Non-QPP Measure Descriptions

Measure Title:
M2S 1: Procedures with statin and antiplatelet agents prescribed at discharge

Measure Description:
This measure estimates the frequency of procedures where a statin and antiplatelet agent was prescribed at discharge for any of the following procedures: Infra-Inginal Bypass, Supra-Inginal Bypass, Peripheral Vascular Intervention, Carotid Artery Stent, Carotid Endarterectomy, Thoracic and Complex EVAR, Endovascular AAA Repair, and Open AAA Repair.

Denominator:
All procedures captured in the Vascular Quality Initiative® Infra-Inginal Bypass, Supra-Inginal Bypass, Peripheral Vascular Intervention, Carotid Artery Stent, Carotid Endarterectomy, Thoracic and Complex EVAR, Endovascular AAA Repair, and Open AAA Repair Registries where the patient was at least 18 years old at the time of the procedure, and the patient was discharged alive during the reporting period.

Denominator Exceptions: N/A

Denominator Exclusions:
Exclude procedures associated with patients who were not discharged on statins or antiplatelets due to a documented medical reason.

Numerator:
All procedures captured in the Vascular Quality Initiative® Infra-Inginal Bypass, Supra-Inginal Bypass, Peripheral Vascular Intervention, Carotid Artery Stent, Carotid Endarterectomy, Thoracic and Complex EVAR, Endovascular AAA Repair, and Open AAA Repair Registries where the patient was at least 18 years old at the time of the procedure, and the patient was discharged alive during the reporting period where a statin and antiplatelet agent were prescribed at discharge.

Rationale:
Many patients undergoing vascular surgical procedures are not on appropriate medical therapy. Statin and antiplatelet therapy preoperatively and at discharge are associated with reduced mortality and improved survival after vascular surgery.

Supporting Evidence:

Medical management (MM) with antiplatelet (AP) and statin therapy is recommended for most patients undergoing vascular surgery. Importantly, discharge on AP and statin therapy was
associated with improved 5-year survival, compared with discharge on neither medication. Discharge on a single medication was associated with intermediate survival at 5 years.


Statin and antiplatelet therapy preoperatively and at discharge was associated with reduced 30-day mortality and an absolute 18% improved 5-year survival after vascular surgery. However, one-third of patients are sub optimally managed in real world practice. This demonstrates an opportunity for quality improvement that can substantially improve survival after vascular surgery.

Measure Type: Process

Number of Multiple Performance Rates (If Applicable): N/A

Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Effective Clinical Care

NQF Number: N/A
**Measure Title:**
M2S 2: Amputation-free survival assessed at least 9 months following Infra-Inguinal Bypass for intermittent claudication

**Measure Description:**
This measure estimates freedom from amputation or death at least 9 months after Infra-Inguinal Bypass for intermittent claudication.

**Denominator:**
All procedures captured in the Vascular Quality Initiative® Infra-Inguinal Bypass Registry associated with patients who were at least 18 years old at the time of the procedure and with indication of claudication for the leg undergoing treatment during the calendar year prior to the reporting interval with at least 9 month follow up.

**Denominator Exceptions:** N/A

**Denominator Exclusions:**
Procedures associated with patients lost to follow up are excluded from this measure. Procedures associated with missing key data elements that are required to calculate the outcomes measure will be excluded from this measure.

**Numerator:**
All procedures captured in the Vascular Quality Initiative® Infra-Inguinal Bypass Registry associated with patients who were at least 18 years old at the time of the procedure and with indication of claudication for the leg undergoing treatment during the calendar year prior to the reporting interval with at least 9 month follow up and who were alive and without minor or major amputation reported at a follow-up visit at least 9 months after the treatment.

**Rationale:**
Patients with symptomatic peripheral arterial disease (PAD) secondary to atherosclerosis may present with intermittent claudication or critical limb ischemia. The first line of treatment for intermittent claudication due to PAD should be supervised walking exercise program to increase collateral circulation; however due to a uncertainty of the effect of this treatment revascularization, by an endovascular or surgical approach has often been performed to improve quality of life and limb salvage. The proposed measure will therefore serve as an indicator of both appropriateness and overall outcome.

**Supporting Evidence:**

**Measure Type:** Outcome

**Number of Multiple Performance Rates (If Applicable):** N/A

**Inverse Measurement:** No

**Proportion Measure Scoring:** Yes

**Continuous Measure Scoring:** No

**NQS Domain:** Patient Safety

**NQF Number:** N/A
Measure Title:
M2S 3: Infra-inguinal bypass for claudication patency assessed at least 9 months following surgery

Measure Description:
This measure estimates the frequency of graft patency assessment one year Infra-Inguinal Bypass.

Denominator:
All Infra-inguinal bypass procedures performed for claudication in Vascular Quality Initiative® Infrainguinal Bypass Registry during the calendar year prior to the reporting interval associated with patients who were at least 18 years old at the time of the procedure.

Denominator Exceptions: N/A

Denominator Exclusions:
Procedures associated with patients lost to follow up are excluded from this measure.

Numerator:
All Infra-inguinal bypass procedures performed for claudication in Vascular Quality Initiative® Infrainguinal Bypass Registry during the calendar year prior to the reporting interval associated with patients who were at least 18 years old at the time of the procedure and with at least 9 month follow up where graft patency was assessed at follow-up.

Rationale:
Patients with symptomatic peripheral arterial disease (PAD) secondary to atherosclerosis may present with intermittent claudication or critical limb ischemia. The first line of treatment for intermittent claudication due to PAD should be supervised walking exercise program to increase collateral circulation; however due to a uncertainty of the effect of this treatment revascularization, by an endovascular or surgical approach has often been performed to improve quality of life and limb salvage. The proposed measure will therefore serve as an indicator of both appropriateness and overall outcome.

Supporting Evidence:

Graft patency rates vary widely among series. One study in which graft patency was assessed by using MRA disclosed an 84% limb salvage rate and a 78% primary graft patency rate at 21 months of follow-up. Prosthetic grafts carry a primary 3-year patency rate of 39% and a 3-year secondary patency rate of 59%, with a 25% risk of amputation within 3 years.


Infrainguinal reconstructive surgery is associated with a mortality rate of somewhat less than 5% despite the high-risk nature of these patients.


Measure Type: Process

Number of Multiple Performance Rates (If Applicable): N/A

Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Communication and Care Coordination

NQF Number: N/A
Measure Title:  
M2S 4: Amputation-free survival assessed at least 9 months following Supra-Inguinal Bypass for claudication

Measure Description:  
This measure estimates freedom from amputation or death at least 9 months after Supra-Inguinal Bypass for claudication.

Denominator:  
All procedures captured in the Vascular Quality Initiative® Supra-Inguinal Bypass Registry associated with patients who were at least 18 years old at the time of the procedure and with indication of claudication in at least one leg during the calendar year prior to the reporting interval with at least 9 month follow up.

Denominator Exceptions: N/A

Denominator Exclusions:  
Procedures associated with patients lost to follow up are excluded from this measure. Procedures associated with missing key data elements that are required to calculate the outcomes measure will be excluded from this measure.

Numerator:  
All procedures captured in the Vascular Quality Initiative® Supra-Inguinal Bypass Registry associated with patients who were at least 18 years old at the time of the procedure and with indication of claudication in at least one leg during the calendar year prior to the reporting interval with at least 9 month follow up who were alive and without minor or major amputation reported after the treatment.

Rationale:  
Patients with symptomatic peripheral arterial disease (PAD) secondary to atherosclerosis may present with intermittent claudication or critical limb ischemia. The first line of treatment for intermittent claudication due to PAD should be supervised walking exercise program to increase collateral circulation; however due to a uncertainty of the effect of this treatment revascularization, by an endovascular or surgical approach has often been performed to improve quality of life and limb salvage. The proposed measure will therefore serve as an indicator of both appropriateness and overall outcome.

Supporting Evidence:  


Measure Type: Outcome
Number of Multiple Performance Rates (If Applicable): N/A

Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Patient Safety

NQF Number: N/A
Measure Title:
M2S 5: Amputation-free survival at assessed at least 9 months following Peripheral Vascular Intervention for intermittent claudication

Measure Description:
This measure estimates freedom from amputation or death at least 9 months after Peripheral Vascular Intervention for intermittent claudication.

Denominator:
All procedures captured in the Vascular Quality Initiative® Peripheral Vascular Intervention Registry associated with patients who were at least 18 years old at the time of the procedure and with indication of claudication for the treated leg(s) undergoing Peripheral Vascular Intervention below external iliac during the calendar year prior to the reporting interval with at least at least 9 month follow up.

Denominator Exceptions: N/A

Denominator Exclusions:
Procedures associated with patients lost to follow up are excluded from this measure. Procedures associated with missing key data elements that are required to calculate the outcomes measure will be excluded from this measure.

Numerator:
All procedures captured in the Vascular Quality Initiative® Peripheral Vascular Intervention Registry associated with patients who were at least 18 years old at the time of the procedure and with indication of claudication for the treated leg(s) undergoing Peripheral Vascular Intervention below external iliac during the calendar year prior to the reporting interval with at least at least 9 month follow up associated with patients who were alive and without minor or major amputation reported after the intervention.

Rationale:
Patients with symptomatic peripheral arterial disease (PAD) secondary to atherosclerosis may present with intermittent claudication or critical limb ischemia which is manifested as ischemic rest pain or tissue loss (lower extremity ulceration or gangrene). Treatment consists of medical optimization of risk factors and revascularization. Revascularization may be accomplished by open surgery or peripheral vascular intervention (PVI). PVI includes angioplasty, stenting/stent-grafting and atherectomy techniques. PVI is increasingly used in the treatment of symptomatic PAD due to its lower morbidity and mortality compared to open surgery. Interventions may be performed on a variety of anatomical levels broad divided into suprainguinal (aorta and iliac arteries) or infrainguinal (femoral, popliteal and infrapopliteal arteries including the tibial and peroneal arteries).

Supporting Evidence:


Measure Type: Outcome

Number of Multiple Performance Rates (If Applicable): N/A

Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Patient Safety

NQF Number: N/A
Measure Title:
M2S 6: Peripheral Vascular Intervention patency assessed at least 9 months following infrainguinal PVI for claudication

Measure Description:
This measure estimates the frequency of patency assessment at least 9 months following infrainguinal Peripheral Vascular Intervention for claudication.

Denominator:
All procedures captured in the Vascular Quality Initiative® Peripheral Vascular Intervention Registry for claudication distal to the external iliac during the calendar year prior to the reporting interval. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Denominator Exceptions: N/A

Denominator Exclusions:
Procedures associated with patients lost to follow up are excluded from this measure.

Numerator:
All procedures captured in the Vascular Quality Initiative® Peripheral Vascular Intervention Registry for claudication distal to the external iliac during the calendar year prior to the reporting interval with at least 9 month follow up where patency was assessed at follow up. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Rationale:
Patients with symptomatic peripheral arterial disease (PAD) secondary to atherosclerosis may present with intermittent claudication or critical limb ischemia which is manifested as ischemic rest pain or tissue loss (lower extremity ulceration or gangrene). Treatment consists of medical optimization of risk factors and revascularization. Revascularization may be accomplished by open surgery or peripheral vascular intervention (PVI). PVI includes angioplasty, stenting/stent-grafting and atherectomy techniques. PVI is increasingly used in the treatment of symptomatic PAD due to its lower morbidity and mortality compared to open surgery. Interventions may be performed on a variety of anatomical levels broad divided into suprainguinal (aorta and iliac arteries) or infrainguinal (femoral, popliteal and infrapopliteal arteries including the tibial and peroneal arteries).

Supporting Evidence:


Measure Type: Process

Number of Multiple Performance Rates (If Applicable): N/A

Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Communication and Care Coordination

NQF Number: N/A
Measure Title:  
M2S 7: Ipsilateral stroke-free survival at assessed at least 9 months following Carotid Artery Stenting for asymptomatic procedures

Measure Description:  
This measure estimates freedom from stroke or death at least 9 months after carotid artery stenting for asymptomatic carotid stenosis.

Denominator:  
All procedures captured in the Vascular Quality Initiative® Carotid Artery Stenting Registry for asymptomatic carotid stenosis during the calendar year prior to the reporting interval with at least 9 month follow up. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Denominator Exceptions: N/A

Denominator Exclusions:  
Procedures associated with patients lost to follow up are excluded from this measure. Procedures associated with missing key data elements that are required to calculate the outcomes measure will be excluded from this measure.

Numerator:  
All procedures captured in the Vascular Quality Initiative® Carotid Artery Stenting Registry for asymptomatic carotid stenosis during the calendar year prior to the reporting interval with at least 9 month follow up which are free of either stroke or death (any cause) after index carotid artery stenting. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Rationale:  
Surgeons performing CAS on asymptomatic patients must select patients at low risk for morbidity or stroke and perform the procedure with a very low complication rate in order to achieve benefit. The proposed measure will therefore serve as an indicator of both appropriateness and overall outcome.

Supporting Evidence:  
Neurologically asymptomatic patients with ≥ 60% diameter stenosis should be considered for CAS for reduction of long-term risk of stroke, provided the patient has a 3- to 5-year life expectancy and perioperative stroke/death rates can be ≤ 3%

Periprocedural and long-term follow-up outcomes (including death and stroke) of CAS appear similar for ACS and SCS.
For asymptomatic carotid stenosis, CAS is associated with a substantially higher risk of postoperative stroke or in-hospital death than CEA even after adjustment for baseline differences in hospital and patient characteristics.

Among individuals undergoing carotid artery revascularization from a large sample of US hospitals, CAS was associated with higher risk of perioperative mortality, stroke, and unfavorable discharges compared with CEA for all ages and clinical presentations.

Measure Type: Outcome

Number of Multiple Performance Rates (If Applicable): N/A

Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Effective Clinical Care

NQF Number: N/A
Measure Title:
M2S 8: Ipsilateral stroke-free survival assessed at least 9 months following isolated Carotid Endarterectomy for asymptomatic procedures

Measure Description:
This measure estimates freedom from stroke or death at least 9 months after carotid endarterectomy for asymptomatic carotid artery stenosis.

Denominator:
All procedures captured in the Vascular Quality Initiative® Carotid Endarterectomy Registry for carotid endarterectomy for asymptomatic disease during the calendar year prior to the reporting interval with at least 9 month follow up. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Denominator Exceptions: N/A

Denominator Exclusions:
Procedures associated with patients lost to follow up are excluded from this measure. This measure also excludes combined Coronary Artery Bypass Graft procedures, and excludes combined proximal or distal intervention or other combined arterial procedures. Procedures associated with missing key data elements that are required to calculate the outcomes measure will be excluded from this measure.

Numerator:
All procedures captured in the Vascular Quality Initiative® Carotid Endarterectomy Registry for carotid endarterectomy for asymptomatic disease during the calendar year prior to the reporting interval with at least 9 month follow up which are free of either stroke or death (any cause) after index carotid endarterectomy. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Rationale:
Asymptomatic carotid bifurcation stenosis poses a potentially significant public health problem. In the US, stroke is the leading cause of disability and the 3rd leading cause of death. This report seeks to measure the freedom from stroke or death at one year for carotid endarterectomy, as prevention of stroke and overall survival are important metrics in surgical treatment of carotid disease.

Supporting Evidence:
This measure is supported by large body of evidence including several randomized clinical trials (e.g. ACAS), systematic reviews, and clinical practice guidelines.


**Measure Type:** Outcome

**Number of Multiple Performance Rates (If Applicable):** N/A

**Inverse Measurement:** No

**Proportion Measure Scoring:** Yes

**Continuous Measure Scoring:** No

**NQS Domain:** Effective Clinical Care

**NQF Number:** N/A
Measure Title:
M2S 9: Imaging-based maximum aortic diameter assessed at least 9 months following Thoracic and Complex EVAR procedures

Measure Description:
This measure estimates the frequency of imaging-based maximum aortic diameter assessment at least 9 months after Thoracic and Complex EVAR procedures.

Denominator:
All Thoracic and Complex EVAR procedures captured in Vascular Quality Initiative® Thoracic and Complex EVAR Registry during the calendar year prior to the reporting interval. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Denominator Exceptions: N/A

Denominator Exclusions:
Procedures associated with patients lost to follow up are excluded from this measure. This measure also excludes conversion to open.

Numerator:
All Thoracic and Complex EVAR procedures captured in Vascular Quality Initiative® Thoracic and Complex EVAR Registry during the calendar year prior to the reporting interval with at least 9 month follow up, where the follow up included a CT scan or MR documenting the maximum aortic diameter. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Rationale:
Endovascular repair of thoracic and complex aneurysms is a viable alternative to open repair; however, endoleak and sac expansion are not uncommon. Imaging to monitor sac diameter remains critical to follow the long-term efficacy of this type of repair. The proposed measure will therefore serve as an indicator of optimal follow-up of these patients after this procedure.

Supporting Evidence:

Blair R, Collins A, Harkin DW. Complex EVAR for abdominal aorto-iliac aneurysm (AAIA) is associated with high rate of endoleak and less aortic sac shrinkage compared to conventional EVAR for AAA. Ir J Med Sci. 2014 Oct 17.


Measure Type: Process
Number of Multiple Performance Rates (If Applicable): N/A

Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Communication and Care Coordination

NQF Number: N/A
Measure Title:
M2S 10: Survival at least 9 months after elective repair of small thoracic aortic aneurysms

Measure Description:
This measure estimates freedom from death at least 9 months after elective repair of small thoracic aortic aneurysms in patients without a known connective tissue disorder.

Denominator:
All elective procedures for aneurysm repair captured in the Vascular Quality Initiative® Thoracic and Complex EVAR Registry during the calendar year prior to the reporting interval where maximum aortic diameter is less than or equal to 6 cm for women, or less than or equal to 6.5 cm for men at the index procedure, and with at least at least 9 month follow up. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Denominator Exceptions: N/A

Denominator Exclusions:
Procedures associated with patients lost to follow up that are not dead are excluded from this measure. Procedures associated with patients with a known connective tissue disorder are excluded from this measure. Procedures associated with missing key data elements that are required to calculate the outcomes measure will be excluded from this measure.

Numerator:
All elective procedures for aneurysm repair captured in the Vascular Quality Initiative® Thoracic and Complex EVAR Registry during the calendar year prior to the reporting interval where maximum aortic diameter is less than or equal to 6 cm for women, or less than or equal to 6.5 cm for men at the index procedure, and with at least 9 month follow up and who are alive. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Rationale:
Surgeons should select patients for intervention who have a reasonable life expectancy and who do not have a high surgical risk. One year survival after repair of small thoracic aortic aneurysms is an indicator of patients who were not frail prior to the procedure and who did not experience a major complication. The proposed measure will therefore serve as an indicator of both appropriateness and overall outcome.

Supporting Evidence:

Measure Type: Outcome

Number of Multiple Performance Rates (If Applicable): N/A

Inverse Measurement: No
Proportion Measure Scoring: Yes
Continuous Measure Scoring: No
NQS Domain: Effective Clinical Care
NQF Number: N/A
Measure Title:
M2S 11: Imaging-based maximum aortic diameter assessed at least 9 months following Endovascular AAA Repair procedures

Measure Description:
This measure estimates the frequency of imaging-based maximum aortic diameter assessment at least 9 months after Endovascular AAA Repair procedures.

Denominator:
All Endovascular AAA Repair procedures captured in Vascular Quality Initiative® Endovascular AAA Repair Registry during the calendar year prior to the reporting interval. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Denominator Exceptions: N/A

Denominator Exclusions:
Procedures associated with patients lost to follow up are excluded from this measure. This measure also excludes procedures that were converted to open.

Numerator:
All Endovascular AAA Repair procedures captured in Vascular Quality Initiative® Endovascular AAA Repair Registry during the calendar year prior to the reporting where maximum aortic diameter was assessed during a follow up visit at least 9 months after surgery and who are alive. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Rationale:
Change in maximum aortic diameter is an indicator of disease progression. Multiple guidelines recommend annual lifelong imaging follow-up after EVAR to identify and to correct complications such as endoleaks or residual aortic sac enlargement and thereby prevent death due to aneurysm rupture after EVAR. The proposed process measure will serve as an indicator for appropriate clinical follow-up.

Supporting Evidence:


Measure Type: Process

Number of Multiple Performance Rates (If Applicable): N/A
Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Communication and Care Coordination

NQF Number: N/A
Measure Title:
M2S 12: Survival at least 9 months after elective repair Endovascular AAA Repair of small abdominal aortic aneurysms

Measure Description:
This measure estimates freedom from death at least 9 months after elective repair of small abdominal aortic aneurysms.

Denominator:
All elective procedures for aneurysm repair captured in the Vascular Quality Initiative® Endovascular AAA Repair Registry during the calendar year prior to the reporting interval where maximum aortic diameter is less than or equal to 5.5 cm for women, or less than or equal to 6.0 cm for men at the index procedure, and with at least 9 month follow up. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Denominator Exceptions: N/A

Denominator Exclusions:
Procedures associated with patients lost to follow up are excluded from this measure. Procedures associated with missing key data elements that are required to calculate the outcomes measure will be excluded from this measure.

Numerator:
All elective procedures for aneurysm repair captured in the Vascular Quality Initiative® Endovascular AAA Repair Registry during the calendar year prior to the reporting interval where maximum aortic diameter is less than or equal to 5.5 cm for women, or less than or equal to 6.0 cm for men at the index procedure, and with at least 9 month follow up and who are alive. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Rationale:
Surgeons should select patients for intervention who have a reasonable life expectancy and who do not have a high surgical risk. One year survival after endovascular repair of small abdominal aortic aneurysms is an indicator of patients who were not frail prior to the procedure and who did not experience a major complication. The proposed measure will therefore serve as an indicator of both appropriateness and overall outcome.

Supporting Evidence:
Elective repair is recommended for patients that present with a fusiform AAA ≥5.5 cm in maximum diameter, in the absence of significant co-morbidities.

Measure Type: Outcome

Number of Multiple Performance Rates (If Applicable): N/A
Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Effective Clinical Care

NQF Number: N/A
Measure Title:  
M2S 13: Survival at least 9 months after elective Open AAA repair of small abdominal aortic aneurysms

Measure Description:  
This measure estimates freedom from death at least 9 months after elective open repair of small abdominal aortic aneurysms.

Denominator:  
All elective procedures for aneurysm repair captured in the Vascular Quality Initiative® Open AAA Repair Registry during the calendar year prior to the reporting interval where maximum aortic diameter is less than or equal to 5.5 cm for women, or less than or equal to 6.0 cm for men at the index procedure, and with at least 9 month follow up. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Denominator Exceptions: N/A

Denominator Exclusions:  
Procedures associated with patients lost to follow up are excluded from this measure. Procedures associated with missing key data elements that are required to calculate the outcomes measure will be excluded from this measure.

Numerator:  
All elective procedures for aneurysm repair captured in the Vascular Quality Initiative® Open AAA Repair Registry during the calendar year prior to the reporting interval where maximum aortic diameter is less than or equal to 5.5 cm for women, or less than or equal to 6.0 cm for men at the index procedure, and with at least 9 month follow up who are alive. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Rationale:  
Elective abdominal aortic aneurysm repair is a procedure designed to improve survival by preventing aneurysm rupture. The analysis of rupture risk vs operative risk and the life expectancy of the patient should be considered when choosing a surgical treatment. Operative mortality after surgical repair of small AAA is not significantly less than for larger aneurysms but patients with multiple risk factors such as Age ≥70 years, CHF, COPD, HTN and CRI, should have much larger aneurysms before open repair in order to justify the risk of operation. The proposed measure will therefore serve as an indicator of both appropriateness and overall outcome.

Supporting Evidence:  
Elective repair is recommended for patients that present with a fusiform AAA ≥5.5 cm in maximum diameter, in the absence of significant co-morbidities.
VQI QCDR Non-QPP Measures


**Measure Type:** Outcome

**Number of Multiple Performance Rates (If Applicable):** N/A

**Inverse Measurement:** No

**Proportion Measure Scoring:** Yes

**Continuous Measure Scoring:** No

**NQS Domain:** Effective Clinical Care

**NQF Number:** N/A
Measure Title:
M2S 14: Disease specific patient-reported outcome surveys for Varicose Vein procedures

Measure Description:
This measure estimates the frequency of disease specific patient reported outcomes assessments for varicose vein procedures during the reporting year.

Denominator:
All procedures captured in the Vascular Quality Initiative® Varicose Vein Registry during the reporting year. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Denominator Exceptions: N/A

Denominator Exclusions:
Exclude procedures where the patient died prior to discharge. Procedures associated with patients lost to follow up are excluded from this measure.

Numerator:
All procedures captured in the Vascular Quality Initiative® Varicose Vein Registry during the reporting year that had patient reported outcomes recorded both prior to treatment and post-treatment during the reporting year. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

Rationale:
Patient Reported Outcomes, such as the Chronic Venous Insufficiency Quality of Life Questionnaire (CIVIQ) the VEINES-QOL/Sym, and the ATTRACT QOL, enquiring about lower limb symptoms, lifestyle factors and satisfaction with disease treatments help determine clinical outcomes, patient satisfaction and retreatment rates. The proposed measure will therefore serve as an indicator of both appropriateness and overall outcome.

Supporting Evidence:
From the National Quality Forum’s 2013 report, Patient Reported Outcomes (PROs) in Performance Measurement:

Patient engagement is increasingly acknowledged as a key component of a comprehensive strategy, (along with performance improvement and accountability), to achieve a high quality, affordable health system. Emerging evidence affirms that patients who are engaged in their care tend to experience better outcomes and choose less costly but effective interventions.

Historically, with the exception of collecting feedback on satisfaction or experience with care, patients remain an untapped resource in assessing the quality of healthcare and of long-term support services. Patients are a valuable and, arguably, the authoritative source of information on outcomes beyond experience with care. These include health-related quality of life, functional status, symptom and symptom burden, and health behaviors.
Patient Reported Outcome Measures (PROMs) are standardized instruments that capture patients’ self-assessment of their health and can provide timely information on patient health status, function and symptoms over time that can be used to improve patient-centered care and inform clinical decision-making.


**Measure Type:** Process

**Number of Multiple Performance Rates (If Applicable):** N/A

**Inverse Measurement:** No

**Proportion Measure Scoring:** Yes

**Continuous Measure Scoring:** No

**NQS Domain:** Person and Caregiver-Centered Experience Outcomes

**NQF Number:** N/A
**Measure Title:**
M2S 15: Appropriate management of retrievable IVC filters

**Measure Description:**
This measure estimates the frequency of IVC Filter procedures that used a retrievable filter, and were managed appropriately. Appropriate management is described as one of the following: (1) the IVC Filter was removed during the reporting year, or (2) the IVC filter was attempted to be removed during the reporting year, or (3) the patient was evaluated within three months of filter placement, and removal was not clinically indicated.

**Denominator:**
All retrievable IVC filter procedures captured in the Vascular Quality Initiative® Inferior Vena Cava Filter Registry during the reporting year. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

**Denominator Exceptions:** N/A

**Denominator Exclusions:**
Procedures associated with patients who could not be contacted for follow-up, or who refused filter removal at evaluation, will be excluded from the measure.

**Numerator:**
All retrievable IVC filter procedures captured in the Vascular Quality Initiative® Inferior Vena Cava Filter Registry during the reporting year associated with one of the following: (1) the IVC Filter was removed during the reporting year, (2) the IVC filter was attempted to be removed during the reporting year, or (3) the patient was evaluated within three months of filter placement, and removal was not clinically indicated. All procedures are associated with patients who were at least 18 years old at the time of the procedure.

**Rationale:**
The proposed measure will serve as an indicator of appropriate care for patients who have received retrievable IVC filters in the reporting period. The use of IVC filters has increased dramatically over the past 15 years. Published retrieval rates vary widely.

**Supporting Evidence:**


Sutphin PD, Reis SP, McKune A, Ravanzo M, Kalva SP, Pillai AK. Improving Inferior Vena Cava Filter Retrieval Rates with the Define, Measure, Analyze, Improve, Control Methodology. J Vasc Interv Radiol. 2015

**Measure Type:** Process

**Number of Multiple Performance Rates (If Applicable):** N/A

**Inverse Measurement:** No

**Proportion Measure Scoring:** Yes

**Continuous Measure Scoring:** No

**NQS Domain:** Effective Clinical Care

**NQF Number:** N/A
Measure Title:
M2S 16: Absence of unplanned reoperation after major lower extremity amputation

Measure Description:
The percentage of patients undergoing closed lower extremity major amputation (below, thru or above knee) for ischemic arterial disease who do NOT have unplanned reoperation during the same admission for infection, bleeding or amputation revision.

Denominator:
All procedures captured in the Vascular Quality Initiative® Lower Extremity Amputation Registry where the patient was at least 18 years old at the time of the procedure and underwent closed lower extremity major amputation (below, thru or above knee) for ischemic arterial disease during the reporting period.

Denominator Exceptions: N/A

Denominator Exclusions:
To both numerator and denominator: Patients undergoing initial open amputation (who always require reoperation) and patients undergoing amputation for non-arterial etiologies, including trauma, neoplasm, infection and diabetic neuropathy without underlying arterial disease.

Numerator:
All procedures captured in the Vascular Quality Initiative® Lower Extremity Amputation Registry where the patient was at least 18 years old at the time of the procedure and underwent closed lower extremity major amputation (below, thru or above knee) for ischemic arterial disease who did NOT have unplanned reoperation during the same admission for infection, bleeding or amputation revision during the reporting period.

Rationale:
Major closed amputations (below and above knee) can be complicated by infection, bleeding or failure to heal, resulting in unplanned reoperation, if the level and conduct of the initial procedure is not appropriate. This outcome measure appropriately identifies proper amputation level selection and proper surgical technique.

Measure Type: Outcome

Number of Multiple Performance Rates (If Applicable): N/A

Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Patient safety

NQF Number: N/A
Measure Title:
M2S 17: Absence of serious technical complications during peripheral arterial intervention

Measure Description:
The percentage of patients undergoing lower extremity interventional treatment for arterial occlusive disease (including balloon angioplasty, stenting and atherectomy) who do NOT experience technical complications (Access site: stenosis, occlusion, AV fistula, hematoma or pseudoaneurism, Thrombosis, Embolization, Perforation, or Dissection (target lesion or other artery) that require admission, or surgical or interventional treatment.

Denominator:
All procedures captured in the Vascular Quality Initiative® Peripheral Vascular Intervention Registry where the patient was at least 18 years old at the time of the procedure and underwent peripheral arterial intervention (balloon angioplasty, stenting or atherectomy) of the lower extremity (abdominal aorta and more distal) arteries for treatment of occlusive disease during the reporting period.

Denominator Exceptions: N/A

Denominator Exclusions:
To both numerator and denominator: Patients undergoing interventional treatment for aneurysm disease or trauma.

Numerator:
All procedures captured in the Vascular Quality Initiative® Peripheral Vascular Intervention Registry where the patient was at least 18 years old at the time of the procedure and underwent peripheral arterial intervention (balloon angioplasty, stenting or atherectomy) of the lower extremity (abdominal aorta and more distal) arteries for treatment of occlusive disease who did NOT experience technical complications (Access site: stenosis, occlusion, AV fistula, hematoma or pseudoaneurism, Thrombosis, Embolization, Perforation, or Dissection (target lesion or other artery) that required hospital admission, or transfusion, surgical or interventional treatment prior to discharge during the reporting period.

Rationale:
Avoiding technical complications is critical to patient safety during peripheral interventional procedures. The complications listed above can result in serious harm, so this outcome measure tracks an important safety issue.

Measure Type: Outcome

Number of Multiple Performance Rates (If Applicable): N/A

Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Patient safety

NQF Number: N/A
Measure Title: M2S-18: Venous clinical severity score (VCSS) assessment before varicose vein treatment

Measure Description: The percentage of patients undergoing surgical or ablation treatment (radiofrequency, laser, mechanochemical, chemical, or embolic adhesive) of truncal, perforator or cluster veins who have venous clinical severity score (VCSS) of the treated leg(s) assessed prior to treatment.

Denominator: All procedures captured in the Vascular Quality Initiative® Varicose Vein Registry where the patient was at least 18 years old at the time of the procedure and underwent surgical or ablation treatment (radiofrequency, laser, mechanochemical, chemical, or embolic adhesive) of truncal, perforator or cluster veins during the reporting period.

Denominator Exceptions: N/A

Denominator Exclusions: None

Numerator: All procedures captured in the Vascular Quality Initiative® Varicose Vein Registry where the patient was at least 18 years old at the time of the procedure and underwent surgical or ablation treatment (radiofrequency, laser, mechanochemical, chemical, or embolic adhesive) of truncal, perforator or cluster veins who had the venous clinical severity score (VCSS) of their treated leg(s) assessed prior to treatment during the reporting period.

Rationale: Pre-treatment assessment using VCSS, which measures 10 clinical signs and symptoms is recommended in practice guidelines to ensure that treatment is not being recommended for patients with minimal venous disease and to allow post-treatment assessment of improvement.

Measure Type: Process

Number of Multiple Performance Rates (If Applicable): N/A

Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Effective clinical care

NQF Number: N/A
Measure Title: M2S 19: Proper patient selection for perforator vein ablation

Measure Description: The percentage of patients undergoing treatment of incompetent lower extremity perforator veins with surgical or ablation treatment (radiofrequency, laser, mechanochemical, chemical, or embolic adhesive) who have CEAP clinical severity of disease C5 or C6 (venous ulcer or history of ulcer), and not less severe C1-4 disease.

Denominator: All procedures captured in the Vascular Quality Initiative® Varicose Vein Registry where the patient was at least 18 years old at the time of the procedure and underwent treatment of incompetent lower extremity perforator veins with surgical or ablation treatment (radiofrequency, laser, mechanochemical, chemical, or embolic adhesive) during the reporting period.

Denominator Exceptions: N/A

Denominator Exclusions: None

Numerator: All procedures captured in the Vascular Quality Initiative® Varicose Vein Registry where the patient was at least 18 years old at the time of the procedure and underwent treatment of incompetent lower extremity perforator veins with surgical or ablation treatment (radiofrequency, laser, mechanochemical, chemical, or embolic adhesive) who had CEAP clinical severity of disease C5 or C6 (venous ulcer or history of ulcer) during the reporting period.

Rationale: Perforator vein treatment has only been shown to benefit patients with venous ulcers or a history of ulcers, but is often applied to patients with less severe disease unnecessarily. This measure documents appropriate patient selection and effective cost reduction.

Measure Type: Process

Number of Multiple Performance Rates (If Applicable): N/A

Inverse Measurement: No

Proportion Measure Scoring: Yes

Continuous Measure Scoring: No

NQS Domain: Effective Clinical Care

NQF Number: N/A