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 2018 (Resolution 2)*

ACR PRACTICE PARAMETER FOR THE PERFORMANCE OF FLUOROSCOPIC CONTRAST ENEMA EXAMINATION IN ADULTS

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 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

The radiographic examination of the colon by single-contrast or double-contrast technique is a proven and useful procedure. The purpose of this examination is to establish the presence or absence of disease and its nature by distending the colonic lumen and the coating of the mucosa of the colon. The goal is to obtain a diagnostic quality study by visualizing the colon in multiple projections with the minimum radiation dose necessary.

II. INDICATIONS [1-3] AND CONTRAINDICATIONS

A. The indications for a fluoroscopic contrast enema examination include, but are not limited to:

1. Diverticular disease
2. Inflammatory bowel disease
3. Colon cancer screening
5. Distal intestinal obstruction syndrome or meconium ileus equivalent in cystic fibrosis patients [5,6]
6. Evaluation of questionable findings on other imaging examinations such as computed tomography (CT)
7. Colonic volvulus
8. Assessing integrity of rectal anastomosis prior to take down of diverting colostomy or ileostomy
9. Assessment of possible colonic fistulae
10. Diseases involving the colon with familial inheritance pattern
11. Perioperative evaluation of the colon for surgical planning and follow-up
12. History of previous colon polyp or neoplasm
13. Bowel fistulas

The fluoroscopic contrast enema may also be helpful in diagnosing almost all disease states intrinsically or extrinsically affecting the colon.

B. Pertinent symptoms for the fluoroscopic contrast enema examination include, but are not limited to:

1. Abdominal pain
2. Diarrhea
3. Constipation
4. Other changes in bowel habits
5. Gastrointestinal bleeding (only if colonoscopy is not available or cannot be performed)
6. Anemia (only if colonoscopy is not available or cannot be performed)
7. Abdominal masses
8. Intestinal obstruction
9. Weight loss
10. Fever or sepsis

C. The possible contraindications for a fluoroscopic contrast enema examination include, but are not limited to:

1. Unexplained pneumoperitoneum or pneumoretroperitoneum
2. Acute colitis, including toxic megacolon
3. Combative, uncooperative patient
4. In the setting of recent endoscopic intervention, there should be a 7-day interval between the fluoroscopic contrast enema examination and the performance of large forceps biopsy through a rigid colonoscope or proctoscope, snare polypectomy, hot biopsy, or biopsy of any size or type in infectious or active inflammatory bowel disease.

For the pregnant or potentially pregnant patient, see the ACR–SPR Practice Parameter for Imaging Pregnant or Potentially Pregnant Adolescents and Women with Ionizing Radiation [7].
III. QUALIFICATIONS OF PERSONNEL

For qualifications of physicians, registered radiologist assistants, radiologic technologist, and other ancillary personnel see the ACR–AAPM Technical Standard for Management of the Use of Radiation in Fluoroscopic Procedures [8].

IV. SPECIFICATIONS OF EXAMINATION

The written or electronic request for a fluoroscopic contrast enema examination should provide sufficient information to demonstrate the medical necessity of the examination and allow for its proper performance and interpretation.

Documentation that satisfies medical necessity includes 1) signs and symptoms and/or 2) relevant history (including known diagnoses). Additional information regarding the specific reason for the examination or a provisional diagnosis would be helpful and may at times be needed to allow for the proper performance and interpretation of the examination.

The request for the examination must be originated by a physician or other appropriately licensed health care provider. The accompanying clinical information should be provided by a physician or other appropriately licensed health care provider familiar with the patient’s clinical problem or question and consistent with the state scope of practice requirements. (ACR Resolution 35, adopted in 2006 – revised in 2016, Resolution 12-b)

A. Colon Preparation

The preparation should consist of an effective combination of dietary restriction, hydration, osmotic laxatives, contact laxatives, and cleansing enemas. These preparations are intended to rid the colon of fecal material and excess fluid as much as possible. In appropriate clinical situations, preparation may be limited and, in the setting of suspected bowel obstruction or colonic volvulus, should be omitted [10-12]. There is also no routine need for colonic preparation in case of existing ileal or colonic diversion.

B. Examination Preliminaries

1. An appropriate medical history should be available, including results of laboratory tests and imaging, endoscopic, and surgical procedures as applicable.

2. The enema tip should be inserted by a physician or a trained assistant (eg, technologist, radiologist assistant, nurse, or physician assistant). A retention cuff may be used. It should be inflated carefully in accordance with the manufacturer’s guidelines and under fluoroscopic guidance and after instillation of a small amount of barium for better visualization of the balloon whenever possible. A retention cuff should be avoided for recent low rectal anastomoses (in rare instances it may be inflated under extreme care and under strict fluoroscopic guidance to avoid anastomotic dehiscence), following pelvic radiation therapy and in chronic inflammatory bowel disease.

3. Medications (eg, glucagon) may be administered to facilitate the examination.

C. Examination Technique

The following fluoroscopic contrast examination procedures should be tailored by the physician to the individual patient, as warranted by clinical circumstances and the condition of the patient, to produce a diagnostic-quality examination [1,10,11].

1. Single-contrast examination
   a. A sufficient volume of an appropriate low-density (ie, 15% to 25% weight/volume) barium suspension or water-soluble iodinated contrast should be administered to provide colonic distention.
In early postsurgical patients, if perforation is suspected or if preparation is contraindicated or not possible for other reasons, water-soluble contrast should be used. Blind-ending colonic segments (eg, rectal remnant following the Hartmann procedure or J-pouch) may also be studied with water-soluble contrast. Water-soluble contrast contains 300 to 370 mg of iodine/mL, equivalent to 60% to 76% density. It may be diluted with water to 20% to 30%, depending on the indication. Water-soluble contrast is also recommended in patients with suspected colonic obstruction or volvulus.

b. For barium studies, kilovoltage of 100 kVp or greater should be used (depending on patient size) during image acquisition. A lower kVp of 70 to 80 optimizes iodine contrast visualization on water-soluble contrast studies.

c. Manual or mechanical compression should be applied as appropriate to all accessible segments of the colon during fluoroscopy.

d. Spot large-format images should demonstrate all fluoroscopically identified suspicious findings as well as those segments of the colon in profile that may not routinely be demonstrated on overhead projections.

e. Images should include frontal and oblique views of the entire filled colon, an angled-beam view of the sigmoid colon, and a lateral view of the rectum. Whenever possible, the lateral rectal view should include an image obtained after the enema tip has been removed.

f. Postevacuation images should be obtained when possible and should always be obtained in the evaluation for leak.

g. The quality assurance indicators specific to the single-contrast enema examination are:
   i. Compression views may be helpful
   ii. Each accessible segment of the colon is seen during fluoroscopy
   iii. Each segment of the entire colon should be seen without overlap, if possible
   iv. Imaging technique should optimize visualization of all segments of the colon
   v. Complete visualization of the entire colon should be ensured through demonstration of ileocecal valve, terminal ileum, or appendix

h. In the setting of distal intestinal obstruction syndrome/meconium ileus equivalent in patients with cystic fibrosis, a water-soluble contrast enema examination can demonstrate the level of the obstruction and possibly be therapeutic. The water-soluble contrast material enema procedure has become an accepted supplement to other nonsurgical therapeutic measures, and multiple enemas with water-soluble contrast agents over several days may be required to mobilize the tenacious stool plugs [5,6]. Repeat enemas in this setting may be performed without fluoroscopic guidance.

2. Double-contrast barium examination
   a. Commercially prepared high-density (80% weight/volume or greater) barium suspension is used.
   b. Kilovoltage of 90 kVp or greater, depending on the patient’s size, is used.
   c. Barium suspension and air are introduced under fluoroscopic control to achieve adequate coating and distention of the entire colon.
   d. The entire colon should be examined fluoroscopically during the course of the examination.
   e. Images should be taken to attempt to demonstrate all segments of the colon in double contrast. Suggested views include the following:
      i. Spot images of the rectum, sigmoid colon, flexures, and cecum in double contrast
      ii. Large-format images, including prone and supine views of the entire colon, an angled-beam view of the sigmoid colon, and a lateral view of the rectum, either cross-table lateral or vertical beam, preferably with the enema tip removed
      iii. Both lateral decubitus views of the entire colon using a horizontal beam (a wedge filter is recommended)
      iv. Erect or semierect flexure views, and postevacuation views, when possible, may be helpful
   f. The quality assurance indicators specific to the double-contrast barium enema examination are as follows:
      i. Adequate barium coating of the entire colon has been achieved
      ii. The colon is well distended with air
      iii. Each segment of the colon is seen in double contrast on at least 2 images taken in different positions, whenever possible
iv. Complete visualization of the entire colon is ensured through demonstration of the ileocecal valve, terminal ileum, or appendix

3. Colostomy or colonic mucous fistula fluoroscopic contrast enema

a. These procedures are indicated when disease is suspected involving a colostomy or colonic mucous fistula or to delineate anatomy in preparation for colostomy revision/takedown. The ostomy should be examined by the radiologist or a trained assistant. An appropriate device should be inserted into the ostomy. Examples of appropriate devices include, but are not limited to:
   i. Foley catheter
   ii. Red rubber catheter
   iii. Cone colostomy tip

   If a Foley catheter is used, the balloon should be inflated on the outside of the stoma and held firmly against the stoma by the patient’s gloved hand. Alternatively the Foley balloon may be inflated under care inside the stoma and under strict fluoroscopic guidance to avoid injury.

b. Low-density barium or water-soluble contrast should be instilled into the ostomy through the device under fluoroscopic observation. The examination should attempt to answer the clinical question and should be recorded on spot radiographic images.

D. Quality Assurance

1. The following quality assurance indicators should be applied as appropriate to all fluoroscopic contrast enema examinations:
   a. Colon preparation should be adequate for the clinical indication.
   b. When examinations are completed, patients should be held in the fluoroscopic area until the physician has reviewed the images.
   c. An attempt should be made to resolve questionable radiologic findings before the patient leaves. Repeat fluoroscopy of the patient should be performed as necessary.

2. The following steps are suggested for a quality assurance and continuing quality improvement program:
   a. Correlation of radiologic, endoscopic, and pathologic findings
   b. In high volume centers, determination of detection rates for colorectal cancer and polyps measuring 1 cm or greater

V. DOCUMENTATION

Reporting should be in accordance with the ACR Practice Parameter for Communication of Diagnostic Imaging Findings [13].

VI. EQUIPMENT SPECIFICATIONS

Examinations should be performed with fluoroscopic image intensification and radiographic equipment that meets all applicable federal and state radiation standards.

Equipment should provide diagnostic fluoroscopic image quality and recording (image, video, or digital) capability. Equipment should be capable of producing kilovoltage greater than 100 kVp. Equipment necessary to compress and isolate regions of the colon should be readily available.
VII. RADIATION SAFETY IN IMAGING

Radiologists, medical physicists, registered radiologist assistants, radiologic technologists, and all supervising physicians have a responsibility for safety in the workplace by keeping radiation exposure to staff, and to society as a whole, “as low as reasonably achievable” (ALARA) and to assure that radiation doses to individual patients are appropriate, taking into account the possible risk from radiation exposure and the diagnostic image quality necessary to achieve the clinical objective. All personnel that work with ionizing radiation must understand the key principles of occupational and public radiation protection (justification, optimization of protection and application of dose limits) and the principles of proper management of radiation dose to patients (justification, optimization and the use of dose reference levels)


Nationally developed guidelines, such as the ACR’s Appropriateness Criteria®, should be used to help choose the most appropriate imaging procedures to prevent unwarranted radiation exposure.

Facilities should have and adhere to policies and procedures that require varying ionizing radiation examination protocols (plain radiography, fluoroscopy, interventional radiology, CT) to take into account patient body habitus (such as patient dimensions, weight, or body mass index) to optimize the relationship between minimal radiation dose and adequate image quality. Automated dose reduction technologies available on imaging equipment should be used whenever appropriate. If such technology is not available, appropriate manual techniques should be used.

Additional information regarding patient radiation safety in imaging is available at the Image Gently® for children (www.imagegently.org) and Image Wisely® for adults (www.imagewisely.org) websites. These advocacy and awareness campaigns provide free educational materials for all stakeholders involved in imaging (patients, technologists, referring providers, medical physicists, and radiologists).

Radiation exposures or other dose indices should be measured and patient radiation dose estimated for representative examinations and types of patients by a Qualified Medical Physicist in accordance with the applicable ACR Technical Standards. Regular auditing of patient dose indices should be performed by comparing the facility’s dose information with national benchmarks, such as the ACR Dose Index Registry, the NCRP Report No. 172, Reference Levels and Achievable Doses in Medical and Dental Imaging: Recommendations for the United States or the Conference of Radiation Control Program Director’s National Evaluation of X-ray Trends. (ACR Resolution 17 adopted in 2006 – revised in 2009, 2013, Resolution 52).

VIII. 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 website (https://www.acr.org/Clinical-Resources/Practice-Parameters-and-Technical-Standards).

ACKNOWLEDGEMENTS

This practice parameter was revised according to the process described under the heading The Process for Developing ACR Practice Guidelines and Technical Standards on the ACR website (https://www.acr.org/Clinical-Resources/Practice-Parameters-and-Technical-Standards) by the Committee on Practice Parameters – Body Imaging (Abdominal) of the ACR Commission on Body Imaging and Committee on Practice Parameters – General, Small, Emergency and/or Rural Practice of the ACR Commission on General, Small, Emergency and/or Rural Practice.

Reviewing Committee
Richard M. Gore, MD, FACR, Chair
Mahmoud M. Al-Hawary, MD
Patrick Gonzales, MD
Ruedi F. Thoeni, MD

Committee on Practice Parameters – General, Small, Emergency and/or Rural Practices
(ACR Committee responsible for sponsoring the draft through the process)

Sayed Ali, MD, Chair
Marco A. Amendola, MD, FACR
Lynn Broderick, MD, FACR
Resmi A. Charalel, MD
Brian D. Gale, MD, MBA
Carolyn A. Haerr, MD
Charles E. Johnson, MD
Candice Johnstone, MD
Padmaja A. Jonnalagadda, MD
Steven E. Liston, MD, MBA, FACR
Tammam Nehme, MD
Samir S. Shah, MD
Jennifer L. Tomich, MD

Committee on Body Imaging (Abdominal)
(ACR Committee responsible for sponsoring the draft through the process)

Ruedi F. Thoeni, MD, Chair
Mahmoud M. Al-Hawary, MD
Mark E. Baker, MD, FACR
Lindsay Busby MD, MPH
Barry D. Daly, MD, MB, BCh
Isaac R. Francis, MD, FACR
Patrick Gonzales, MD
Richard M. Gore, MD, FACR
Jay P. Heiken MD, FACR
Frank H. Miller, MD, FACR
Donald G. Mitchell, MD, FACR
Eric M. Rubin, MD
Scott D. Stevens, MD, FACR
William E. Torres, MD, FACR

Robert S. Pyatt, Jr, MD, FACR, Chair, 2Commission on General, Small, Emergency and/or Rural Practice
Lincoln Berland, MND, FACR, Chair, Commission on Body Imaging
Jacqueline Anne Bello, MD, FACR, Chair, Commission on Quality and Safety
Matthew S Pollack, MD, FACR, Chair, Committee on Practice Parameters & Technical Standards

Comments Reconciliation Committee
Andy Rosenkrantz, MD, Chair
Debra Dyer, MD, FACR, Co-Chair
Mahmoud M. Al-Hawary, MD
Sayed Ali, MD
Jacqueline Anne Bello, MD, FACR
Travis G. Browning, MD
Priscilla F. Butler, MS, FACR
Timothy A. Crummy, MD
Sandeep P. Deshmukh, MD

Richard Duszak, Jr., MD, FACR
Richard A Geise, PhD, FACR
Patrick Gonzales, MD
Richard M Gore, MD, FACR
Paul A. Larson, MD, FACR
Matthew S. Pollack, MD, FACR
Timothy L. Swan, MD, FACR, FSIR
Ruedi F. Thoeni, MD

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
1991 (Resolution 7)
Amended 1995 (Resolution 24, 53)
Revised 1999 (Resolution 31)
Revised 2002 (Resolution 32)
Amended 2006 (Resolution 17, 34, 35, 36)
Amended 2007 (Resolution 12m)
Revised 2008 (Resolution 37)
Amended 2009 (Resolution 11)
Revised 2013 (Resolution 25)
Amended 2014 (Resolution 39)
Revised 2018 (Resolution 2)