteoporotic vertebral fracture(s) refractory to medical therapy.
   2. Vertebral bodies weakened by neoplasm.
   3. Symptomatic vertebral body microfracture (as documented by magnetic resonance imaging [MRI] or nuclear imaging, and/or lytic lesion seen on computed tomography (CT) without obvious loss of vertebral body height.

When fewer than 95% of vertebral augmentations in an institution are performed for the above indications, it should prompt a review of practices related to selection of patients for this procedure.

B. Absolute Contraindications
   1. Septicemia.
   2. Active osteomyelitis of the target vertebra.
   3. Uncorrectable coagulopathy.
   4. Allergy to bone cement or opacification agent.

C. Relative Contraindications
   1. Radiculopathy in excess of local vertebral pain, caused by a compressive syndrome unrelated to vertebral collapse. Occasionally preoperative vertebroplasty can be performed before a spinal decompressive procedure.
   2. Retropulsion of a fracture fragment causing severe spinal canal compromise.
   3. Epidural tumor extension with significant encroachment on the spinal canal.
   4. Ongoing systemic infection.
   5. Patient improving on medical therapy.
   6. Prophylaxis in osteoporotic patients (unless being performed as part of a research protocol).
   7. Myelopathy originating at the fracture level.

V. QUALIFICATIONS AND RESPONSIBILITIES OF PERSONNEL

A. Physician

In general, the requirements for physicians performing vertebral augmentation may be met by adhering to the recommendations listed below:

1. Certification in Radiology or Diagnostic 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, and must include performance of successful vertebral augmentation procedures in at least 5 patients as the primary operator, under the supervision of a qualified physician, and without major complications.

2. Completion of an approved residency or fellowship program 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 an American Osteopathic Association (AOA) approved residency program that included 6 months of training in cross-sectional imaging, including CT and MR imaging, and 4 months of training in image-guided interventional radiological techniques, including vertebral augmentation, biopsy and drainage procedures, and vascular embolization. This must include performance of successful vertebral augmentations in at least 5 patients as the primary operator, under the supervision of a qualified physician, and without major complications.

3. A physician who did not successfully complete an ACGME approved radiology residency or fellowship program that included the above may still be considered qualified to perform vertebral augmentation provided the following can be demonstrated: the physician must have at least 1 year of experience in performing percutaneous image-guided spine procedures, during which the physician was supervised by a physician with active privileges in these spine procedures. During this year he or she must have performed a minimum of 5 vertebral augmentations as primary operator with outcomes within the quality improvement thresholds of this guideline.

4. Physicians meeting any of the qualifications in 1, 2, or 3 above must have written substantiation that they are familiar with all of the following:

   a. Indications and contraindications for vertebral augmentation.
   b. Periprocedural and intraprocedural assessment, monitoring, and management of the patient, and particularly the recognition and initial management of procedural complications.
   c. Appropriate use and operation of fluoroscopic and radiographic equipment, digital subtraction systems, and other electronic imaging systems.
   d. Principles of radiation protection, hazards of radiation exposure to the patient and the
radiologic personnel, and radiation monitoring requirements.

e. Anatomy, physiology, and pathophysiology of the spine, spinal cord, and nerve roots.

f. Pharmacology of contrast agents and implanted materials and recognition and treatment of potential adverse reactions to these substances.

g. Technical aspects of performing this procedure.

The written substantiation should come from the chief of interventional radiology, the chief of neuroradiology, the chief of interventional neuroradiology, or the chair of the department of the institution in which the physician will be providing these services. Substantiation could also come from a prior institution in which the physician provided the services, but only at the discretion of the current interventional, Neurointerventional, or neuroradiology chief, or the chair who solicits the additional input.

5. Physicians must possess certain fundamental knowledge and skills that are required for the appropriate application and safe performance of vertebral augmentation:

a. In addition to a basic understanding of spinal anatomy, physiology, and pathophysiology, the physician must have sufficient knowledge of the clinical and imaging evaluation of patients with spinal disorders to determine those for whom vertebral augmentation is indicated.

b. The physician must fully appreciate the benefits and risks of vertebral augmentation and the alternatives to the procedure.

c. The physician is required to be competent in the use of fluoroscopy, CT, and MRI or interpretation of images in the modalities used to evaluate potential patients and guide the vertebral augmentation procedure.

d. The physician should be able to recognize, interpret, and act immediately on image findings.

e. The physician must have the ability, skills, and knowledge to evaluate the patient’s clinical status and to identify those patients who might be at increased risk, who may require additional perioperative care, or who have relative contraindications to the procedure.

f. The physician must be capable of providing the initial clinical management of complications of vertebral augmentation, including administration of basic life support, treatment of pneumothorax, and recognition of spinal cord compression.

g. Training in radiation physics and safety is an important component of these requirements. Such training is important to maximize both patient and physician safety. It is highly recommended that the physician have adequate training in and be familiar with the principles of radiation exposure, the hazards of radiation exposure to both patients and radiologic personnel, and the radiation monitoring requirements for the imaging methods listed above.

Some methods of vertebral augmentation may require specialized training and experience, and such needs should be assessed before a physician contemplates using any method.

Maintenance of Competence

Physicians should perform a sufficient number of vertebral augmentation procedures to maintain their skills, with acceptable success and complication rates as laid out in this guideline. Continued competence depends on participation in a quality improvement program that monitors these rates. Regular attendance at postgraduate courses that provide continuing education on diagnostic and technical advances in vertebral augmentation is necessary.

Continuing Medical Education

The physician’s continuing education should be in accordance with the ACR Practice Guideline for Continuing Medical Education (CME).

B. Nonphysician Practitioners

Physician assistants and nurse practitioners can be valuable members of the interventional radiology team but not as primary operator. These nonphysician practitioners can function as independent members of the team but not as primary operator. See the ACR-SIR-SNIS Practice Guideline for Interventional Clinical Practice.

C. Qualified Medical Physicist

A Qualified Medical Physicist is an individual who is competent to practice independently one or more of the subfields in medical physics. The American College of Radiology (ACR) considers certification, continuing education, and experience in the appropriate subfield(s) to

\[1\] At institutions in which there is joint (dual) credentialing across departments doing like procedures, this substantiation of experience should be done by the chairs of both departments to ensure equity of experience among practitioners when their training backgrounds differ [43].
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 the American Board of Medical Physics (ABMP).

The appropriate subfield in medical physics for this guideline is Diagnostic Medical Physics. (Previous medical physics certification categories including Radiological Physics, Diagnostic Radiological Physics, and Diagnostic Imaging Physics are also acceptable.)

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

D. Registered Radiologist Assistant

A registered radiologist assistant is an advanced level radiographer who is certified and registered as a radiologist assistant by the American Registry of Radiologic Technologists (ARRT) after having successfully completed an advanced academic program encompassing an ACR/ASRT (American Society of Radiologic Technologists) radiologist assistant curriculum and a radiologist-directed clinical preceptorship. Under radiologist supervision, the radiologist assistant may perform patient assessment, patient management and selected examinations as delineated in the Joint Policy Statement of the ACR and the ASRT titled “Radiologist Assistant: Roles and Responsibilities” and as allowed by state law. The radiologist assistant transmits to the supervising radiologists those observations that have a bearing on diagnosis. Performance of diagnostic interpretations remains outside the scope of practice of the radiologist assistant. (ACR Resolution 34, adopted in 2006)

E. Radiologic Technologist

The technologist, together with the physician and the nursing personnel, should be responsible for patient comfort. The technologist should be able to prepare and position the patient for the vertebral augmentation procedure and, together with the nurse, monitor the patient during the procedure. The technologist should obtain the imaging data in a manner prescribed by the supervising physician. The technologist should also perform regular quality control testing of the equipment under the supervision of the Qualified Medical Physicist.

The technologist should have appropriate training and experience in the vertebral augmentation procedure and be certified by the American Registry of Radiologic Technologists (ARRT) and/or have an unrestricted state license.

F. Nursing Services

Nursing services are an integral part of the team for perioperative patient management and education and may assist the physician in monitoring the patient during the vertebral augmentation procedure.

VI. SPECIFICATIONS OF THE PROCEDURE

A. Technical Requirements

Vertebral augmentation may be performed with either fluoroscopy or CT imaging guidance. The choice is a matter of operator preference and patient characteristics. In either case, there are several technical requirements to ensure safe and successful vertebral augmentations. These include adequate institutional facilities, imaging and monitoring equipment, and support personnel. The following are minimum requirements for any institution in which vertebral augmentation is to be performed:

1. A procedural suite large enough to allow safe and easy transfer of the patient from bed to procedural table with sufficient space for appropriate positioning of patient monitoring equipment, anesthesia equipment, respirators, etc. There should be adequate space for the operating team to work unencumbered on either side of the patient and for the circulation of other staff within the room without contaminating the sterile conditions.

2. The majority of these procedures are performed under fluoroscopic guidance. A high-resolution image intensifier or flat-panel detector and video system with adequate shielding, capable of rapid imaging in orthogonal planes, and with capabilities for permanent image recording is strongly recommended. The fluoroscope should be compliant with IEC 601-2-43 [45]. Imaging findings are acquired and stored either on conventional film or digitally on computerized storage media. Imaging and image recording must be consistent with the as-low-as-reasonably-achievable (ALARA) radiation safety guidelines.

3. Immediate access to CT and rapid (within 30 or 45 minutes) access to MRI is necessary to evaluate potential complications. This may be particularly important if vertebral augmentation is planned in patients with osteolytic vertebral metastasis and/or with significant pre-existing spinal canal compromise.
4. The facility must provide adequate resources for observing patients during and after vertebral augmentation. Physiologic monitoring devices appropriate to the patient’s needs – including blood pressure monitoring, pulse oximetry, and electrocardiography – and equipment for cardiopulmonary resuscitation must be available in the procedural suite.

B. Surgical and Emergency Support

Although serious complications of vertebral augmentation are infrequent, there should be prompt access to surgical, interventional, and medical management of complications.

C. Patient Care

1. Preprocedural care
   a. The clinical history and findings, including the indications for the procedure, must be reviewed and recorded in the patient’s medical record by the physician performing the procedure. Specific inquiry should be made with respect to relevant medications, prior allergic reactions, and bleeding/clotting status.
   b. The vital signs and the results of physical and neurological examinations must be obtained and recorded.
   c. The indication(s) for the procedure, including (if applicable) documentation of failed medical therapy, must be recorded.
   d. The indication(s) for treatment of the fracture should have documentation of imaging correlation and confirmation.

2. Procedural care
   a. Adherence to the Joint Commission’s current Universal Protocol for Preventing Wrong Site, Wrong Procedure, Wrong Person Surgery™ is required for procedures in non-operating room settings including bedside procedures.

   The organization should have processes and systems in place for reconciling differences in staff responses during the “time out.”
   b. Vital signs should be obtained at regular intervals during the course of the procedure, and a record of these measurements should be maintained.
   c. Patients undergoing vertebral augmentation must have intravenous access in place for the administration of fluids and medications as needed.
   d. If the patient receives sedation, pulse oximetry must be used. Administration of sedation for vertebral augmentation should be in accordance with the ACR–SPR Practice Guideline for Sedation/Analgesia. A registered nurse or other appropriately trained personnel should be present and have primary responsibility for monitoring the patient. A record of medication doses and times of administration should be maintained.

3. Postprocedural care
   a. A procedural note should be written in the patient’s medical record summarizing the course of the procedure and what was accomplished, any immediate complications, and the patient’s status at the conclusion of the procedure (see section VIII.A.2 below). This note may be brief if the formal report will be available within a few hours. This information should be communicated to the referring physician in a timely manner. A more detailed summary of the procedure should be written in the medical record if the formal typed report will not be on the medical record within the same day.
   b. All patients should be at bed rest and observed during the initial postprocedural period. The length of this period will depend on the patient’s medical condition.
   c. During the immediate postprocedural period, skilled nurses or other appropriately trained personnel should monitor the patient’s vital signs, urinary output, sensorium, and motor strength. Neurological status should be assessed frequently at regular intervals. Initial ambulation of the patient must be carefully supervised.
   d. The operating physician or a qualified designee (another physician or a nurse) should evaluate the patient after the initial postprocedural period, and these findings should be summarized in a progress note on the patient’s medical record. The physician or designee must be available for continuing care during hospitalization and after discharge.

VII. EQUIPMENT QUALITY CONTROL

Each facility should have documented policies and procedures for monitoring and evaluating the effective management, safety, and proper performance of imaging and interventional equipment. The quality control program should be designed to maximize the quality of the diagnostic information. This may be accomplished as part of a routine preventive maintenance program.
VIII. QUALITY IMPROVEMENT AND DOCUMENTATION

A. Documentation

Results of vertebral augmentation procedures should be monitored on a continuous basis. Records should be kept of both immediate and long-term results and complications. The number of complications should be documented. Any biopsies performed in conjunction with vertebral augmentation should be followed up to detect and record any false negative and false positive results.

A permanent record of vertebral augmentation procedures should be maintained in a retrievable image storage format.

1. Imaging labeling should include permanent identification containing:
   a. Facility name and location.
   b. Examination date.
   c. Patient’s first and last names.
   d. Patient’s identification number and/or date of birth.

2. The initial progress note and final report should include:
   a. Procedure undertaken and its purpose.
   b. Type of anesthesia used (local, moderate, deep or general).
   c. Listing of level(s) treated and amount of cement injected at each level.
   d. Evaluation of injection site and focused neurologic examination.
   e. Immediate complications, if any, including treatment and outcome.
   f. Radiation dose estimate (or fluoroscopy time and the number of images obtained on equipment that does not provide direct dosimetry information) [46-48].

3. Follow up documentation:
   a. Postprocedure evaluation to assess patient response (pain relief, mobility improvement). Standardized assessment tools such as the SF 36 and the Roland-Morris disability scale may be useful for both preoperative and postoperative patient evaluation.
   b. Evaluation of injection site and focused neurologic examination.
   c. Delayed complications, if any, including treatment and outcome.
   d. Pathology (biopsy) results, if any.
   e. Record of communications with patient and referring physician.
   f. Patient disposition.

Reporting should be in accordance with the ACR–SIR Practice Guideline for the Reporting and Archiving of Interventional Radiology Procedures.

B. Informed Consent and Procedural Risk

Informed consent or emergency administrative consent must be obtained and must comply with the ACR–SIR Practice Guideline on Informed Consent for Image-Guided Procedures [49]. Risks cited should include infection; bleeding; allergic reaction; rib or vertebral fracture; vessel injury; pneumothorax (for appropriate levels); and implanted material displacement into the adjacent epidural or paravertebral veins resulting in worsening pain or paralysis, spinal cord or nerve injury, or pulmonary complication. The potential need for immediate surgical intervention should be discussed. The possibility that the patient may not experience significant pain relief should also be discussed.

C. Success and Complication Rates and Thresholds [1-2,4-21]

Although practicing physicians should strive to achieve perfect outcomes (e.g., 100% success, 0% complications), in practice, all physicians will fall short of this ideal to a variable extent. Therefore, indicator thresholds may be used to assess the efficacy of ongoing quality improvement programs. For the purposes of these guidelines, a threshold is a specific level of an indicator that should prompt a review. Procedure thresholds or overall thresholds refer to a group of indicators for a procedure, e.g., major complications. Individual complications may also be associated with complication-specific thresholds. When measures such as indications or success rates fall below a minimum threshold, or when complication rates exceed a maximum threshold, a review should be performed to determine causes and to implement changes if necessary. For example, if the incidence of fracture of rib or other bone is one measure of the quality of vertebral augmentation, values in excess of the defined threshold (in this case <1%) should trigger a review of policies and procedures within the department to determine the causes and to implement changes to lower the incidence of the complication.

Thresholds may vary from those listed herein; for example, patient referral patterns and selection factors may dictate a different threshold value for a particular indicator at a particular institution. Therefore, setting universal thresholds is very difficult, and each department is urged to alter the thresholds as needed to higher or lower values to meet its own quality improvement program needs.

Complications can be stratified on the basis of outcome. Major complications result in admission to a hospital for
therapy (for outpatient procedures), an unplanned increase in the level of care, prolonged hospitalization, permanent adverse sequelae, or death. Minor complications result in no sequelae, but may require nominal therapy or a short hospital stay for observation (generally overnight; see Appendix A). The complication rates and thresholds described herein refer to major complications.

Routine periodic review of all cases having less than perfect outcomes is strongly encouraged. Serious complications of vertebral augmentation are infrequent. A review is therefore recommended for all instances of death, infection, or symptomatic pulmonary embolus.

Success Rates
When vertebral augmentation is performed for osteoporosis, procedure outcomes can be defined using the criteria by Hodler et al [50] with patients categorized as worse, same, better, or pain/disability gone. For the purpose of this document pain/disability gone is defined as improved. Therefore patients should be categorized as either improved, the same, or worse. This categorization should be determined with the use of a validated measurement tool.

When vertebral augmentation is performed for neoplastic involvement, success is defined as achievement of significant pain relief and/or improved mobility as measured by validated measurement tools.

Table 1: Vertebral Augmentation Success Rates [51-58]

<table>
<thead>
<tr>
<th>Complication</th>
<th>Published Success Rates</th>
<th>Threshold for Review</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neoplastic, all causes</td>
<td>70% to 92%</td>
<td>&lt;60%</td>
</tr>
<tr>
<td>Osteoporosis, all causes</td>
<td>80% to 95%</td>
<td>&lt;70%</td>
</tr>
</tbody>
</table>

Complications

Major complications occur in less than 1% of patients treated for compression fractures secondary to osteoporosis and in less than 5% of treated patients with neoplastic involvement. Published complications rates and suggested thresholds are given below.

Published rates for individual types of complications are highly dependent on patient selection and are based on series comprising several hundred patients, which is a volume larger than most individual practitioners are likely to treat. It is also recognized that a single complication can cause a rate to cross above a complication-specific threshold when the complication occurs in a small volume of patients, e.g., early in a quality improvement program. In this situation, the suggested threshold is more appropriate for use in a quality-improvement program than is the published rate.

Table 2: Specific Complications for Vertebral Augmentation [52-53,59-63]

<table>
<thead>
<tr>
<th>Specific Complication</th>
<th>Published Rates</th>
<th>Thresholds for Review</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transient neurological deficit (within 30 days of the procedure)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Osteoporosis</td>
<td>1%</td>
<td>&gt; 2%</td>
</tr>
<tr>
<td>Neoplasm</td>
<td>10%</td>
<td>&gt;10%</td>
</tr>
<tr>
<td>Permanent neurological deficit (within 30 days of the procedure or requiring surgery)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Osteoporosis</td>
<td>&lt;1%</td>
<td>&gt;1%</td>
</tr>
<tr>
<td>Neoplasm</td>
<td>2%</td>
<td>&gt;5%</td>
</tr>
<tr>
<td>Fracture of rib, sternum or vertebra</td>
<td>1%</td>
<td>&gt;2%</td>
</tr>
<tr>
<td>Allergic or idiosyncratic reaction</td>
<td>&lt;1%</td>
<td>&gt;1%</td>
</tr>
<tr>
<td>Infection</td>
<td>&lt;1%</td>
<td>&gt;0%</td>
</tr>
<tr>
<td>Symptomatic pulmonary material embolus</td>
<td>&lt;1%</td>
<td>&gt;0%</td>
</tr>
<tr>
<td>Significant hemorrhage or vascular injury</td>
<td>&lt;1%</td>
<td>&gt;0%</td>
</tr>
<tr>
<td>Symptomatic hemothorax or pneumothorax</td>
<td>&lt;1%</td>
<td>&gt;0%</td>
</tr>
<tr>
<td>Death</td>
<td>&lt;1%</td>
<td>&gt;0%</td>
</tr>
</tbody>
</table>

The overall procedure threshold for all complications resulting from vertebral augmentation performed for osteoporosis is 2%, and when performed for neoplastic indications it is 10%.
IX. RADIATION SAFETY IN IMAGING

Radiologists, medical physicists, radiologic technologists, and all supervising physicians have a responsibility to minimize radiation dose to individual patients, to staff, and to society as a whole, while maintaining the necessary diagnostic image quality. This concept is known as “as low as reasonably achievable (ALARA).”

Facilities, in consultation with the medical physicist, should have in place and should adhere to policies and procedures, in accordance with ALARA, to vary examination protocols to take into account patient body habitus, such as height and/or weight, body mass index or lateral width. The dose reduction devices that are available on imaging equipment should be active; if not; manual techniques should be used to moderate the exposure while maintaining the necessary diagnostic image quality. Periodically, radiation exposures should be measured and patient radiation doses estimated by a medical physicist in accordance with the appropriate ACR Technical Standard. (ACR Resolution 17, adopted in 2006 – revised in 2009, Resolution 11.)

X. QUALITY CONTROL AND IMPROVEMENT, SAFETY, INFECTION CONTROL, 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 web site (http://www.acr.org/guidelines).

ACKNOWLEDGEMENTS

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Appendix A

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Classification of Complications by Outcome

Minor Complications

A. No therapy, no consequence.
B. Nominal therapy, no consequence; includes overnight admission for observation only.

Major Complications

C. Require therapy, minor hospitalization (<48 hours).
D. Require major therapy, unplanned increase in level of care, prolonged hospitalization (>48 hours).
E. Have permanent adverse sequelae.
F. Result in death

*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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