The IAC Standards for Carotid Stenting Accreditation
Table of Contents

Introduction .................................................................................................................................................................................... 3

Part A: Organization........................................................................................................................................................................ 4

Section 1A: Personnel and Supervision........................................................................................................................................ 4
  STANDARD – Medical Director .................................................................................................................................................... 4
  STANDARD – Medical Staff ....................................................................................................................................................... 5
  STANDARD – Interventional Technologist Technical/Administrative Director ................................................................. 6
  STANDARD – Interventional Nurse Technical/Administrative Director .............................................................................. 7
  STANDARD – Technical Staff (Interventional Technologist[s]) ............................................................................................... 8
  STANDARD – Technical Staff (Interventional Nurse[s]) ........................................................................................................ 9
  STANDARD – Neurological Assessment Examiner(s) ............................................................................................................ 10
  STANDARD – Ancillary Personnel ....................................................................................................................................... 11
  STANDARD – Medical Physicist ............................................................................................................................................ 11

Section 2A: Facility ........................................................................................................................................................................ 13
  STANDARD – Examination Areas ............................................................................................................................................ 13
  STANDARD – Interpretation Areas ......................................................................................................................................... 14
  STANDARD – Storage Space .................................................................................................................................................. 14
  STANDARD – Equipment and Instrumentation .................................................................................................................... 14
  STANDARD – Equipment and Instrumentation Quality Control ......................................................................................... 15
  STANDARD – Quality Control Documentation .................................................................................................................... 16

Section 3A: Administrative .......................................................................................................................................................... 17
  STANDARD – Patient Confidentiality ....................................................................................................................................... 17
  STANDARD – Patient or Other Customer Complaints ........................................................................................................ 17
  STANDARD – Primary Source Verification .......................................................................................................................... 17

Part B: Process ............................................................................................................................................................................... 18

Section 1B: Procedures and Protocols ......................................................................................................................................... 18
  STANDARD – Procedure Overview ................................................................................................................................. 18
  STANDARD – Procedure Requirements ............................................................................................................................ 18
  STANDARD – Procedure Volumes .................................................................................................................................... 20

Part C: Quality Improvement ................................................................................................................................................... 21

Section 1C: Quality Improvement Program ............................................................................................................................. 21
  STANDARD – QI Program ...................................................................................................................................................... 21
  STANDARD – QI Documentation ....................................................................................................................................... 21

Section 2C: Quality Improvement Measures ............................................................................................................................ 22
  STANDARD – QI Measures ................................................................................................................................................. 22
  STANDARD – Reporting of Outcomes ............................................................................................................................... 22

Section 3C: Quality Improvement Meetings ............................................................................................................................. 26
  STANDARD – QI Meetings .................................................................................................................................................... 26

Selected Bibliography ..................................................................................................................................................................... 27

Appendix .................................................................................................................................................................................... 28
Introduction

The Intersocietal Accreditation Commission (IAC) accredits imaging facilities specific to carotid stenting procedures. IAC accreditation is a means by which facilities can evaluate and demonstrate the level of patient care they provide.

This program is designed to accredit facilities that perform stenting of the extracranial carotid artery by ensuring that the facility meets benchmarks for quality based on resources, training and outcomes. Carotid stents may be appropriately placed for many indications (See Appendix), but the most common indication, and the indication for which outcome data is most widely available, is treatment of carotid bifurcation disease secondary either to atherosclerosis, post endarterectomy restenosis or radiation induced stenosis. Therefore, the outcome benchmarks used in this program are intended to be applied only to cases treated for these indications. A facility that is able to meet the outcome benchmarks for these most common indications will most likely provide adequate outcomes for carotid stenting performed for less common indications (See Appendix).

A carotid stenting facility consists of at least one designated fluoroscopy system, a qualified physician, a nurse and an interventional technologist. Each facility must have a Medical Director and Technical/Administrative Director. The facility must meet the organizational requirements defined in this document. There may be additional physicians, interventional technologists and other professional and/or ancillary personnel. When more than one technical member is employed, a Technical/Administrative Director is responsible for supervision of the technical staff. If stenting is performed in more than one location within one facility, the facility is encouraged to apply for all locations within that facility under the overall direction of a Medical Director(s). All operators and all cases under the direction of the Medical Director(s) must be included in the application for accreditation.

The intent of the accreditation process is two-fold. It is designed to recognize facilities that provide quality carotid stenting services. It is also designed to be used as an educational tool to improve the overall quality of the facility.

These accreditation Standards are the minimum standards for accreditation of carotid stenting facilities. Standards are the minimum requirements to which an accredited facility is held accountable.

Standards that are highlighted are content changes that were made as part of the April 15, 2019 revision. These Standards are effective immediately. Facilities applying for accreditation after April 15, 2019 must comply with these new highlighted Standards.

All absolute requirements appear in bolded text. In addition to all standards listed below, the facility, including all staff, must comply at all times with all federal, state and local laws and regulations, including but not limited to laws relating to licensed scope of practice, facility operations and billing requirements.
Part A: Organization

Section 1A: Personnel and Supervision

STANDARD – Medical Director

1.1A The Medical Director must be a licensed physician.

1.1.1A Medical Director Required Training and Experience

The Medical Director must demonstrate an appropriate level of training and experience by meeting the following:

1.1.1.1A board certified in his/her specialty; and

1.1.1.2A have clinical expertise in the management of extracranial carotid occlusive disease but need not personally perform cervical/extracranial carotid angioplasty and/or stenting.

1.1.2A Medical Director Responsibilities

The Medical Director is responsible for all cervical/extracranial carotid angioplasty and stenting services provided, including compliance, radiation safety, outcomes, quality control, quality of care and appropriateness of care provided. The Medical Director responsibilities include but are not limited to:

1.1.2.1A Compliance with all facility policies/procedures/protocols and will review and update all manuals periodically as necessary (minimum every three years) or as new policies are introduced. This review must be documented via signature (or initials) and date on the reviewed document or manual.

1.1.2.2A Active oversight of radiation safety within the facility as evidenced by membership on the institution’s radiation safety committee or periodic review of radiation safety issues and documentation. The radiation protection program content and implementation must be reviewed at least annually.

1.1.2.3A Delegation, when appropriate, of the supervision of radiation safety standards to the Technical/Administrative Director, radiation safety officer or health physics consultant. Records of radiation safety must be kept on file in accordance with local requirements and available for inspection.

1.1.2.4A The review and oversight of the clinical practice of cervical/extracranial carotid angioplasty and stenting services.

1.1.2.5A The Medical Director must provide oversight and documentation of comprehensive Quality Improvement (QI) Program. Reference Section 1C: Quality Improvement Program.

Comment: The Medical Director may supervise the entire operation of the facility or delegate specific operations but is responsible for assuring compliance of medical and technical staff to the standards outlined in this document.
1.1.3A Continuing Medical Education (CME) Requirements

1.1.3.1A The Medical Director must obtain at least 15 hours of Category I CME credits, relevant to cerebrovascular disease that includes but is not limited to content that is directly related to the performance of cervical/extracranial carotid angioplasty and/or stenting and/or carotid atherosclerotic disease every three years. Radiation safety training must be part of the CME and not be less than one hour of the 15 hours required. If the Medical Director performs these procedures, he/she must meet the qualifications and maintenance of qualifications of the medical staff.

Comment: If the Medical Director has successfully attained one or more of the following within the three years prior to the application date, the CME requirement will be considered fulfilled:

i. completion of an Accreditation Council for Graduate Medical Education (ACGME) approved (or similarly recognized) residency or fellowship; or
ii. attaining certification by an American Board of Medical Specialties (ABMS) recognized board.

1.1.3.2A Documentation of CME credits must be kept on file and available for inspection.

STANDARD – Medical Staff

1.2A All members of the medical staff must be licensed physicians.

1.2.1A Medical Staff Required Training and Experience

The medical staff must demonstrate an appropriate level of training and experience by meeting the following:

1.2.1.1A The medical staff member(s) must meet one of the published national society training standards pertaining to cervical/extracranial carotid angioplasty and stenting and be credentialed by the health care facility to perform cervical/extracranial carotid angioplasty and stenting. The currently acceptable national society training standards are:


iv. Other national society training standards may be considered appropriate subject to review and approval by the IAC Carotid Stenting Board of Directors.

1.2.1.2A All physicians (including the Medical Director, if applicable) performing cervical/extracranial carotid angioplasty and stenting must be privileged by clear and concise requirements as outlined by their hospital privileging committee that include periodic review and documentation of credentialed staff.
1.2.2A Continuing Medical Education (CME) Requirements

1.2.2.1A The medical staff members must obtain at least 20 hours of Category 1 CME credits, relevant to percutaneous therapeutic endovascular intervention and cerebrovascular disease every three years. Of these, 10 credit hours should be relevant to cervical/extracranial carotid angioplasty and stenting. Radiation safety training must be part of the CME and not be less than one hour of the 20 hours required.

Comment: If the medical staff member has successfully attained one or more of the following within the three years prior to the application date, the CME requirement will be considered fulfilled:

i. completion of an Accreditation Council for Graduate Medical Education (ACGME) approved (or similarly recognized) residency or fellowship; or
ii. certification by an American Board of Medical Specialties (ABMS) recognized board.

1.2.2.2A Documentation of CME credits must be kept on file and available for inspection.

STANDARD – Interventional Technologist Technical/Administrative Director

1.3A The Technical/Administrative Director must be either an interventional technologist or interventional nurse (1.4A) and meet the required training and experience qualifications as outlined.

1.3.1A Interventional Technologist Technical/Administrative Director Required Training and Experience

The Interventional Technologist Technical/Administrative Director must demonstrate an appropriate level of training and experience by meeting one the following criteria:

1.3.1.1A A registered radiologic technologist with the American Registry of Radiologic Technologists (ARRT) or the Canadian Association of Medical Radiation Technologists (CAMRT) with post primary certification in one of the following:

i. Cardiac-Interventional Radiography RT(CI);
ii. Vascular-Interventional Radiography RT(VI);
iii. Cardiovascular-Interventional Radiography RT(CV); or
iv. Registered Cardiovascular Invasive Specialist (RCIS).

1.3.1.2A A registered radiologic technologist with the American Registry of Radiologic Technologists (ARRT) or the Canadian Association of Medical Radiation Technologists (CAMRT) with a minimum of five years of experience performing interventional, vascular or cardiology procedures. A letter from the Medical Director or supervising physician verifying the training, experience and competency in performance and supervision of CAS procedures is required.

Comment: In the event that the Technical Director applying under pathway 1.3.1.2A no longer works in this capacity, the newly appointed Technical Director must meet training pathway 1.3.1.1A.

1.3.2A Interventional Technologist Technical/Administrative Director Responsibilities

The Interventional Technologist Technical/Administrative Director responsibilities may include, but are not limited to:

1.3.2.1A the day-to-day operations of the facility;
1.3.2.2A the delegation, when necessary, of specific responsibilities to the technical and/or ancillary staff; and
1.3.2.3A verification of documentation of proper training and, at least annually, assessment of the competence of technical staff and/or any ancillary staff who report to the Technical/Administrative Director.

1.3.3A Continuing Education (CE) Requirements

1.3.3.1A The Interventional Technologist Technical/Administrative Director must obtain at least 15 hours of accredited CE in percutaneous interventional procedures or patient management every three years. Radiation safety training must be part of the CE and not be less than one hour of the 15 hours required.

1.3.3.2A All CE hours must be approved (i.e., Recognized Continuing Education Evaluation Mechanism (RECEEM), American Registry of Radiologic Technologists (ARRT)-Category A, American Society of Radiologic Technologists (ASRT), American Medical Association (AMA), American Nurses Credentialing Center (ANCC)-Category I).

Comment: If the Technical/Administrative Director has successfully attained an appropriate technical credential within the three years, prior to the application date, the CE requirement hours will be considered fulfilled.

1.3.3.3A Documentation of CE credits must be kept on file and available for inspection.

STANDARD – Interventional Nurse Technical/Administrative Director

1.4A The Technical/Administrative Director must be either an interventional technologist (1.3A) or interventional nurse and meet the required training and experience qualifications as outlined.

1.4.1A Interventional Nurse Technical/Administrative Director Required Training and Experience

1.4.1.1A The Interventional Nurse Technical/Administrative Director must demonstrate an appropriate level of training and experience by meeting one of the following criteria:

i. Registered Nurse (RN);
ii. advanced health care degree or Bachelor of Science in nursing (BSN) preferred; or
iii. certification in interventional nursing specialty such as Cardiac Vascular Nursing (CVRN), Cardiac Vascular Invasive Specialist (CVIS) or Certified Radiology Nurse (CRN).

1.4.1.2A Critical care or emergency room experience is required.

1.4.1.3A At least six months of critical care or emergency room nursing is required.

1.4.1.4A Basic Life Support (BLS) and Advanced Cardiac Life Support (ACLS) certification is required.

1.4.2A Interventional Nurse Technical/Administrative Director Responsibilities

The Interventional Nurse Technical/Administrative Director responsibilities may include, but are not limited to:

1.4.2.1A the day-to-day operations of the facility;

1.4.2.2A the delegation, when necessary, of specific responsibilities to the technical and/or ancillary staff; and
1.4.2.3A  verification of documentation of proper training and, at least annually, assessment of the competence of technical staff and/or any ancillary staff who report to the Technical/Administrative Director.

1.4.3A  Continuing Education (CE) Requirements

1.4.3.1A  The interventional nursing staff must obtain at least 15 hours of accredited CE in percutaneous interventional procedures, neurologic assessment and/or patient management, every three years. Radiation safety training must be part of the CE and not be less than one hour of the 15 hours required.

1.4.3.2A  All CE hours must be American Nurses Credentialing Center (ANCC) approved. At least one contact hour in moderate sedation is required annually.
Comment: If the nursing staff member has successfully attained an appropriate specialty certification (CVRN, CVIS or CRN) within the three years prior to the application date, the CE requirement will be considered fulfilled.

1.4.3.3A  Documentation of CE credits must be kept on file and available for inspection.

STANDARD – Technical Staff (Interventional Technologist[s])

1.5A  Interventional technologist(s) at the facility must meet the following qualifications:

1.5.1A  Interventional Technologist(s) Required Training and Experience

The interventional technologist(s) must meet one of the following criteria:

1.5.1.1A  A registered radiologic technologist with the American Registry of Radiologic Technologists (ARRT) or the Canadian Association of Medical Radiation Technologists (CAMRT) with post primary certification in one of the following:

i.  Cardiac-Interventional Radiography RT(CI);
ii.  Vascular-Interventional Radiography RT(VI);
iii.  Cardiovascular-Interventional Radiography RT(CV); or
iv.  Registered Cardiovascular Invasive Specialist (RCIS).

1.5.1.2A  A registered radiologic technologist [RT(R)] with a minimum of one year of full-time equivalent experience as an interventional technologist under the direct supervision of personnel meeting pathway 1.3.1A, as indicated above. A clinical rotation in interventional, cardiology, vascular or invasive procedures as part of their educational program may be counted for up to six months of clinical experience.

1.5.2A  Interventional Technologist(s) Responsibilities

The interventional technologist(s) responsibilities may include, but are not limited to:

1.5.2.1A  reporting to the Technical/Administrative Director;

1.5.2.2A  reviewing and/or recording pertinent patient history and supporting clinical data;

1.5.2.3A  obtaining a record of anatomical, pathological and/or physiological data for interpretation by the physician;

1.5.2.4A  positioning of the patient, selection of radiation exposure parameters, imaging of the patient and archiving of the images;
1.5.2.5A maintaining a high degree of awareness of all radiation and patient safety issues involved with any invasive procedure;

1.5.2.6A demonstrating a thorough understanding and working knowledge of normal and abnormal anatomy, physiology, radiation safety, interventional supplies and equipment operation;

1.5.2.7A recognizing and resolving equipment problems and discrepancies, anticipating patient needs and concerns and communicating the appropriate care needed;

1.5.2.8A using professional judgment and critical thinking when performing procedures;

1.5.2.9A scrubbing in and assisting the physician in the procedure when necessary;

1.5.2.10A circulating within the procedure room and procuring equipment needed for any given procedure; and

1.5.2.11A performing other procedures and duties, as assigned.

1.5.3A Continuing Education (CE) Requirements

1.5.3.1A The interventional technologist staff must obtain at least 15 hours of accredited CE in percutaneous interventional procedures or patient management, every three years. Radiation safety training must be part of the CE and not be less than one hour of the 15 hours required.

1.5.3.2A All CE hours must be approved CE (i.e., RECEEM, ARRT-Category A, ASRT, AMA Category I).

Comment: If the interventional technologist staff member has successfully attained an appropriate technical credential within the three years prior to the application date, the CE requirement will be considered fulfilled.

1.5.3.3A Documentation of CE credits must be kept on file and available for inspection.

STANDARD – Technical Staff (Interventional Nurse[s])

1.6A Interventional nurse(s) at the facility must meet the following qualifications:

1.6.1A Interventional Nurse(s) Required Training and Experience

1.6.1.1A The interventional nurse(s) must meet one of the following criteria:

i. Registered Nurse (RN);

ii. advanced health care degree or Bachelor of Science in Nursing (BSN) preferred; or

iii. certification in interventional nursing specialty such as Cardiac Vascular Nursing (CVRN), Cardiac Vascular Invasive Specialist (CVIS) or Certified Radiology Nurse (CRN).

1.6.1.2A Critical care or emergency room experience is required.

1.6.1.3A At least six months of critical care or emergency room nursing is required.

1.6.1.4A Basic Life Support (BLS) and Advanced Cardiac Life Support (ACLS) certification is required.
1.6.2A Interventional Nurse(s) Responsibilities

The interventional nurse(s) responsibilities may include, but are not limited to:

1.6.2.1A administering and monitoring moderate sedation;
1.6.2.2A performing neurological assessment;
1.6.2.3A knowing relevant radiation safety;
1.6.2.4A monitoring and assessing clinical status of patient;
1.6.2.5A cardiovascular and hemodynamic monitoring and management; or
1.6.2.6A advising patient care team and treating patient appropriately.

1.6.3A Continuing Education (CE) Requirements

1.6.3.1A The interventional nursing staff must obtain at least 15 hours of accredited CE in percutaneous interventional procedures, neurologic assessment and/or patient management, every three years. Radiation safety training must be part of the CE and not be less than one hour of the 15 hours required.

1.6.3.2A All CE hours must be American Nurses Credentialing Center (ANCC) approved. At least one contact hour in moderate sedation is required annually.

Comment: If the nursing staff member has successfully attained an appropriate specialty certification CVRN, CVIS or CRN within the three years prior to the application date, the CE requirement will be considered fulfilled.

1.6.3.3A Documentation of CE credits must be kept on file and available for inspection.

STANDARD – Neurological Assessment Examiner(s)

1.7A Neurological assessment examiner(s) at the facility must meet the following qualifications:

1.7.1A Neurological Assessment Examiner(s) Required Training and Experience

The neurological assessment examiner(s) must demonstrate an appropriate level of training and experience by meeting the following criteria:

1.7.1.1A Independent neurologic assessments (to include NIHSS and mRS) must be performed by a physician, physician extender, nurses or other qualified health care professionals as allowed by state law (i.e., physician assistant (PA) or nurse practitioner (NP) with neurological expertise). The independent examiner is defined as the person performing the assessment that is not the operator or any other person listed on the procedure documentation. Optimally, these examinations are performed by a neurologist.

1.7.1.2A The National Institutes of Health Stroke Scale (NIHSS) and Modified Rankin Scale (mRS) must be performed by personnel meeting requirements in 1.7.1.1A performed by physicians that have completed formal training and demonstrated competency in performing NIH Stroke and Modified Rankin scales. All examiners must have current NIHSS and Modified Rankin certification by a nationally recognized certification organization recognized by organizations such as the National Stroke Association or the American Stroke Association. Following initial certification, initial recertification is required at six months and then annually.
i. Documentation of current NIHSS and Modified Rankin certification must be kept on file and available for inspection.

STANDARD – Ancillary Personnel

1.8A The facility must ensure that adequately trained and experienced ancillary personnel are available to perform safe and effective patient care appropriate for the level of service as designated by the Medical or Technical/Administrative Director. The specific needs of a facility must be determined by an evaluation of the types and volumes of procedures as well as facility configuration.

1.8.1A Ancillary personnel may consist of, but are not limited to:

1.8.1.1A anesthesia personnel;
1.8.1.2A technical assistants;
1.8.1.3A clerical and administrative assistants;
1.8.1.4A computer support staff; or
1.8.1.5A equipment support staff (i.e., biomedical, x-ray service).

1.8.2A All ancillary personnel within the department must be supervised by the Medical Director or a qualified designee. The supervisor must document/verify proper training, at least annually and current competence of the ancillary personnel appropriate to the assigned duties.

STANDARD – Medical Physicist

1.9A A qualified medical physicist must be appointed for the facility and meet the following qualifications:

1.9.1A Medical Physicist Required Training and Experience

The medical physicist(s) must meet one of the following criteria:

1.9.1.1A Board certification by the American Board of Radiology (ABR), the American Board of Medical Physics (ABMP) or the Canadian College of Medical Physics (CCMP) in a discipline that includes diagnostic imaging is recommended.

1.9.1.2A A physicist who has passed Part 2 of the ABR examination in a discipline of medical physics that includes diagnostic imaging is acceptable. Full certification by a recognized board as outlined above is required prior to the next accreditation cycle.

1.9.1.3A Licensed or certified in accordance with state and local regulations. Full certification by a recognized board as outlined above is required prior to the next accreditation cycle. Individuals listed in the National QMP Registry maintained by the Conference of Radiation Control Program Directors for a subspecialty of medical physics in diagnostic imaging are acceptable.

1.9.2A Medical Physicist Responsibilities

The medical physicist(s) responsibilities may include, but are not limited to:

1.9.2.1A Perform initial and annual surveys for equipment performance evaluation including:

i. radiation output measurements;
ii. system quality control tests;
iii. image quality performance measurements; and
iv. analyze all data with appropriate recommendations.

1.9.2.2A Provide a written summary of all assessment and evaluations performed.
1.9.2.3A Provide guidance for any patient and/or staff dosimetry issues.
1.9.2.4A Provide radiation training for facility physicians and staff as required.
1.9.2.5A Other personnel, deemed by the medical physicist as competent to perform the assigned tasks, may assist the medical physicist in the collection of data under the direct supervision of the medical physicist. The medical physicist must review and approve all such data. The medical physicist remains personally responsible for the performance quality of the assigned tasks.
1.9.2.6A It is recommended that the physicist observe at least one (interventional) procedure per year.

1.9.3A Continuing Education (CE) Requirements

1.9.3.1A The medical physicist must obtain at least 15 credits hours of CE approved by the Commission on Accreditation of Medical Physics Education Program (CAMPEP) in diagnostic imaging including fluoroscopy, every three years. The 15 CAMPEP hours should include education in radiation dosimetry, radiation protection and equipment performance related to the use of fluoroscopy. The medical physicist should regularly perform a sufficient number of radiation measurements, dosimetric calculations and equipment performance evaluations of fluoroscopic equipment to maintain competence in the performance of these activities.

Comment: If the medical physicist has successfully attained board certification within the three years prior to the application date, the CE requirement will be considered fulfilled.

1.9.3.2A Documentation of CAMPEP credits must be kept on file and available for inspection.
Section 2A: Facility

STANDARD – Examination Areas

2.1A Adequate facilities must be provided for all operations of the facility so that patient comfort, safety, dignity and privacy are ensured as well as staff comfort and safety. Procedure areas must have sufficient space, be well maintained and clean. There should be adequate space for the interventional personnel to freely access the patient and for all staff to maintain the sterile field. Physical space requirements include, but are not limited to:

2.1.1A waiting, reception and patient/staff bathrooms;

2.1.2A patient education, consultation and examination areas;

2.1.3A readily accessible hand washing/sanitation for staff;

2.1.4A performance of pre-test/post-procedures within appropriate proximity of the procedure area including adequate space for performing resuscitation in case of emergency;

2.1.5A there must be immediate access to computed tomography or magnetic resonance imaging to allow evaluation of possible complications;

2.1.6A adequate space, facility configuration and doorways for the emergency transport of patients from patient care areas and for emergency exit of staff; and

2.1.7A the procedure room/area must have or meet:

2.1.7.1A General anesthesia capability.

2.1.7.2A Radiation shielded barriers that meet state and federal requirements.

2.1.7.3A Patient post care areas.

2.1.7.4A Room utilities: Adequate utilities based upon the types of procedures and workload. These utilities include water taps, lighting, electrical outlets, emergency power, telephones, heating/cooling and ventilation.

2.1.7.5A Room lighting: Overhead and task lighting must be adequate to perform carotid stent procedures and for clinical evaluation and treatment of the patient. The overhead lighting must be able to be dimmed during fluoroscopy. It is recommended that the overhead lighting be controlled by a foot pedal used by the operating physician.

2.1.7.6A Room power: The facility must have a plan that outlines the response to unexpected power loss or computer function, such as movement of the patient to another carotid stent capable procedure room in the immediate vicinity.

i. When normal power is not available, emergency power should be capable of providing a minimum of 10 minutes of fluoroscopy, and at least one hour of backup power for the computers, monitoring equipment and ancillary equipment.

ii. For systems ordered after July 2011, there should be sufficient emergency power supply to run fluoroscopy for a duration of one hour and to run the remainder of the x-ray system components including lighting, for a minimum of 24 hours.

iii. Utilization of emergency power must be visible by the operator at the normal working position.
iv. X-ray equipment and computers should not require rebooting during transition between normal and emergency power or during power line instabilities.

STANDARD – Interpretation Areas

2.2A Adequate designed space must be provided for the interpretation of exam results and preparation of reports.

STANDARD – Storage Space

2.3A Adequate space must be provided for:

2.3.1A the storage must ensure confidentiality of data and should be safe from fire/flood;

2.3.2A patient records, reports and digital data storage areas;

2.3.3A administration records and support areas; and

2.3.4A equipment/supply storage areas.

STANDARD – Equipment and Instrumentation

2.4A Equipment and instrumentation used in the performance of carotid artery stenting must be appropriate, in good working condition and should substantially comply with the International Electrotechnical Commission (IEC) 60601-2-43 interventional standard.

2.4.1A All equipment and instrumentation must be routinely inspected for safety and proper functionality and records of the inspections kept on file. Equipment and instrumentation must include at a minimum the following:

2.4.1.1A A fixed (recommended) or portable, single or biplane (recommended) angiography system that must meet the following specifications:

2.4.1.2A digital subtraction imaging of at least 3 frames per second, with the ability to do program change of rate during the image acquisition run;

2.4.1.3A high quality, subtracted digital radiographs;

2.4.1.4A road-mapping with ability to refer back to an unsubtracted live image;

2.4.1.5A last image hold;

2.4.1.6A pulsed fluoroscopy;

2.4.1.7A dose measurement capability;

2.4.1.8A digital Imaging and Communications in Medicine (DICOM) compatible digital image storage with capability of storing uncompressed images on portable format without loss of image resolution;

2.4.1.9A ability to display and review prior relevant images during the procedure;

2.4.1.10A minimum detector diameter of 9 inches;

2.4.1.11A minimum spatial resolution of matrix of 1000 x 1000;
2.4.1.12A minimum contrast resolution to see the 1.5 mm hole in a standard phantom (see page 4, section 4.b. (low contrast performance) of Guidance Document Fluoro QA Guide posted on www.intersocietal.org/carotid/seeking/sample_documents.htm);

2.4.1.13A image monitor performance using the Society of Motion Picture and Television Engineers (SMPTE) pattern;

2.4.1.14A for equipment installed before 2006 that does not display cumulative dose and or dose area product (DAP), documentation of fluoroscopy time and the number of images per procedure is acceptable; and

2.4.1.15A radiation use must be consistent with the “as low as reasonably achievable” principle or ALARA radiation safety guidelines.

2.4.2A Emergency equipment and supplies (response cart or kit) including, but not limited to:

   2.4.2.1A oxygen/suction;
   2.4.2.2A defibrillator/automated external defibrillator (AED);
   2.4.2.3A external pacemaker equipment; and
   2.4.2.4A emergency drugs (including a master list with verification of expiration date).

2.4.3A Monitoring equipment must be available to perform:

   2.4.3.1A intravascular pressure;
   2.4.3.2A non-invasive blood pressure;
   2.4.3.3A pulse oximetry;
   2.4.3.4A electrocardiogram (ECG); and
   2.4.3.5A capnography (CO2) monitoring is recommended for use with moderate sedation.

2.4.4A Low or iso osmolar contrast must be used for intravascular injections.

2.4.5A Power or automated injectors (as applicable).

2.4.6A Adequate disposable supplies must be immediately available. These include U.S. Food and Drug Administration (FDA) approved catheters, wires, stents, balloons and embolic protection devices. Non-FDA approved devices may also be used as permitted by law.

2.4.7A Ancillary equipment as appropriate (e.g., monitoring equipment, blood coagulation testing equipment, workstations, picture archiving communication system (PACS), radiation protection for personnel (aprons and thyroid shields, portable shield either on wheels or suspended from ceiling).

STANDARD – Equipment and Instrumentation Quality Control

2.5A There must be a comprehensive Quality Improvement (QI) program to provide a standard of measurement for system performance and the documentation of any variance thereof.

   2.5.1A A QI Committee should be appointed to provide oversight to the equipment and instrumentation quality control (QC).
2.6A Fluoroscopic system QC testing must include a comprehensive evaluation of the system components, image performance and radiation output limits as outlined in the FDA Code of Federal Regulations (CFR) Title 21 subchapter J, Parts 1010 and 1020 and applicable FDA guidance documents.6

2.7A Image quality requirements, radiation output limits, and other fluoroscopic performance requirements must also comply with the health-code regulations of the state in which the facility is located.

2.8A The site-appointed qualified medical physicist must complete the performance evaluations at equipment installation and annually, unless state regulations require more frequent testing. Equipment performance evaluations are recommended semi-annually to include radiation output measurements, system quality control tests and image quality performance measurements.

2.9A The site-appointed medical physicist must perform a radiation safety survey to ensure that occupational workers and members of the public are shielded according to state regulation. This must be performed prior to installation of each new angiographic imaging system. A documented radiation safety survey of the interventional suite and adjacent areas that has been accepted by the State Radiation Program fulfills this requirement.

2.10A A radiation safety survey must be performed on all renovated or newly constructed interventional suites and adjacent areas. This must be performed prior to first patient use. This survey must confirm that the levels of radiation protection are in conformance with the State Radiation Program.

2.11A All spaces outside the procedure rooms should provide adequate protection for full time occupancy by non-radiation workers. This recommendation includes the control room.

2.12A Preventive maintenance (PM) service is recommended periodically per the manufacturers’ recommendations for each angiographic system at the facility.

2.13A Ancillary equipment (e.g., monitoring equipment, blood coagulation testing equipment, injectors, workstations, PACS, lead aprons, suction, oxygen lines, etc.) should also be included in a PM program.

2.14A The emergency response cart or kit must be checked at least monthly, with documentation to assure that all expected items are present, and the supplies are not expired.

2.15A There must be a process to check inventory of disposable supplies (e.g., catheters, wires, balloons, stents, embolic protection devices, contrast, portable oxygen tank) on a regular basis to assure that these supplies are readily available during a procedure.

STANDARD – Quality Control Documentation

2.16A All QC results must be documented and reviewed.

2.16.1A Documentation of the physicists’ evaluation, preventative maintenance and quality control tests performed, and service records for all angiographic systems and ancillary equipment must be maintained at the facility and available for review. The reports must be signed and dated by the person(s) performing the tests.

2.16.2A Results of all QC tests must be documented, archived and stored on film, in digital format, or on other suitable media according to state requirements (if applicable).
Section 3A: Administrative

STANDARD – Patient Confidentiality

3.1A All facility personnel must ascribe to professional principles of patient-physician confidentiality as legally required by federal, state, local or institutional policy or regulation.

STANDARD – Patient or Other Customer Complaints

3.2A There must be a policy in place outlining the process for patients or other customers to issue a complaint/grievance in reference to the care/services they received at the facility and how the facility handles complaints/grievances.

STANDARD – Primary Source Verification

3.3A There must be a policy in place identifying how the facility verifies the medical education, training, appropriate licenses and certifications of all physicians as well as, the certification and training of all technical staff members and any other direct patient care providers.

Sample documents are available for each of the required policies listed in Section 3A on the IAC Carotid Stenting website at www.intersocietal.org/carotid/seeking/sample_documents.htm.
Part B:
Process

Section 1B: Procedures and Protocols

STANDARD – Procedure Overview

1.1B Carotid stenting is a procedure in evolution and the procedure overview described below is not intended to be a comprehensive list of requirements to perform a case, nor does it list every step necessary for every patient. It represents an overview of the general steps to perform a typical elective case in order to provide a context for the overall requirements of this accreditation program. A facility may find it helpful to use this description to create an institutional template to be used as a reference when analyzing outcomes.

1.1.1B The facility must assure that appropriate staff members with BLS and ACLS certification are present during the procedure.

1.1.2B Appropriate staff must be available to assist the patient should an adverse event occur during the procedure and/or during recovery.

STANDARD – Procedure Requirements

1.2B Prior to performance of the procedure:

1.2.1B An adequate supply of devices approved by the FDA for marketing or investigational use must be available. This includes, but is not limited to: interventional guide wires, diagnostic catheters, guiding catheters, angioplasty balloons, distal protection devices and carotid stents.

1.2.2B Appropriate pharmacologic agents must be readily available for use during the procedure.

1.2.3B Proper identification of the patient and planned procedure (e.g., target vessel) must be carried out prior to puncture according to national patient safety goals and the proper patient name or identification (ID) must be present on the imaging system.

1.2.4B History and physical exam must be performed and should be in the chart and include documentation of relevant medications, allergies and bleeding disorders.

1.2.5B Independent neurologic assessment, National Institutes of Health Stroke Scale (NIHSS) and modified Rankin Scale (mRS) scores must be documented.

1.2.5.1B Patients undergoing carotid stenting will undergo independent neurologic assessments prior to and following the procedure to document the functional performance status and neurological examination and evaluate for any adverse neurological events.

1.2.6B Facility testing should be carried out including hemoglobin, hematocrit, platelet count, blood urea nitrogen (BUN), creatinine (within 30 days of the procedure) and pregnancy test (in women of childbearing age).

1.2.7B Imaging assessment of the cerebral parenchyma must be performed for patients with recent symptoms (stroke or cerebral transient ischemic attack [TIA]).
1.2.8B Imaging of the intra and extracranial carotid and vertebral circulation should be performed for both symptomatic and asymptomatic patients using either catheter or noninvasive angiography.

1.2.9B The current standard treatment for antiplatelet agents/antithrombotic therapy should have been administered. At this time, the current standard is a dual antiplatelet regimen consisting of aspirin and either clopidogrel or ticlopidine.

1.2.10B The facility must have a process to address intra-procedural complications, including development of a new neurological deficit during the procedure.

1.3B During the performance of the procedure:

1.3.1B Cardiac pacing supplies and equipment if necessary must be available.

1.3.2B Cardiovascular medications must be available.

1.3.3B Physiologic monitoring must include continuous ECG, blood pressure and pulse oximetry. Capnography may be used (if appropriate).

1.3.4B Intravenous access for administration of fluids and medications must be in place.

1.3.5B Radiation should be monitored during the procedure.

1.3.6B The stent delivery sheath or guiding catheter should be connected to an airless flush system.

1.3.7B Baseline ipsilateral carotid and cerebral angiography must be performed. The percent of carotid artery stenosis must be measured using NASCET criteria. Baseline ipsilateral carotid and cerebral angiography must be performed. The use of electronic calipers, (if available on the system) is required to quantitatively measure the percent of the stenosis to validate the clinical necessity for the procedure. Quantitative electronic measurements must use the post bulbar internal carotid artery diameter as the reference vessel, as per NASCET4.

1.3.8B Adequate anti-coagulation should be confirmed with activated clotting time (ACT) > 250 seconds prior to crossing the lesion.

1.3.9B Embolic protection device should be placed (if feasible).

1.3.10B Pre-dilation of lesion should be considered (if needed).

1.3.11B Deploy the stent.

1.3.12B Post-dilation should be considered (if needed).

1.3.13B Remove embolic protection following stent deployment.

1.3.14B Perform post stent deployment carotid and cerebral angiogram to check for stent patency, and to evaluate for distal emboli.

1.4B Following the performance of the procedure:

1.4.1B Perform and document post-procedure basic neurologic evaluation to assess for new neurologic deficits prior to moving the patient off the table.

1.4.1.1B The facility must have a protocol in place to address post-procedure neurologic deficits.

1.4.2B Assessment of blood pressure and the status of the puncture site.
1.4.2.1B Blood pressure must be controlled post-procedure according to the facility protocol.

1.4.3B A post-procedure note in the patient’s chart must be generated summarizing the procedure and addressing any immediate complications and the patient’s status at the end of the procedure.

1.4.3.1B Radiation usage as recorded by the angiographic system (i.e., fluoro time, DAP, mGy/cm) during the procedure must be documented in the final procedure report.

1.4.4B The patient must be moved to an appropriate setting such as a neuro critical care/ intensive care/step down unit with the equipment and trained personnel necessary to perform vascular, hemodynamic and neurological monitoring and stroke assessment.

1.4.5B Document post-procedure independent neurologic assessment (including a NIHSS and modified Rankin score) within approximately 24 hours and also within 60 days.

1.4.5.1B If there is any worsening from the pre procedure NIHSS (> 2 point change) or modified Rankin score or new neurological deficit, a neurologic consultation must be performed by a neurologist or if a neurologist is not available a physician with neurologic expertise.

1.4.6B Imaging of the stent must be performed within 60 days after the carotid artery stent procedure. Duplex ultrasound is the recommended imaging study.

STANDARD – Procedure Volumes

1.5B The procedure volume must be sufficient to maintain proficiency in procedure performance.

1.5.1B The facility must have specific privileging requirements for individual operators to perform both carotid/cerebral angiography and carotid stenting.

1.5.2B The facility must have performed at least 25 carotid stent procedures over the preceding three-year period for the facility to be considered eligible for accreditation.

1.5.2.1B If the facility has performed fewer than 25 carotid stent procedures over the preceding three-year period at least one operator must have performed 15 carotid stent cases (either in training or during post training experience as the primary operator) preferably with an embolic protection device in the past three years with adequate outcomes as defined in Quality Improvement (QI) Standards 2.4C and 2.5C.

1.5.3B The facility QI Program must include all elective cases.

1.5.3.1B Reporting of additional emergent cases which may involve additional intracranial or cervical advanced techniques should be given consideration.

Comment: The application review will recognize and take into consideration that a higher complication rate may be reported for the emergent cases.
Part C: Quality Improvement

Section 1C: Quality Improvement Program

STANDARD – QI Program

1.1C The facility must have a Quality Improvement (QI) Program and conduct internal quality assessment and improvement at regular intervals that are appropriate for the facility’s stated purpose and include carotid artery stenting (See Appendix). Typically, assessments are an ongoing process with monthly or quarterly review of results.

1.1.1C The performance of all medical, technical and ancillary staff must be assessed as part of the QI Program.

1.1.2C There must be a program in place to assess and evaluate patient and personnel radiation dose.

1.1.3C Adherence to National Patient Safety Goals must be documented.¹

1.1.4C The program must show evidence of improvement activities or, if an assessment confirms acceptable quality of a measure, the program must demonstrate improvement by selecting a new or an additional area for assessment.

1.1.5C The program should have pre-defined indicators of quality and pre-defined thresholds that indicate the need for corrective action. Comparisons with external benchmarks are desirable.

1.1.6C Participation in a national registry of carotid stent procedures is strongly recommended.

1.1.7C The program should include a review of procedural indications, safety and complications as well as standardized and recognized clinical outcome measures.

STANDARD – QI Documentation

1.2C QI documentation (policies, reports, records, etc.) must be maintained at the facility and made available to all personnel.

1.2.1C The Medical Director and appropriate staff must review and maintain minutes or reports of QI evaluations and document (as applicable) corrective measures taken.

1.2.2C The facility must have a mechanism in place to track each carotid stent procedure performed including but not limited to patient identification, date of birth, date of procedure, the clinical indication, pre- and post-procedure independent neurologic assessment including NIHSS and mRS, use of embolic protection device use, degree of stenosis improvement post-procedure and any procedure complications that occur within 60 days after the procedure.

1.2.3C Documentation of a written process to contact all patients for the follow-up neurological assessment within 60 days after the procedure.
Section 2C: Quality Improvement Measures

STANDARD – QI Measures

2.1C For purposes of Quality Improvement (QI) and reporting of outcomes, the carotid stent procedure begins when the guide catheter or sheath for stent placement has been inserted into the patient. Deployment is considered successful if the stent has been placed across the target lesion per the instructions for use.

2.1.1C The QI Program must include clinical indications including risk category and outcome measures. All of the quality components as outlined below in items 2.6C, 2.7C, 2.8C and 2.9C should also be included (as appropriate).

2.1.2C Outcome data is most widely available for treatment of carotid bifurcation disease secondary either to atherosclerosis, post endarterectomy restenosis or radiation induced stenosis. Therefore, the outcome benchmarks used in this program are intended to be applied only to cases treated for these indications. However, all cases should be included in outcome monitoring.

2.1.3C Outcomes data must be consistent with national benchmarks and where there are no benchmarks the data must be used to internally improve processes and procedures. For facilities not meeting the benchmarks described below in items 2.4C and 2.5C, a plan for improvement must be submitted and documented improvement provided within 12-18 months before full accreditation will be considered.

2.1.4C A process to confirm the accuracy of the percentage of stenosis reported for symptomatic and asymptomatic patients warranting the intervention must be in place.

2.1.4.1C If quantitative electronic measurements are used to determine the stenosis percentage, there must be a process to assess differences between the subjectively reported and the electronic measurements. Ideally, the quantitative (electronic) and subjective measures of stenosis severity should be very similar or identical.

2.1.4.2C If subjective measurements are used to determine the need for treatment, deviations between the subjective and electronic measurement warrant a documented explanation and where appropriate, documented corrective action.

2.1.4.3C 90% of patients should meet facility defined clinical and degree of stenosis indications as defined in the QI program.

STANDARD – Reporting of Outcomes

2.2C Initial Accreditation

2.2.1C Many facilities may have low volumes of procedures when they apply for initial accreditation. This low volume makes statistical analysis of complication rates unreliable and may penalize a low volume facility that has a complication early in its experience. In addition, a new facility may not have performed any procedures (although at least one individual physician must have a minimum experience of 15 procedures in order to apply for accreditation). Similar to the technique used to evaluate the outcomes of vascular surgeons performing carotid endarterectomy® facilities applying for initial accreditation will be evaluated on a minimum of 100 patients (50 symptomatic and 50 asymptomatic) in the previous three years. If the facility has not performed 100 procedures in the three years prior to initial accreditation, the facility will be provided up to 50 complication-free hypothetical procedures in each indication category. For example, a facility with a low volume will be given up to 100 hypothetical procedures (50 symptomatic and 50 asymptomatic) without a death or stroke.
2.3C **Reaccreditation**

2.3.1C After initial accreditation the facility will be evaluated only on cases performed since the initial accreditation. Each facility will be provided one time with 50 complication-free hypothetical procedures in each indication category. The facility will then create a running total of 50 procedures in each indication category. The outcomes must be tracked both by actual procedures as well as actual procedures plus hypothetical procedures. The overall combined analysis is what will be used to determine if a facility meets the benchmarks. However, complications and complication rates in actual procedures must be reviewed to prevent the hiding of an unacceptably high complication rate within the moving 50. For each actual procedure performed a hypothetical procedure will be dropped from the total until all hypothetical procedures are removed and outcomes are then based solely on actual procedures performed by the facility.

2.3.1.1C After the facility has performed 50 procedures in an indication category, analysis of outcomes for the facility for that category will be based on the total actual procedures performed since the initial accreditation, and the ability to use any hypothetical cases will be removed.

2.3.1.2C After a facility has performed 100 actual procedures in an indication category it will be evaluated on the running total of the most recent 100 procedures in that category.

2.4C **Outcome Measures: Benchmarks by Indications for Procedures**

2.4.1C Elective Symptomatic Carotid Stenosis – defined as experiencing acute TIA symptoms or completed ischemic stroke within six months of the intervention, but neurologically stable for at least 24 hours.

2.4.1.1C Benchmark: < 6% all stroke and death within 30 days of the procedure. Adverse events noted on the first neurological assessment post procedure (21-60 days) must be counted as a complication. (See Appendix for definition of stroke)

Comment: In order to apply for accreditation, if the facility exceeds the benchmark of 6% but does not exceed 10%, a plan for improvement must be submitted and documented improvement provided within 12-18 months before full accreditation will be considered. Benchmarks must be met within three years.

2.4.2C Elective Asymptomatic Carotid Stenosis

2.4.2.1C Benchmark: < 3% all stroke and death within 30 days of the procedure. Adverse events noted on the first neurological assessment post procedure (21-60 days) must be counted as a complication. (See Appendix for definition of stroke)

Comment: In order to apply for accreditation, if the facility exceeds the benchmark of 3% but does not exceed 4%, a plan for improvement must be submitted and documented improvement provided within 12 months before full accreditation will be considered. Benchmarks must be met within three years.

2.5C **Outcome Measures: Technical**

2.5.1C **Peri/Immediate Post-Procedure**

2.5.1.1C Successful Stent Deployment – Deployment is considered successful if the stent has been deployed across the target lesion per the instructions for use.

i. Benchmark = > 95% of the cases have successful stent deployment.
2.5.1.2C Improvement in degree of stenosis – post procedural % angiographic stenosis by North American Symptomatic Carotid Endarterectomy Trial (NASCET) methodology.4

i. Benchmark = > 95% of the cases have improvement in the degree of stenosis.

2.5.1.3C Measurement of neurological outcome by an independent examiner (defined as the person performing the assessment that is not the operator or any other person listed on the procedure documentation).

i. Modified Rankin Scale at 24 hours (or discharge)
ii. NIHSS at 24 hours
iii. Neurological Consult – If there is any worsening from the pre procedure NIHSS (> 2 point change) or mRS or new neurological deficit, a neurologic consultation must be performed by a neurologist or if not available a physician with neurologic expertise.
   • Benchmark = 100% at 24 hours for pre and post-procedure measurement of NIHSS and mRS.

2.5.1.4C Post-procedure follow-up must be performed within 60 days.

i. 80% of patients will have 60-day outcomes follow-up data.
   • Imaging of the stent must be performed within 60 days for patency. Duplex ultrasound is the recommended imaging study.
   Comment: If stent occlusions occur in two or more patients at post-procedure follow-up (i.e., within 60 days) within a three-year review cycle then a process must be in place for review and reporting of causative factors and methods for improvement.

ii. Independent neurological assessment: Measurement of neurological outcome within 60 days.
   • Modified Rankin scale
   • NIHSS
   • Neurological Consult: If there is any worsening from the pre procedure NIHSS (> 2 point change) or Rankin or new neurological deficit, a neurologist or if a neurologist is not available a physician with neurologic expertise.

2.6C Administrative Quality Assessment

2.6.1C The QI Program must be in place to assess and improve the administrative quality of the facility’s operation. Administrative areas that may be assessed include, but are not limited to:

2.6.1.1C scheduling back logs;
2.6.1.2C patient wait times;
2.6.1.3C accuracy of patient information during scheduling;
2.6.1.4C completeness of documentation;
2.6.1.5C late reports;
2.6.1.6C time from completion of procedure to distribution of final report;
2.6.1.7C patient satisfaction and feedback;
2.6.1.8C referring physician satisfaction and feedback; and
2.6.1.9C patient education: on individual risk factors, smoking cessation, signs and symptoms of stroke and calling 911, importance of follow-up after discharge, review of discharge medications including importance of adherence to antithrombotic therapy.

2.7C Technical Quality Assessment

2.7.1C The QI Program must include assessment of the technical quality of the images and procedures being performed. Areas that may be assessed include but are not limited to:

2.7.1.1C image quality;
2.7.1.2C image display/labeling; and
2.7.1.3C documentation of adverse technical events such as equipment or device failure.

2.8C Physician Performance Quality Assessment

2.8.1C The QI Program must include assessment of the performance of physicians regarding the quality of medical practice (such as report accuracy, appropriateness of care, effectiveness of performing the procedure) and physician behaviors (communication and professionalism). Areas that may be assessed include but are not limited to:

2.8.1.1C peer review;
2.8.1.2C correlation of interpretation with other diagnostic studies, pathology/surgical results and/or patient outcomes;
2.8.1.3C time from completion of procedure to distribution of final report;
2.8.1.4C referring physician satisfaction and feedback; and
2.8.1.5C patient satisfaction and feedback.

2.9C Patient Radiation Dose Review/Evaluation

2.9.1C The QI Program must include an assessment of patient dose as compared to published guidelines. At a minimum the program must meet the criteria outlined in Guidelines for Patient Radiation Dose Management.13

Comment: The radiation dose thresholds outlined in the reference are trigger values set at a prudently low value such that the possibility of an injury at the threshold level is highly unlikely. There is no implication that exceeding a threshold will always cause an injury.

2.9.1.1C When radiation dose thresholds are exceeded, the facility must have a process in place for patient monitoring and follow-up.

2.9.1.2C All steps in the evaluation and follow-up process must be documented.

2.9.2C Other Diagnostic Angiographic Procedures – Facility should have processes in place for monitoring cerebral diagnostic arteriography with complications not to exceed published guidelines.10
Section 3C: Quality Improvement Meetings

STANDARD – QI Meetings

3.1C Quality Improvement (QI) meetings must be documented.

3.1.1C All relevant personnel assessed in the QI Program must participate in periodic facility meetings to review findings and determine actions for improvement of performance. At a minimum, these meetings must occur at least every six months.

3.1.2C Every stroke and death must be reviewed during these meetings.

3.1.3C All relevant personnel must be included in periodic facility meetings to provide in-service education containing relevant topics. Topics should include safety procedures, technical information and improvements to be made based on quality assessments and other information.
Selected Bibliography


9. National Stroke Association (NSA) [www.stroke.org/we-can-help/healthcare-professionals/improve-your-skills/tools-training-and-resources/training/nih?pagename=nihs;](www.stroke.org/we-can-help/healthcare-professionals/improve-your-skills/tools-training-and-resources/training/nih?pagename=nihs; American Stroke Association (ASA) [professional.heart.org](professional.heart.org)


Appendix

QUALITY IMPROVEMENT MEASURES: INDICATIONS AND ADVERSE EVENTS

I) INDICATIONS FOR CAROTID ANGIOPLASTY AND/OR STENTING

Definition: Symptomatic carotid stenosis is defined as a carotid stenosis associated with an ipsilateral cerebral or retinal TIA or infarction within the past six months.

Comment: Centers for Medicare and Medicaid Services (CMS) reimbursement policy requires patients to meet specific indications (www.cms.gov/medicare-coverage-database/details/ncd-details.aspx?NCDId=201). Currently, CMS policy requires that all patients are high surgical risk (see below) and have a pre-procedure mRS < 2. For symptomatic patients current CMS reimbursement policy requires > 70% angiographic diameter stenosis using NASCET criteria. For patients participating in an approved trial, the stenosis may be > 50%. For asymptomatic patients, current CMS reimbursement policy requires patients to be enrolled in an approved trial and have > 80% angiographic diameter stenosis using NASCET criteria.

High Risk Category as defined by CMS:
1. congestive heart failure (CHF) class III/IV;
2. left ventricular ejection fraction (LVEF) < 30%;
3. unstable angina;
4. contralateral carotid occlusion;
5. recent myocardial infarction (MI);
6. previous CEA with recurrent stenosis;
7. prior radiation treatment to the neck; and
8. other conditions that were used to determine patients at high risk for CEA in the prior carotid artery stenting trials and studies, such as ARCHER, CABERNET, SAPPHIRE, BEACH and MAVERIC II.

IAC Carotid Stenting recognizes that not all patients fall into one of the above categories and CMS may alter reimbursement policy as more data are available. Other potential indications are listed below. However, the presence of one or more of these potential anatomic and medical conditions in a patient does not imply that carotid angioplasty and/or stenting is indicated for that patient. Appropriate therapy is determined by symptoms, severity of stenosis and an overall assessment of the risks and benefits of the procedure compared to alternative medical and interventional therapies.

1. Stenosis that is surgically difficult to access (e.g., high bifurcation requiring mandibular dislocation).
2. Stenosis in a patient with significant medical disease that would make the patient high risk for surgery
3. Stenosis and one of the following conditions:
   a. significant tandem lesion that may require endovascular therapy;
   b. radiation-induced stenosis;
   c. restenosis following CEA;
   d. refusal to undergo CEA following proper informed consent;
   e. stenosis secondary to arterial dissection;
   f. stenosis secondary to fibromuscular dysplasia (stents rarely, if ever, indicated);
   g. stenosis secondary to vasculitis (stents rarely, if ever, indicated).
4. Stenosis associated with contralateral carotid artery occlusion requiring treatment before undergoing cardiac surgery.
5. Severe underlying carotid artery stenosis revealed after recanalization of carotid occlusion following thrombolysis for acute stroke (presumed to be the etiology of the treated occlusion) or to enable thrombolysis for acute stroke.
6. Pseudoaneurysm.
II) ADVERSE EVENTS

1. All stroke: For purposes of IAC Carotid Stenting Quality Improvement (QI) and comparing facility outcomes to national benchmarks, a stroke is defined as an ischemic or hemorrhagic brain injury causing a neurologic deficit that persists for more than 24 hours.
   a. cerebral infarction/ ischemic stroke;
   b. intracranial hemorrhage/ hemorrhagic stroke;
   c. unknown type of stroke (no imaging performed).
2. TIA (symptoms < 24 hours) with or without neuroimaging evidence of acute infarction is not considered a stroke for IAC Carotid Stenting QI and comparison to national benchmarks, although it may be considered a stroke for research reporting purposes.
3. Other neurologic (edema/hyperperfusion syndrome).
4. All death.
5. MI.
6. Other cardiac event:
   a. heart failure or pulmonary edema;
   b. arrhythmia requiring cardioversion, pacemaker insertion, or ICD insertion;
   c. hypotension requiring parenteral medications for > 24 hours post-procedure.
7. Renal failure with new requirement for dialysis.
8. Infection related to procedure requiring antibiotics.
9. Angiographic complications:
   a. dissection requiring treatment;
   b. urgent surgery required for technical problems with stent deployment or placement;
   c. intracranial embolization.
11. Arterial access:
    a. pseudoaneurysm requiring treatment with thrombin injection and/or compression during hospitalization;
    b. access site injury requiring open surgical repair.
12. Vessel thrombosis, peripheral embolization, or ischemia of an extremity.
13. Other:
    a. unexpected intubation or resuscitation;
    b. contrast reaction.