

# Vascular Surgery Data Dictionary

Blue Cross Blue Shield of Michigan Vascular Surgery Registry Data Collection Definitions

Definitions updated 2.17.2021

Tuble of Contents	Tabl	le of	Conten	ts
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Physician	8
Fellow ID/Second Operator	8
Procedure Date & Start Time	8
Procedure Date & End Time	8
Status of Procedure	8
Staged Procedure	9
Consultations	9
Imaging Studies (w/in past 6 months)	9
Labs - Pre Procedure	10
Pre Procedure Creatinine	10
Pre Procedure Hemoglobin (Hgb)	11
Pre Procedure BNP	11
Pre Procedure Troponin	11
Pre Procedure COVID-19	12
Labs - Post Procedure	13
Post Procedure Peak Creatinine	13
Post Procedure Nadir Hemoglobin	13
Post Procedure COVID-19	14
Labs - Other	15
Albumin	15
Indications	15
Indication Type	15
Indications for EVAR/AAA Revascularization	15
Asymptomatic	15
Abdominal/Back Pain	15
Rapidly Increasing Aneurysm Diameter	16
Unfit for open AAA repair	16
Unfit for general anesthesia	16
Infection	16
Size of Iliac Aneurysm	16
Correction of Endoleak	16
Concomitant Iliac Occlusive Disease	17
Lower Extremity Ischemia/Emboli	17
Documented Patient Anxiety Levels	17
Penetrating Ulcer	17
Complication from Prior Procedure	17
Trauma	17
Mycotic Aneurysm	18
Pre-procedure Smoking Cessation	18
Indications for Lower Extremity Revascularization (Open Bypass / Open Thrombectomy)	18
Claudication	18
https://bmc2.ora/print/book/export/html/78	2/61

Rest Pain	
Threatened Bypass Graft	19
Acute Limb Ischemia	19
Failed Endovascular Procedure	19
Infection (Lower Extremity Revascularization)	19
Facilitation of Procedure	
Pre-Procedure Exercise Therapy	
Impaired Ability to Work	
Peripheral Aneurysm Repair	
Increased Stent Velocity	
Increased Stent Graft Velocity	
Wound (WIfI)	21
Ischemia (WIfI)	21
Foot Infection (WIfI)	22
Complication from Prior Procedure	
Trauma	
Pre-procedure Smoking Cessation	23
Indications for Upper Extremity Revascularization (Open Bypass / Open Thrombectomy)	23
Ulcer/Gangrene	23
Acute Limb Ischemia	23
Angina/Abnormal Cardiac Stress Test	24
BP discrepancy	24
Arm Claudication	24
Peripheral Aneurysm Repair	24
Complication from Prior Procedure	24
Trauma	25
Pre-procedure Smoking Cessation	25
Procedure Details	25
Open Bypass/Revascularization	
Graft Insertion	
Redo Procedure	
Vein Graft	27
Vein Graft Harvest	27
Number of Vein Segments	27
Minimal Vein Graft Diameter	
Prosthetic Graft	
Vein Cuff	
Intra Operative Graft Patency	
Intra Operative Graft Revision	
Closure for Open Exposure	
Open Abdominal Aortic Aneurysm (OAAA)	
Prior Aortic Surgery (OAAA)	

2/16/2021 https://bmc2.org/print/book/export/html/78	20
Aneurysm anatomy (OAAA)	
Lowest Pre-Intubation Blood Pressure (for ruptured AAA repairs only) (OAAA)	
Mental Status (for ruptured AAA repairs only) (OAAA)	
Cardiac Arrest (for ruptured AAA repairs only) (OAAA)	
Timetrame: Symptoms to Incision (for ruptured AAA repairs only) (OAAA)	
I imetrame: Admission to Incision (for ruptured AAA repairs only) (OAAA)	
Abdomen Explored (OAAA)	
Conversion from Endovascular Repair (OAAA)	
Exposure (OAAA)	
Distal Anastomosis (OAAA)	
Graft Body Diameter (OAAA)	
Graft Type (OAAA)	
Renal Status (OAAA)	
Anastomotic Felt Reinforcement (OAAA)	
Hypogastric ligated/occluded (OAAA)	
Proximal Clamp Position (OAAA)	
Inferior Mesenteric Artery at Completion (OAAA)	
Renal/Visceral Ischemic Time (OAAA)	
Intra Operative Graft Revision (OAAA)	
Cold Renal Perfusion (OAAA)	
Mannitol administered during procedure (OAAA)	
Closure for Open Exposure (OAAA)	
Endovascular Aneurysm Repair (EVAR)	
Prior Aortic Surgery (EVAR)	
Maximum AAA Diameter (EVAR)	
Iliac Aneurysm (EVAR)	
Aneurysm location (EVAR)	
Aneurysm anatomy (EVAR)	
Contained rupture (EVAR)	
Infrarenal Neck Diameter (EVAR)	
Infrarenal Neck Length (EVAR)	
Ruptured AAA (EVAR)	
Lowest Pre-Intubation Blood Pressure (for ruptured AAA repairs only) (EVAR)	
Mental Status (for ruptured AAA repairs only) (EVAR)	
Cardiac Arrest (for ruptured AAA repairs only) (EVAR)	
Timeframe: Symptoms to Incision (for ruptured AAA repairs only) (EVAR)	
Timeframe: Admission to Incision (for ruptured AAA repairs only) (EVAR)	

2/16/2021 Abdomon Explored (EV(AP)	https://bmc2.org/print/book/export/html/78	22
Abdomen Explored (EVAR)	·······	ວວ ວວ
Graft Rody Diameter (EVAR)		)) 22
Bight Distal Soal Zono Diameter	······	21
L off Distal Seal Zone Diameter	······	21 21
Graft Type (EV/AP)	······	21 21
Graft Configuration	······	25
Additional graft components (EV/AR)	······	35
Renal Status	~	,, 35
Hypogastric Coiled/Plugged	~	36
Hypogastric Intentionally Covered	~	36
Hypogastric Unintentionally Covered	~	36
Arterial Injury		36
Intra-Operative Revision Needed		70 75
Endoleak at Completion	~	יי דג
	······	יי דג
Additional Planned Procedures	······	יי דג
	······································	יר גע גע
Open Thrombostomy (OT)	······	20
Timeframe: Presentation to incision (OT)	······································	20
Prior lytic procedure (OT)	······	20
Vessel location (OT)	~	20
	~	20
Completion angio (OT)	~	20
Concomitant endarterectomy (OT)	~	20
Data Fields for All Procedure Types	~	20
Anesthesia Type	~	20 20
Antibiotics Pre Procedure	/	10 10
Skin Preparation		10 10
Contraindicated to Chlorbevidine & Alcoh	ol Skin Preparation	10 10
Glucose (peak)		10
Nadir body temperature	2	41
Crystalloids	2	 41
Estimated Blood Loss (EBL)	2	 41
ASA (American Society of Anesthesiologi	sts) Class	 41
Contrast Types	2	 42
Total IV Contrast Used	2	42
Heparin Administered		
Peak Intra Procedure Activated Clotting T	ime (ACT)	- 43
End of procedure ACT		- 43
Locations		43
Vessel Location		43

Lesion Segment Area	
PVI Procedure Performed	44
Bypass Graft	44
Graft Type	
Graft Origin	
Graft Insertion	45
Lesion Length	45
Heavy Calcium	45
In-stent Restenosis	45
Thrombus	45
Pre Stenosis % (0-100)	
Post Stenosis % (0-100)	
Final Balloon Diameter	
Stents	
Stent Name	
Stent Diameter	
Stent Length	
Vascular Access	
Vascular Access Site(s)	
Vascular Access Type	
Vessel Accessed	
Access Guidance	
Access Approach	
Sheath Size	
Sheath Removed	
Vascular Closure Type	
Sheath Removal Time	
Outcomes During Procedure	
Death/Cause (ODP)	
Dissection (Not Repaired)	
Myocardial Infarction (MI) (ODP)	
Cardiac Arrest	
Embolus (ODP)	
Thrombus (ODP)	
Stent/Graft Thrombosis	51
Vessel Perforation	51
TIA/Stroke (ODP)	51
Transfusion (ODP)	51
Vascular Access Complications (ODP)	
Vascular Surgery Emergent (ODP)	
Amputation (in Lab/OR)	
Compartment Syndrome (in Lab/OR)	53

2/16/2021	
Outcomes	Pos

0/2021	https://bindz.org/print/book/export/ntint/76
utcomes Post Procedure	
Death/Cause (OPP)	
Stay in ICU	
Vasopressors Post Operatively	
Respiratory	
Myocardial Injury	
Dysrhythmia	
Congestive Heart Failure (CHF)	
TIA/Stroke (OPP)	
Infection/Sepsis	
New Requirement for Dialysis	
Transfusion (OPP)	
Symptomatic Prior to Transfusion	
Vascular Access Complications (OPP)	
Compartment Syndrome	
Embolus (OPP)	
Thrombus (OPP)	
Stent/Graft Thrombosis	
Amputation	
Return to Operating Room (OPP)	
Bowel Ischemia	
Was the LOS >2 days after EVAR? (OPP).	

## **Procedure Information for Vascular Intervention**

### Physician

### **Data Abstraction Instructions:**

Choose the attending physician from the drop down list, or create a physician identification if not already listed.

#### Selections:

· Choose physician

### Supporting Definitions:

If physician is not available in the drop down, enter the physician's information to create a physician ID.

Required:

Yes

### Fellow ID/Second Operator

#### Data Abstraction Instructions:

Use the drop down box to choose a physician. If not available in drop down, enter the fellow's or second operator's information to create a physician ID.

#### Selections:

• Enter Fellow ID/Second Operator

### Supporting Definitions:

This is an optional field if your site chooses to track it's fellows or second operators.

Required: No

### **Procedure Date & Start Time**

### **Data Abstraction Instructions:**

Enter the date of the current procedure (mm/dd/yyyy) and enter the time the procedure was initiated. (Military time)

#### Selections:

· Enter Date & Time

### Supporting Definitions:

The time the procedure started is defined as the time which local anesthetic was first administered for vascular access, or the time of the first attempt at vascular access for the peripheral intervention (use whichever is earlier), or incision time for open vascular surgery procedures. Indicate the time (hours:minutes) using military 24-hour clock, beginning at midnight (0000 hours). If an arterial sheath is already in place, use the time of the introduction of a catheter or the time the sheath was exchanged.

### **Required:**

Yes

### **Procedure Date & End Time**

#### **Data Abstraction Instructions:**

Enter the date and time the procedure ends.

#### Selections:

Enter Date & Time

#### Supporting Definitions:

The time the procedure ended is defined as the time the primary operator leaves the room for peripheral interventions. End time for open surgical procedures is defined as the time when all instrument and sponge counts are completed; all dressings and drains are secured; and the physicians/surgeons have completed all procedure-related activities on the patient. Should the patient expire in the procedure area, indicate the time the patient was pronoun ced. Indicate the time (hours:minutes) using military 24-hour clock, beginning at midnight (0000 hours).

Required: Yes

### **Status of Procedure**

#### Data Abstraction Instructions:

Indicate status of the procedure using the following categories.

- · Elective: The procedure could be deferred without increased risk of compromised vascular outcome. This should include the elective or scheduled patients.
- Urgent: Required operation within 72 hours, but > 12 hours of symptoms.
- Emergent: Required operation within 12 hours of symptoms.

#### Required: Yes

### **Staged Procedure**

### **Data Abstraction Instructions:**

Indicate if the intervention is the part of a staged procedure.

### Selections:

- Yes
- No

### Supporting Definitions:

Interventions planned for subsequent procedures at the time of the initial procedure are considered staged procedure, e.g., initial procedure left superficial femoral angioplasty (SFA) with a plan for right SFA in future. Patients undergoing lysis procedures are considered staged procedures. A plan for a staged procedure can be developed with the patient as an outpatient or an inpatient. Enter a new case for any staged procedure (For the first case, mark No for staged procedure, for the second procedure mark Yes).

The following are NOT staged procedures: a subsequent intervention due to restenosis, a diagnostic angiogram with intervention planned on a separate day, a coronary procedure followed by a peripheral procedure, a return attempt after a failed PVI.

### **Required:**

Yes

### Consultations

### Data Abstraction Instructions:

Indicate whether the patient had any of the following physician consultations within the last 6 months up to the date of procedure. These consults are done in anticipation of a pending procedure.

### Selections:

- Cardiology Consultation
  - Yes
    - ∘ No
- Pulmonology Consultation
  - Yes
  - ∘ No
- Primary Care/Internal Medicine
  - ∘ Yes
  - ◇ No
- Hematology Consultation
   Yes
  - No
- Renal Consultation
  - ◊ Yes
  - No

### Supporting Definitions:

#### Choose all that apply.

If a consultation is ordered <u>before</u> the procedure, but the patient was seen after the procedure, you can still claim it as a pre-procedure consultation. Post procedure consults do not qualify.

For discharges with multiple procedures, consults should only be applied to procedures which they immediately precede (they do not apply to all procedures within the hospitalization). For example, a cardiac consult followed by two procedures: procedure # 1consult = yes, procedure #2 consult = no.

#### Required: Yes

### Imaging Studies (w/in past 6 months)

### **Data Abstraction Instructions:**

For each of the listed studies, indicate if the study was performed to diagnose PAD, CAD, or as part of a functional assessment within prior 6 months. Also, for each of the listed studies that were performed, indicate if the study was normal or abnormal or include value where applicable. If study was not performed, record the study was not done.

For patients who have multiple procedures during the same hospitalization, do not use the same pre-procedure imaging studies as used prior to the first procedure. Only enter imaging studies performed prior to the specific procedure being entered.

Enter all available data for ABIs, TBIs, and toe pressures that are valid for the present procedure (include both right and left, regardless of the operative side). For Chest X-Ray enter the documentation of the physical condition of the lungs. For example, if the radiologist documents that an ET tube is in place and ther are no structural abnormalities noted in the lungs. Enter Normal.

Selections:

- ABI's Compressible (Compressible </= 1.4)</li>
   Yes
  - No
  - Right Pre Procedure ABI • Yes, enter value • No
- Left Pre Procedure ABI
   Yes, enter value
- ∘ No
- Right Pre Procedure TBI
   Yes, enter value
  - o res, en o No
- Left Pre Procedure TBI
- Yes, enter value
  - ∘ No
- Right Pre Procedure Toe Pressure
   ves, enter value (mm Hg)
   No
- Left Pre Procedure Toe Pressure
  - Yes, enter value (mm Hg)
  - No
- Vein Mapping:
  - Yes
  - No
- Duplex Ultrasound:
  - Yes
  - ∘ No
    - If yes, select option
      - Normal
      - Abnormal
- Computerized Tomographic Angiography (CTA):
  - ◊ Yes
  - No
    - If yes, select option

      Normal
      - Abnormal
- Magnetic Resonance Imaging/Magnetic Resonance Angiography (MRI/MRA):
  - o Yes ◇ No
    - NO
    - If yes, select option

      Normal
      - Abnormal
- Contrast Cineangiography:
  - Yes
  - No
    - If yes, select option
    - Normal
  - Abnormal
- Cardiac Stress Test:
  - Yes
  - ∘ No
    - If yes, select option
    - NormalAbnormal
  - Electrocardiogram:
- Electrocardiogram:
   Yes
  - No
    - If yes, select option
      - Normal
      - Abnormal
- Chest X-Ray:
  - Yes
  - No
    - If yes, select option
      - Normal
    - Abnormal

Required: Yes

### Labs - Pre Procedure

### Pre Procedure Creatinine

**Data Abstraction Instructions:** 

Use the last value between 30 days prior to arrival and current procedure. If there is no value, mark not drawn.

### Selections:

- Yes
- Enter valueNot drawn

### Supporting Definitions:

When multiple lab values are available pre-procedure, enter the value closest to the procedure start time. If a patient has multiple procedures within a hospitalization, the pre-procedure labs should be abstracted after the previous procedure; if no labs are drawn from the end of the previous procedure to the start of the current procedure, enter "Not drawn".

Required: Yes Suffix: mg/dl Minimum: 0.1 Maximum: 15 Soft Minimum: 0.3

### Pre Procedure Hemoglobin (Hgb)

### **Data Abstraction Instructions:**

Use the last value between 30 days prior to arrival and current procedure. If there is no value, mark "Not drawn."

#### Selections:

- Yes
- Enter value
   Not drawn
- Not ar

### Supporting Definitions:

When multiple lab values are available pre-procedure, enter the value closest to the procedure start time. If a patient has multiple procedures within a hospitalization, the pre-procedure labs should be abstracted after the previous procedure; if no labs are drawn from the end of the previous procedure to the start of the current procedure, enter "Not drawn".

#### **Required:**

Yes Suffix: g/dl Minimum: 2 Maximum: 20 Soft Minimum: 5 Soft Maximum: 18

### **Pre Procedure BNP**

#### **Data Abstraction Instructions:**

Indicate if a BNP value was obtained within the 30 days prior to the procedure. If more than one value exists, use the one closest to the procedure start time.

### Selections:

Pre Procedure BNP

- Yes
  - o Value\_\_\_\_pg/mL

### No

### Supporting Definitions:

When multiple lab values are available pre-procedure, enter the value closest to the procedure start time. If a patient has multiple procedures within a hospitalization, the pre-procedure labs should be abstracted after the previous procedure; if no labs are drawn from the end of the previous procedure to the start of the current procedure, enter "Not drawn".

Required: Yes Suffix: pg/mL

### Pre Procedure Troponin

**Data Abstraction Instructions:** 

#### https://bmc2.org/print/book/export/html/78

Indicate if a Troponin I, Troponin T, Troponin I HS (High Sensitivity) or Troponin T HS value was obtained within the 30 days prior to the procedure. If more than one value exists, use the value closest to the procedure start time.

#### Selections:

Pre Procedure Troponin

- Yes
  - Pre procedure troponin I
    - Yes
      - Enter lab value \_\_\_\_\_
        Pick unit of lab value from I
        - Pick unit of lab value from list
          - ng/dLng/mL
          - ng/IIIL ■ ng/L
          - pg/mL
          - pg/m
  - NoPre procedure troponin T
    - Yes
      - Enter lab value \_\_\_\_\_
        Pick unit of lab value from list
        - k unit of lai
        - ng/dL
        - ng/mLng/L
        - ng/L ■ pg/mL
        - pg/i
  - Pre procedure troponin I HS
    - Yes

No

- Enter lab value \_\_\_\_\_
  - Pick unit of lab value from list
    - ng/dL
    - ng/mL
    - ng/Lpg/mL
- NoPre procedure troponin T HS
  - Yes
    - Enter lab value
    - Pick unit of lab value from list
      - ng/dL
      - ng/mL
      - ng/L
      - pg/mL
- No

### Not Drawn

### Supporting Definitions:

When multiple lab values are available pre-procedure, enter the value closest to the procedure start time. If a patient has multiple procedures within a hospitalization, the pre-procedure labs should be abstracted after the previous procedure; if no labs are drawn from the end of the previous procedure to the start of the current procedure, enter "Not Drawn" for the initial Troponin field.

Required: Yes Suffix: ng/dL, ng/mL, ng/L, pg/mL

### **Pre Procedure COVID-19**

### **Data Abstraction Instructions:**

#### Pre Procedure COVID-19

Indicate the COVID-19 status known at procedure start. If there are multiple results available within 7 days of the procedure, please enter the COVID-19 result that is closest to the procedure start time.

**Positive COVID-19**<sup>\*</sup> = within 7 days of procedure start, patient has tested positive for COVID-19 documented by clinician and/or laboratory specimen result (lower respiratory, upper respiratory, serology, sputum, stool, urine, oropharyngeal or nasopharyngeal swab). Enter Positive if a patient tested positive prior to 7 days of the procedure start and does not meet the criteria for recovered.

If Positive COVID-19 result, enter the date and time of Positive Result Posted. If a date is documented but the time is not documented enter the date and then enter 00:00 for the time. If date and time are not documented, enter Not Documented.

**Negative COVID-19**\* = within 7 days of procedure start, patient has tested negative for COVID-19 documented by clinician and/or laboratory specimen result (lower respiratory, upper respiratory, serology, sputum, stool, urine, oropharyngeal or nasopharyngeal swab).

### If Negative COVID-19 result, enter the date and time of Negative Result Posted. If a date is documented but the time is not documented enter the date and then enter 00:00 for the time. If date and time are not documented, enter Not Documented.

Investigating COVID-19 = possibility patient is infected is being investigated (specimen for testing may or may not have been obtained yet)

COVID-19 Not Suspected\*\* = there is no documentation that the patient is suspected to have COVID-19 and they have not been tested.

Recovered COVID-19 = The patient was previously diagnosed with COVID-19 infection (lab or clinical criteria) and is no longer contagious as defined by:

Test-based strategy:

At least three (3) days (72 hours) have passed since recovery defined as resolution of fever and improvement of respiratory symptoms, AND Two (2) consecutive negative COVID-19 laboratory tests >= 24 hours apart

OR

Symptom-based strategy:

- At least three (3) days (72 hours) have passed since recovery defined as resolution of fever and improvement of respiratory symptoms, AND
- At least ten (10) days have passed since symptom onset

#### OR

- If tested positive for COVID-19 and never exhibited symptoms:
- At least 10 days have passed since first positive COVID-19 test, OR
- Two (2) consecutive negative COVID-19 laboratory tests >= 24 hours apart

#### Timeframe: Prior to procedure start.

\*Please include any of the viruses associated with the COVID-19 pandemic (2019-nCOV, MERS-CoV, SARS-CoV, SARS-CoV2, and other associated viruses identified by the CDC).

\*\*If procedure performed prior to the pandemic, please select COVID Not Suspected.

#### Selections:

- Positive COVID-19 (Within 7 Days of Procedure Start)
  - Documented
     Enter Date/Time: MM/DD/YYYY HH:MM
  - Not Documented
- · Negative COVID-19 (Within 7 Days of Procedure Start)
  - Documented
    - Enter Date/Time: MM/DD/YYYY HH:MM
- Not Documented
- Investigating COVID-19
- COVID-19 Not Suspected
- Recovered COVID-19

#### Supporting Definitions:

World Health Organization. (2020, March 2). WHO.int. Laboratory testing for coronavirus disease 2019 (COVID-19) in suspected human cases. Interim guidance.

https://www.who.int/emergencies/diseases/novel-coronavirus-2019/technical-guidance/laborfleratory-guidance/

Centers for Disease Control and Prevention. (2020, May 2). CDC.gov. https://www.cdc.gov/coronavirus/2019-ncov/hcp/disposition-hospitalized-patients.html

Centers for Disease Control and Prevention. (2020, May 3). CDC.gov. https://www.cdc.gov/coronavirus/2019-ncov/hcp/disposition-in-home-patients.html

### Required:

Yes

### Labs - Post Procedure

### **Post Procedure Peak Creatinine**

### **Data Abstraction Instructions:**

Record the highest level recorded from the end of the procedure to the next procedure or discharge, whichever occurs first. If there is no value drawn post procedure, mark "Not drawn." For extended hospitalizations, greater than 30 days, use the highest creatinine prior to day 30 after the procedure.

### Selections:

- Yes
- Enter value
- Not drawn

### Supporting Definitions:

The next procedure is any procedure utilizing contrast, or any open surgical procedure. If the "next procedure" is a complication from the entered procedure, continue to abstract the peak creatinine beyond it.

If only one value is available post procedure through discharge, that value will be used for both the post procedure peak creatinine and the discharge value.

Requirea:
Yes
Suffix:
mg/dl
Minimum:
0.1
Maximum:
15
Soft Minimum:
0.3

### Post Procedure Nadir Hemoglobin

#### **Data Abstraction Instructions:**

Record the lowest level recorded from the end of the procedure to the next procedure or discharge, whichever occurs first. If there is no value drawn post procedure, mark "Not drawn."

#### Selections:

Required:

- Yes
- Enter value
- Not drawn

#### Supporting Definitions:

The next procedure is any invasive procedure that could potentially result in significant blood loss. If the "next procedure" is a complication from the entered procedure, continue to abstract the nadir hemoglobin beyond it.

If only one value is available post procedure through discharge, that value will be used for the post procedure nadir hemoglobin **and** the discharge hemoglobin.

Yes		
Suffix		
a/dl		
g/ui		
Minimum:		
2		
Maximum:		
20		
Soft Minimum:		
5		
Soft Maximum:		
18		

### Post Procedure COVID-19

#### **Data Abstraction Instructions:**

Indicate the COVID-19 status for this discharge/episode of care. If there are multiple results available post procedure through discharge, please enter the first positive result available post procedure. If there are no positive results, please enter the first negative result available post procedure.

**Positive COVID-19\*** = patient has a positive test result for COVID-19\* post procedure. Specimen\*\* may have been obtained, either prior to or post procedure and was positive after the procedure. If a specimen\*\* obtained during this hospitalization is positive for COVID-19\* at time of abstraction, please select this field even if results were posted after discharge.

If Positive COVID-19 result, enter the date and time of Positive Result Posted. If a date is documented but the time is not documented enter the date and then enter 00:00 for the time. If date and time are not documented, enter Not Documented.

**Negative COVID-19\*** = patient has negative test result for COVID-19 post procedure. Specimen obtained, either prior to or post procedure and was negative after the procedure. If a specimen\*\* obtained during this hospitalization is negative for COVID-19\* at time of abstraction, please select this field even if results were posted after discharge.

If Negative COVID-19 result, enter the date and time of Positive Result Posted. If a date is documented but the time is not documented enter the date and then enter 00:00 for the time. If date and time are not documented, enter Not Documented.

No COVID-19\* Result = Specimen\*\* obtained, but no result at time of abstraction.

No COVID-19\* Specimen\*\* = patient was not tested for COVID-19 during this hospitalization.

No Change in COVID Status = patient was identified as "Positive COVID-19"," Negative COVID-19", or "Recovered COVID-19" prior to the first procedure and there has been no change in patient condition prompting further testing.

Timeframe: Post procedure through discharge from hospital. While the result timeframe you are addressing is post procedure, the actual specimen may have been obtained prior to the VS or carotid procedure. If patient has multiple procedures, please select response based on COVID-19 status after last procedure.

\*Please include any of the viruses associated with the COVID-19 pandemic (2019-nCOV, MERS-CoV, SARS-CoV, SARS-CoV2, and other associated viruses identified by the CDC)

\*\*Specimen sites: lower respiratory, upper respiratory, serology, sputum, stool, urine, oropharyngeal or nasopharyngeal swab.

- Positive COVID-19
  - Documented
    - Enter Date/Time: MM/DD/YYYY HH:MM
  - Not Documented
- Negative COVID-19
   Documented
  - Enter Date/Time: MM/DD/YYYY HH:MM
  - Not Documented
  - No COVID-19 Result
- COVID-19\* Specimen
- No Change in COVID Status

### Supporting Definitions:

World Health Organization. (2020, March 2). WHO.int. Laboratory testing for coronavirus disease 2019 (COVID-19) in suspected human cases. Interim guidance.

https://www.who.int/emergencies/diseases/novel-coronavirus-2019/technical-guidance/laboratory-guidance/

### Required:

### Yes

### Labs - Other

### Albumin

### **Data Abstraction Instructions:**

Record the most recent value within 6 months of procedure. If no value is available pre-procedure, a value drawn during the current hospitalization may be used (if multiple values are available, enter the value closest to the procedure). If there is no value, mark "Not drawn."

### Selections:

- Yes
  - Enter value
- Not drawn

### Supporting Definitions:

Reference range 3.5 - 4.9 g/dL

Required: Yes Minimum: 0 Maximum: 7

### Indications

### Indication Type

#### Selections:

- EVAR/AAA Revascularization
- Lower Extremity Revascularization
- Upper Extremity Revascularization
- Mesenteric Revascularization
- Renal Revascularization

Required: Yes

### Indications for EVAR/AAA Revascularization

### Asymptomatic

### **Data Abstraction Instructions:**

Indicate if patient presents for this hospitalization without showing any symptoms of AAA.

### Selections:

- Yes
- No

### Supporting Definitions:

The size of the aneurysm has reached a point where surgery is determined necessary.

### Required:

Yes

### Abdominal/Back Pain

### **Data Abstraction Instructions:**

Indicate if patient presented with abdominal and/or back pain attributed to the aneurysm.

No

Required: Yes

### **Rapidly Increasing Aneurysm Diameter**

### Data Abstraction Instructions:

Indicate if there has been an increase in aneurysm diameter by 0.5 cm within 6 months to one year as determined by CTA.

#### Selections:

- Yes
- No

### Supporting Definitions:

There has been rapid growth of the aneurysm necessitating surgery.

Required: Yes

Suffix: cm

### Unfit for open AAA repair

### **Data Abstraction Instructions:**

Endovascular repair performed because patient was considered too high risk by surgeon for open repair.

### Selections:

Yes

No

### Required:

Yes

### Unfit for general anesthesia

#### **Data Abstraction Instructions:**

Endovascular repair performed because patient was considered too high risk by surgeon or anesthesiologist for general anesthesia.

### Selections:

- Yes
- No

### Required:

Yes

### Infection

### **Data Abstraction Instructions:**

Indicate if the procedure was performed due to an infection from a previous procedure.

### Selections:

- Yes
- No

### Required:

Yes

### Size of Iliac Aneurysm

**Data Abstraction Instructions:** 

Indicate if the primary reason for AAA repair is the size of an iliac aneurysm.

### Selections:

- Yes
- No

### Required:

Yes

### Correction of Endoleak

### **Data Abstraction Instructions:**

Please indicated if the procedure is being perform to correct an endoleak from a prior procedure.

Required: Yes

### **Concomitant Iliac Occlusive Disease**

### **Data Abstraction Instructions:**

Indicate of the procedure is being performed with the presence of concomitant iliac occlusive disease.

### Supporting Definitions:

If the patient has documented ABIs, please enter them in the pre-imaging fields.

**Required:** 

Yes

### Lower Extremity Ischemia/Emboli

### **Data Abstraction Instructions:**

Indicate if the procedure is being performed due to the presence of lower extremity ischemia/emboli due to the aneurysm.

Required: Yes

### **Documented Patient Anxiety Levels**

#### **Data Abstraction Instructions:**

Indicate if the aneurysm repair is being performed due to documented patient anxiety about aneurysm presence/growth/rupture.

#### Supporting Definitions:

There must be supporting documentation in the medical record to select yes.

**Required:** 

#### Yes

### **Penetrating Ulcer**

#### **Data Abstraction Instructions:**

Indicate if there is a penetrating ulcer present in the aneurysm and it is part of the indication for repair.

If a penetrating ulcer is present, indicate the size (mm).

### Selections:

- Yes
   Enter value in mm
   Not documented
- No

Required: Yes Minimum: 0

Maximum: 30

### **Complication from Prior Procedure**

#### **Data Abstraction Instructions:**

Indicate if the patient had a complication from a prior procedure.

#### Selections:

- Yes
- No

#### Supporting Definitions:

The patient had a dissection, perforation or another complication from a prior procedure (such as a cardiac or pulmonary angiogram) that required vascular repair (Percutaneous or Surgical).

### **Required:**

Yes Trauma

### Data Abstraction Instructions:

Indicate if the vascular procedure was performed to repair injury that resulted from trauma (motor vehicle accident, fire, fall etc.).

Yes
 No

#### • 140

Required: Yes

### Mycotic Aneurysm

#### Data Abstraction Instructions:

Indicate if the procedure was performed to repair an infected abdominal aortic aneurysm. When choosing the Indication of Mycotic Aneurysm for a primary AAA repair, do not enter the Indication of Infection.

#### Required:

Yes

### **Pre-procedure Smoking Cessation**

#### **Data Abstraction Instructions:**

Indicate if the patient received physician delivered advice, a prescription for nicotine replacement, and/or a referral for smoking cessation services submitted on the patient's behalf prior to admission to the hospital. These interventions would be implemented to prepare the patient for the current procedure. Choose all that apply.

#### Selections:

- Yes
  - Physician delivered advice
    - Nicotine replacement therapy (NRT)
    - Referral to smoking counseling services
- No

### Supporting Definitions:

Yes = Enter Yes for Smoking Cessation at Discharge if Yes was entered for Current Smoker under Patient History / Comorbidity, and one of the 3 steps were implemented prior to admission to the hospital.

If the physician recommended smoking cessation prior to admission to the hospital, and the patient refused, enter Yes smoking cessation counseling. There must be adequate documentation to support this claim.

Physician delivered advice = A surgeon, advanced practice personnel (PA, NP), or resident has a conversation with the patient and recommends that the patient stops smoking.

A recommendation to stop smoking offered by a nurse, respiratory therapist, or student does not count as physician delivered advice.

Nicotine replacement therapy (NRT) = NRT may include a nicotine patch, gum, lozenge, or other pharmacologic assistance (Varenicline or Bupropion) prior to admission to the hospital.

If a patient refuses nicotine replacement therapy, and there is physician documentation that NRT was offered and documentation that the patient refused, you can choose NRT.

**Referral to smoking counseling services** = A referral to a smoking cessation program or class is submitted on the patient's behalf. Giving the patient the Michigan Tobacco Quit line phone number and/or website is not sufficient.

If a physician, mid-level provider, or resident does an assessment and then puts in a referral to RT or a dedicated smoking cessation nurse to provide smoking cessation education, you can choose referral to smoking counseling services.

No = Enter No for Smoking Cessation at Discharge if No was entered for Ever Smoked or Current Smoker under Patient History / Comorbidity. Enter No if the patient is a current smoker; however, none of the 3 steps were implemented prior to admission to the hospital.

#### **Required:**

Yes

### Indications for Lower Extremity Revascularization (Open Bypass / Open Thrombectomy)

### Claudication

#### **Data Abstraction Instructions:**

Indicate if the patient has leg pain caused by poor circulation, inhibiting patient's ability to walk distances.

#### Selections:

- Yes
- No

### Supporting Definitions:

Refers to cramping pains in the legs (usually the calf muscles, but may be in the thigh muscles) caused by poor circulation of the blood in the arteries to the leg muscles during exercise. True claudication is relieved with rest from exercise. If the patient has arm claudication from subclavian stenosis, do not include.

### Required:

Yes

### **Rest Pain**

### **Data Abstraction Instructions:**

Indicate if the patient has severe pain in the foot and toes that is not readily controlled by analgesics that is made worse by elevation of the leg and relieved by dependency.

### Selections:

- Yes
- No

Required:

Yes

### **Threatened Bypass Graft**

### **Data Abstraction Instructions:**

Indicate if the procedure performed is to maintain patency of a previously placed bypass graft. This may be demonstrated via abnormal duplex scans, abnormal non-invasive imaging, or increased velocities identified during surveillence visits.

#### Selections:

- Yes

   Symptomatic
   Asymptomatic
- No

#### **Required:**

Yes

#### Acute Limb Ischemia

#### **Data Abstraction Instructions:**

Indicate if there is any sudden decrease in limb perfusion that causes a potential threat to limb viability.

### Selections:

- Yes
- No

#### Supporting Definitions:

Critical limb ischemia is NOT acute limb ischemia. Critical limb ischemia is the disease process with the presence of rest pain or tissue loss/ulcer. Acute limb ischemia is an urgent or emergent situation in which intervention is necessary to prevent the loss of limb/function. It may be indicated by sudden onset pain, pallor, or numbness/paralysis.

#### **Required:**

Yes

### Failed Endovascular Procedure

### Data Abstraction Instructions:

Indicate if the procedure was performed for a failed endovascular intervention.

### Selections:

YesNo

### Supporting Definitions:

A failed endovascular procedure is one performed on the ipsilateral limb (same side as current procedure) within the same vascular bed within the last 30 days.

Example: Two weeks prior, patient had a left SFA stent. Now presents for a Left Common Femoral to Popliteal bypass.

### Required: Yes

### Infection (Lower Extremity Revascularization)

### Data Abstraction Instructions:

Indicate if the procedure was performed due to an infection from a prior procedure.

### Selections:

- Yes
- No

### Supporting Definitions:

### https://bmc2.org/print/book/export/html/78

This captures vascular surgeries performed due to an infected graft, wound, or other sources of infection from a prior procedure; not those indicated for ulcers/wound healing.

### Required:

Yes

### **Facilitation of Procedure**

### **Data Abstraction Instructions:**

Indicate if the PVI procedure performed was to facilitate a different endovascular procedure (EVAR, TAVR, etc.).

#### Selections:

- Yes
- No

Required:

Yes

### Pre-Procedure Exercise Therapy

### Data Abstraction Instructions:

Indicate if there is documentation that the patient was on or failed some type of pre-procedure exercise program prior to the hospitalization. If so, indicate whether it was informal or structured.

### Selections:

- Yes
  - Structured/Supervised
- Home Based/InformalNo

Required: Yes

#### Impaired Ability to Work

#### **Data Abstraction Instructions:**

Indicate if the procedure is performed due to an inability to work.

### Selections:

- Yes
- No

### Required:

Yes

### Peripheral Aneurysm Repair

### Data Abstraction Instructions:

Indicate if the procedure is being performed for repair of a peripheral aneurysm and whether the patient is experiencing symptoms.

### Selections:

- Yes
  - Symptomatic
- AsymptomaticNo

### Required:

Yes

### **Increased Stent Velocity**

### Data Abstraction Instructions:

Indicate if the procedure performed is due to increased velocities in a pre-existing stent and whether the patient is experiencing symptoms. This may be demonstrated via abnormal duplex scans, abnormal non-invasive imaging, or increased velocities identified during surveillance visits.

#### Selections:

Yes

 Symptomatic
 Asymptomatic

 No

### Required:

Yes

### **Increased Stent Graft Velocity**

**Data Abstraction Instructions:** 

### https://bmc2.org/print/book/export/html/78

Indicate if the procedure performed is due to increased velocities in a pre-existing stent graft and whether the patient is experiencing symptoms. This may be demonstrated via abnormal duplex scans, abnormal non-invasive imaging, or increased velocities identified during surveillance visits.

#### Selections:

- Yes
  - Symptomatic
- AsymptomaticNo

Required:

Yes

### Wound (WIfl)

### **Data Abstraction Instructions:**

Indicate if the patient has a wound present and to what degree.

Grade	Ulcer	Gangrene
0	No ulcer	No gangrene
Clinical description	: ischemic rest pain (requires typical symptoms + ischemia grade 3); i	no wound.
1	Small, shallow ulcer(s) on distal leg or foot; no exposed bone, unless limited to distal phalanx	No gangrene
Clinical description	: minor tissue loss. Salvageable with simple digital amputation (1 or 2	digits) or skin coverage.
2	Deeper ulcer with exposed bone, joint or tendon; generally not involving the heel; shallow heel ulcer, without calcaneal involvement	Gangrenous changes limited to digits
Clinical description	: major tissue loss salvageable with multiple $(\geq 3)$ digital amputations	or standard TMA ± skin coverage.
3	Extensive, deep ulcer involving forefoot and/or midfoot; deep, full thickness heel ulcer ± calcaneal involvement	Extensive gangrene involving forefoot and /or midfoot; full thickness heel necrosis ± calcaneal involvement
Clinical description flap coverage or	: extensive tissue loss salvageable only with a complex foot reconstruction complex wound management needed for large soft tissue defect	on or nontraditional TMA (Chopart or Lisfranc):

#### Selections:

Yes

- Grade 1: Minor tissue loss; small shallow ulceration
- Grade 2: Major tissue loss; deeper ulceration with exposed bone, joint, or tendon
- Grade 3: Extensive ulcer/gangrene;
- Not Documented
- No

### Required:

Yes

### Ischemia (WIfI)

### Data Abstraction Instructions:

Indicate the degree of ischemia present.

Grade	ABI	Ankle systolic pressure	$TP, TcPO_2$
0	≥0.80	>100 mm Hg	≥60 mm Hg
1	0.6-0.79	70-100 mm Hg	40-59 mm Hg
2	0.4-0.59	50-70 mm Hg	30-39 mm Hg
3	≤0.39	<50 mm Hg	<30 mm Hg

ABI, Ankle-brachial index; PVR, pulse volume recording; SPP, skin perfusion pressure; TP, toe pressure; TePO2, transcutaneous oximetry.

Patients with diabetes should have TP measurements. If arterial calcification precludes reliable ABI or TP measurements, ischemia should be documented by TcPO<sub>2</sub>, SPP, or PVR. If TP and ABI measurements result in different grades, TP will be the primary determinant of ischemia grade.

Flat or minimally pulsatile forefoot PVR = grade 3.

### Selections:

Yes

- Grade 0Grade 1
- Grade 2

- Grade 3
- Not Documented

No

Required:

#### Yes

### Foot Infection (WIfl)

### **Data Abstraction Instructions:**

Indicate if the patient has a foot infection and to what degree.

Grade	Clinical Description	IDSA	IWGDF Class
0	wound without purulence or manifestations of infection	uninfected	1
1	>2 manifestations of infection (erythema or purulence, pain tenderness, warmth or induration) any cellulitis or crythema extends < 2cm around ulcer; infection is limited to skin or subcutaneous tissues; no local complications or systemic illness	mild	2
2	Infection in patient who is systemically and metabolically stable but has ≥1 of the following: cellulitis extending 2cm, lymphangitis; spread beneath fascia; deep tissue abscess; gangrene; muscle, tendon, joint or bone involvement	moderate	3
3	Infection in patient with systemic or metabolic toxicity	severe	4

### Selections:

- Yes
  - · Grade 0: Wound without purulence or manifestations of infection
  - Grade 1: >2 manifestations of infection (erythema, purulence, pain, warmth, etc.) 0
    - · Grade 2: Infection in patient who is systemically stable but has one or more of the following; cellulitis extending 2cm, spread beneath fascia, deep tissue abscess, gangrene, muscle/tendon/bone involvement
  - Grade 3: Infection in patient with systemic or metabolic toxicity
  - Not Documented
- No

### **Required:**

Yes

### **Complication from Prior Procedure**

### **Data Abstraction Instructions:**

Indicate if the patient had a complication from a prior procedure.

### Selections:

- Yes
- No

### Supporting Definitions:

The patient had a dissection, perforation or another complication from a prior procedure (such as a cardiac or pulmonary angiogram) that required vascular repair (Percutaneous or Surgical).

### **Required:**

Yes

### Trauma

### **Data Abstraction Instructions:**

Indicate if the vascular procedure was performed to repair injury that resulted from trauma (motor vehicle accident, fire, fall etc.).

### Selections:

 Yes No

Required: Yes

### **Pre-procedure Smoking Cessation**

### **Data Abstraction Instructions:**

Indicate if the patient received physician delivered advice, a prescription for nicotine replacement, and/or a referral for smoking cessation services submitted on the patient's behalf prior to admission to the hospital. These interventions would be implemented to prepare the patient for the current procedure. Choose all that apply.

#### Selections:

- Yes
  - Physician delivered advice
  - Nicotine replacement therapy (NRT)
     Referral to smoking counseling services
- Referral
   No

#### Supporting Definitions:

Yes = Enter Yes for Smoking Cessation at Discharge if Yes was entered for Current Smoker under Patient History / Comorbidity, and one of the 3 steps were implemented prior to admission to the hospital.

If the physician recommended smoking cessation prior to admission to the hospital, and the patient refused, enter Yes for smoking cessation counseling. There must be adequate documentation to support this claim.

Physician delivered advice = A surgeon, advanced practice personnel (PA, NP), or resident has a conversation with the patient and recommends that the patient stops smoking.

A recommendation to stop smoking offered by a nurse, respiratory therapist, or student does not count as physician delivered advice.

Nicotine replacement therapy (NRT) = NRT may include a nicotine patch, gum, lozenge, or other pharmacologic assistance (Varenicline or Bupropion) prior to admission to the hospital.

If a patient refuses nicotine replacement therapy, and there is physician documentation that NRT was offered and documentation that the patient refused, you can choose NRT.

**Referral to smoking counseling services** = A referral to a smoking cessation program or class is submitted on the patient's behalf. Giving the patient the Michigan Tobacco Quit line phone number and/or website is not sufficient.

If a physician, mid-level provider, or resident does an assessment and then puts in a referral to RT or a dedicated smoking cessation nurse to provide smoking cessation education, you can choose referral to smoking counseling services

No = Enter No for Smoking Cessation at Discharge if No was entered for Ever Smoked or Current Smoker under Patient History / Comorbidity. Enter No if the patient is a current smoker; however, none of the 3 steps were implemented prior to admission to the hospital.

#### **Required:**

Yes

### Indications for Upper Extremity Revascularization (Open Bypass / Open Thrombectomy)

### **Ulcer/Gangrene**

### Data Abstraction Instructions:

Indicate if the patient has an ulcer, gangrene, or if tissue loss is present.

### Selections:

Yes

No

Required: Yes

### Acute Limb Ischemia

### **Data Abstraction Instructions:**

Indicate if there is any sudden decrease in limb perfusion that causes a potential threat to limb viability.

### Selections:

- Yes
- No

### Supporting Definitions:

Critical limb ischemia is NOT acute limb ischemia. Critical limb ischemia is the disease process with the presence of rest pain or tissue loss/ulcer. Acute limb ischemia is an urgent or emergent situation in which intervention is necessary to prevent the loss of limb/function. It may be indicated by sudden onset pain, pallor, or numbness/paralysis.

### Required:

Yes

### **Angina/Abnormal Cardiac Stress Test**

#### **Data Abstraction Instructions:**

Indicate if patient has had episodes of angina, or if patient has had an abnormal cardiac stress test.

#### Selections:

- Yes
- No

### Supporting Definitions:

Indication for revascularization is cardiac ischemia secondary to impaired blood flow to coronary artery bypass graft (e.g. left subclavian stenosis in a patient with an in situ left internal mammary artery bypass graft).

For informational purposes, one of the following criteria are necessary:

- Angina at rest (usually prolonged >20 mins)
- New onset (less than two months) exertional angina of at least Canadian cardiovascular Society Classification (CCSC) class III
- Recent (less than two months) acceleration of angina reflected by an increase in severity of at least one CCSC class to at least CCSC class III. The patient
  must also NOT have any biochemical evidence of myocardial necrosis.

#### Required: Yes

#### ------

### BP discrepancy

### Data Abstraction Instructions:

Indicate if the patient has a more than 50 mm difference in systolic BP between L and R arms.

### Selections:

- Yes
- No

### Supporting Definitions:

This may be seen in subclavian stenosis.

#### Required: Yes

### 165

### Arm Claudication

### **Data Abstraction Instructions:**

Indicate if the patient has arm pain caused by poor circulation.

### Selections:

- Yes
- No

### Supporting Definitions:

Refers to cramping pains in the arms caused by poor circulation of the blood in the arteries to the arm muscles during exercise. True claudication is relieved with rest from exercise.

#### **Required:**

Yes

# Peripheral Aneurysm Repair

### Data Abstraction Instructions:

Indicate if the procedure is being performed for repair of a peripheral aneurysm and whether the patient is experiencing symptoms.

### Selections:

Yes

 Symptomatic
 Asymptomatic

 No

### Required:

Yes

### **Complication from Prior Procedure**

### **Data Abstraction Instructions:**

Indicate if the patient had a complication from a prior procedure.

- Selections:
  - YesNo

#### Supporting Definitions:

The patient had a dissection, perforation or another complication from a prior procedure (such as a cardiac or pulmonary angiogram) that required vascular repair (Percutaneous or Surgical).

#### **Required:**

Yes

#### Trauma

#### **Data Abstraction Instructions:**

Indicate if the vascular procedure was performed to repair injury that resulted from trauma (motor vehicle accident, fire, fall etc.).

#### Selections:

- Yes
- No

### **Required:**

Yes

### Pre-procedure Smoking Cessation

### **Data Abstraction Instructions:**

Indicate if the patient received physician delivered advice, a prescription for nicotine replacement, and/or a referral for smoking cessation services submitted on the patient's behalf prior to admission to the hospital. These interventions would be implemented to prepare the patient for the current procedure. Choose all that apply.

#### Selections:

### Selections:

- Yes
  - Physician delivered advice
  - Nicotine replacement therapy (NRT)
  - · Referral to smoking counseling services
- No

#### Supporting Definitions:

Yes = Enter Yes for Smoking Cessation at Discharge if Yes was entered for Current Smoker under Patient History / Comorbidity, and one of the 3 steps were implemented prior to admission to the hospital.

If the physician recommended smoking cessation prior to admission to the hospital, and the patient refused, enter Yes for smoking cessation counseling. There must be adequate documentation to support this claim.

Physician delivered advice = A surgeon, advanced practice personnel (PA, NP), or resident has a conversation with the patient and recommends that the patient stops smoking.

A recommendation to stop smoking offered by a nurse, respiratory therapist, or student does not count as physician delivered advice.

Nicotine replacement therapy (NRT) = NRT may include a nicotine patch, gum, lozenge, or other pharmacologic assistance (Varenicline or Bupropion) prior to admission to the hospital.

If a patient refuses nicotine replacement therapy, and there is physician documentation that NRT was offered and documentation that the patient refused, you can choose NRT.

**Referral to smoking counseling services** = A referral to a smoking cessation program or class is submitted on the patient's behalf. Giving the patient the Michigan Tobacco Quit line phone number and/or website is not sufficient.

If a physician, mid-level provider, or resident does an assessment and then puts in a referral to RT or a dedicated smoking cessation nurse to provide smoking cessation education, you can choose referral to smoking counseling services.

No = Enter No for Smoking Cessation at Discharge if No was entered for Ever Smoked or Current Smoker under Patient History / Comorbidity. Enter No if the patient is a current smoker; however, none of the 3 steps were implemented prior to admission to the hospital.

#### **Required:**

Yes

### **Procedure Details**

Choose the type of Vascular Surgery procedure to be performed.

- Open Bypass/Revascularization
- Open Abdominal Aortic Aneurysm
- Endovascular Aneurysm Repair (EVAR)

### **Open Bypass/Revascularization**

#### **Qualifying Procedures:**

- · Upper Extremity Bypass or Lower Extremity Bypass for the indication stenosis or aneurysm repair
- Bypass that includes a Graft Origin (where the graft was proximally attached to the artery) and a Graft Insertion (the distal attachment to the artery)
- A procedure where the aneurysm is cut out and replaced with an interposition graft
- · Bypass of a previous bypass
- · An Open Bypass revision where the old graft is removed and replaced with a new graft
- · An Open Bypass revision where a new graft is connected to an old graft
- A qualifying Open Bypass where the primary incision was made, but the procedure was not completed
- Patch angioplasty, open endarterectomy, open thromboembolectomy, aneurysm, or pseudoaneurysm repair as a concomitant procedure to an open bypass. You would enter these concomitant procedures under the Locations section of the website.

#### Open Bypass procedures that do not qualify:

- Visceral bypass
- · Carotid-subclavian bypass
- Carotid-carotid bypass
- Renal bypass
- Mesenteric bypass
- · Revision is done at one end of the graft (for example, a kink was fixed)
- AV fistula procedure, repair or intervention
- Dialysis graft procedure, repair or intervention
- · A qualifying open bypass where the procedure was aborted BEFORE the primary incision was made
- A bypass graft revision that did not have a new graft implanted
- · Patch on anastomosis

### **Graft Origin**

### **Data Abstraction Instructions:**

Indicate if the patient had an open bypass, and if yes select origin point on vessel using the vessel artery map.

### Selections:

· Select artery name from list

### Supporting Definitions:

This is the location where arterial flow enters the graft.

### Required:

Yes

### Graft Insertion

### Data Abstraction Instructions:

Select insertion point on vessel. If two insertion sites are used, indicate both sites.

### Selections:

· Select artery name from list

### Supporting Definitions:

This is the location where arterial flow exits the graft.

### Required:

Yes

### Redo Procedure

### **Data Abstraction Instructions:**

Indicate if the procedure being performed is a redo of a previous bypass.

### Selections:

Yes

### Supporting Definitions:

The intent is to capture bypass procedures that are performed to revascularize the same arterial bed as a previous bypass. It does not necessarily require the same origin and insertion.

#### **Required:**

Yes

### Vein Graft

### **Data Abstraction Instructions:**

Indicate if a vein graft was used for bypass. If yes, indicate type of vein graft.

### Selections:

Yes

- Reversed (greater saphenous vein) GSV
- In Situ GSV
- Non-reversed transposed GSV
- Lesser saphenous
- Cephalic
- Basilic
- Allograft
- Composite
- Other

• No

### Supporting Definitions:

Reversed GSV = an incision is made at the proximal and distal ends of the vein. The vein is reversed to allow blood to flow through.

In Situ GSV = The GSV can be used In Situ, in its native position in the vascular bed where only the origin and insertion are maneuvered to create an anastomosis to the artery.

Non-reversed transposed GSV = tTe GSV can be Non-Reversed Transposed, or moved, to create a bypass.

Lesser saphenous = a venous blood vessel that runs up the length of the leg. It originates from the junction formed between two small veins in the foot, the fifth toe's dorsal vein and the dorsal venous arch.

Cephalic = The cephalic vein courses through both the forearm and arm and terminates by draining into the axillary vein.

Basilic = The basilic vein lies in the deep subcutaneous tissue at the antecubital crease and pierces the brachial fascia in the distal third of the upper arm.

Allograft = A donor cadaver vein.

Composite = A combination of vein graft and prosthetic graft.

Other = Options that are not in the list above, such as Xenograft.

Required: Yes

#### Vein Graft Harvest

#### **Data Abstraction Instructions:**

Indicate the type of vein that was harvested from the patient during the current open bypass surgery. If a prosthetic graft, allograft or xenograft is implanted, enter Not Harvested.

#### Selections:

- Open
- Endoscopic
- · Not harvested

### Supporting Definitions:

**Open** = A long incision or skip incisions are made into the skin.

Endoscopic = small puncture incisions and an endoscope with a CO2 balloon, a dissector tool, a harvesting tool and a cutting tool using cautery to cut side branches. Not harvested = A prosthetic graft, allograft or xenograft was used.

### Required:

No

### Number of Vein Segments

### Data Abstraction Instructions:

Indicate the number of vein segments used in bypass procedure.

- One
- Two
- Three or more

Required: Yes

### **Minimal Vein Graft Diameter**

### **Data Abstraction Instructions:**

Indicate the minimal vein graft diameter of the graft being used for bypass. Value can be found in duplex venous imaging pre-operatively or dictated by the physician.

### Selections:

· Record Value in mm

### Required:

No Suffix: mm Minimum: 0.5 Maximum: 10

#### Prosthetic Graft

### Data Abstraction Instructions:

Indicate the type of prosthetic graft used. Select all that apply.

### Selections:

Yes

 Dacron
 PTFE
 Composite with vein

 No

Required:

Yes

#### Vein Cuff

#### **Data Abstraction Instructions:**

Indicate if a vein cuff was used at anastomosis.

#### Selections:

- Yes
- No

### Supporting Definitions:

Prosthetic grafts can be anastomosed directly to the recipient artery. However, several techniques incorporating vein into the distal anastomosis have been utilized in an effort to improve long-term patency.

#### **Required:**

Yes

### **Intra Operative Graft Patency**

### **Data Abstraction Instructions:**

Indicate the method used to assess graft patency at the end of the procedure and if it was normal or abnormal. This should indicate flow through the graft itself and is not related to outflow vessels.

### Selections:

- Yes
  - Doppler
  - DuplexAngiogram
    - Normal
    - Abnormal
- No

### Required:

Yes

### Intra Operative Graft Revision

### **Data Abstraction Instructions:**

Indicate if intraoperative graft revision was performed.

### Selections:

Yes

No

### Supporting Definitions:

A graft revision is a part of the procedure that is unplanned after the initial graft is sewn into place and tested for patency. For example: revision for thrombus after poor doppler signals, kinking, or anastomosis revision.

#### **Required:**

Yes

### Closure for Open Exposure

### Data Abstraction Instructions:

Identify all that apply for incision closure. Include all layers of closure - muscle, subcutaneous, and skin.

#### Selections:

- Suture
  - Absorbable
- Permanent
   Staples
- Delayed
- Other

Required: Yes

#### 163

### **Open Abdominal Aortic Aneurysm (OAAA)**

#### **Qualifying Procedures:**

- · Open Infrarenal, Juxtarenal, and Suprarenal AAA repair
- · Ruptured AAA. Even if the patient expires after the primary incision was made
- · A qualifying OAAA where the primary incision was made, but the procedure was not completed
- · An EVAR that was converted to an OAAA during the same OR time
- An Open AAA that was performed for the Indication of Penetrating Ulcer without the presence of an aneurysm.
- Patch angioplasty, open endarterectomy, open thromboembolectomy, or pseudoaneurysm repair as a concomitant procedure to an open AAA. You would enter
  these concomitant procedures under the Locations section of the website

### OAAA procedures that do not qualify:

- An open aneurysm repair in the thoracic abdominal aorta (above the diaphragm)
- An OAAA that is done for the indication of aortic stenosis or pseudoaneurysm repair
- A qualifying OAAA where the procedure was aborted BEFORE the primary incision was made
- · Patch on anastomosis

### **Prior Family History of AAA**

#### **Data Abstraction Instructions:**

Indicate if relative (parent, sibling, aunt, uncle, child) has/had a history of AAA.

#### Selections:

- Yes
- No

Required:

Yes

## Prior Aortic Surgery (OAAA)

### **Data Abstraction Instructions:**

Enter the year, and type of repair, of the prior procedure (yyyy).

# Selections:

- Yes Enter year
- Select option
  - AAA (Infrarenal)
  - SAAA (Suprarenal)
     Bypass
  - 0 Bypass
  - Other (Endarterectomy or Other)

No

Required: No Minimum: 1900 Maximum: 9999

### Maximum AAA Diameter (OAAA)

### **Data Abstraction Instructions:**

Enter AAA largest diameter in millimeters. For ruptured AAA procedures, use not documented only if value not available.

### Selections:

- Documented
  - Enter value in mm
- Not documented

### Supporting Definitions:

Use largest diameter. If multiple imaging modality, use most accurate in following hierarchy: CT > MRI > ECHO > Arteriogram.

For AAA repair performed due only to iliac aneurysm size (no abdominal aortic aneurysm), enter Not Documented.

Required: Yes Suffix: mm Minimum: 25 Maximum: 200

### Iliac Aneurysm (OAAA)

### **Data Abstraction Instructions:**

Indicate if the patient had an iliac aneurysm. Choose if Unilateral, Bilateral or None. If yes, enter value for maximum diameter in millimeters.

### Selections:

- Yes
  - Unilateral
    - Bilateral
       Enter value in mm
- No

#### Supporting Definitions:

Iliac diameter > 1.5 cm. Maximum diameter of largest iliac artery, common or internal.

Required: Yes Suffix: mm Minimum: 10 Maximum: 100

### Aneurysm location (OAAA)

#### **Data Abstraction Instructions:**

Enter the aneurysm location as described in the pre-imaging or surgical dictation.

### Selections:

- Infrarenal
- Juxtarenal
- Suprarenal
- Not documented

Required:

Yes

### Aneurysm anatomy (OAAA)

### **Data Abstraction Instructions:**

Enter the aneurysm anatomy as described in the pre-imaging or surgical dictation.

#### Selections:

- Fusiform
- Saccular
- · Not documented

### Required:

Yes

### Contained rupture (OAAA)

### **Data Abstraction Instructions:**

Indicate if the aneurysm was described as a contained rupture on pre-imaging or in the physician dictation.

#### Required: Yes

### Ruptured AAA (OAAA)

### **Data Abstraction Instructions:**

Indicate if the anueuysm was ruptured at the time of the procedure. This includes free ruptures with extravasation of blood/contrast on pre-imaging or in the physician dictation.

#### Selections:

- Yes
- No

### Supporting Definitions:

Contained ruptures do not qualify.

### Required:

Yes

### Lowest Pre-Intubation Blood Pressure (for ruptured AAA repairs only) (OAAA)

### **Data Abstraction Instructions:**

Indicate the lowest pre-intubation BP, after arrival at hospital, use systolic pressure.

### Selections:

- Yes
  - Enter value in mmHg
- Not documented

Required: Yes Suffix: mmHg Minimum: 0 Maximum: 200

### Mental Status (for ruptured AAA repairs only) (OAAA)

### Data Abstraction Instructions:

Indicate the patient's mental status on arrival to the operating room.

### Selections:

- Yes
  - Normal (alert and oriented)
  - Disoriented to person, place, or time
  - Unconscious
- Not documented

### **Required:**

### Yes

### Cardiac Arrest (for ruptured AAA repairs only) (OAAA)

### Data Abstraction Instructions:

Indicate if the patient was in cardiac arrest on arrival to the operating room.

### Selections:

- Yes
- No

```
Required:
Yes
```

### res

### Timeframe: Symptoms to Incision (for ruptured AAA repairs only) (OAAA)

### Data Abstraction Instructions:

Record the time from the patients first symptoms until the surgical incision in hours.

- Yes
  - Enter value in hours

#### **Data Abstraction Instructions:**

Record the time from when the patient was admitted to the surgical incision in hours.

#### Selections:

- Yes
  - Enter value in hours
     Not documented

### Required:

Yes Suffix: hours

Minimum: 0 Maximum:

24

### Abdomen Explored (OAAA)

### **Data Abstraction Instructions:**

Indicate if the abdomen was explored by the surgeon to evacuate hematoma, but not to repair rupture.

### Selections:

- Yes
- No

#### **Required:**

Yes

### Conversion from Endovascular Repair (OAAA)

### **Data Abstraction Instructions:**

Indicate if the endovascular aneurysm repair had to be converted to an open procedure If yes, the time frame for the conversion.

### Selections:

Yes

 Immediate
 > 1 day to 30 days
 > 30 days

 No

### Supporting Definitions:

Include EVARs converted to open procedures even after multiple years.

### Required:

Yes

### Exposure (OAAA)

### **Data Abstraction Instructions:**

Indicate the exposure used for AAA repair.

### Selections:

- Transperitoneal
- Retroperitoneal

### Supporting Definitions:

Transperitoneal indicates through the peritoneum - look for midline incision. Retroperitoneal indicates behind the peritoneum - look for lateral positioning, flank incision.

### Required:

Yes

### **Distal Anastomosis (OAAA)**

### **Data Abstraction Instructions:**

Indicate most distal extent of either right or left limb if bifurcated.

### Selections:

- Aorta
- Common Iliac Artery (CIA)
- External Iliac Artery (EIA)
- Common Femoral Artery (CFA)
- Graft Not Utilized

### Required:

Yes

### Graft Body Diameter (OAAA)

### **Data Abstraction Instructions:**

Body size is diameter of most proximal portion of graft.

#### Selections:

- Documented
- Enter value in mm
- Not documented
- Graft Not Utilized

#### **Required:**

Yes Suffix: mm Minimum: 8 Maximum: 40

### Graft Type (OAAA)

#### **Data Abstraction Instructions:**

Indicate type of graft used for repair. Choose all that apply.

### Selections:

- Dacron
- PTFE
- Allograft
- Other
- Graft Not Utilized

### Required:

Yes

### Renal Status (OAAA)

### **Data Abstraction Instructions:**

Indicate renal status and interventions, check all that apply.

### Selections:

- Yes
  - Patent, No Intervention
  - Chronically Occluded
  - Purposely Occluded
  - De-Branch/Bypass
  - Stent
  - Chimney
  - Fenestrated/scallop
  - o Side Branch From Graft
  - Accessory Renal Artery Covered
  - No o
  - Not documented

### Supporting Definitions:

- Patent, No Intervention: No intervention required to maintain renal artery patency
- Chronically Occluded: Renal artery was chronically occluded prior to procedure
- Purposely Occluded: Renal artery was intentionally occluded during the procedure
- . De-Branch/Bypass: Additional intra-operative surgery to bypass renal or visceral vessels
- Stent Only: Renal stent placed to maintain renal artery patency
- Chimney: Placement of a bare metal/covered stent that maintains renal artery patency where the graft occludes the orifice of the renal(s)
- . Fenestrated/scallop: Hole or orifice in the graft to maintain renal or visceral vessel patency
- Side Branch From Graft: Custom made grafts with an additional smaller graft off the main body to maintain renal or visceral vessel patency
- Accessory renal covered: Multiple arteries to the kidney are present and one (or more) is covered by the graft

#### https://bmc2.org/print/book/export/html/78

Additional covered stents can be (and usually are) added to the fenestrated grafts at the scalloped and side branches to maintain renal patency. The side arm branches may or may not reach beyond the aneurysm by themselves and a covered graft may bridge between the main body and renal artery.

#### **Required:**

Yes

#### Anastomotic Felt Reinforcement (OAAA)

### **Data Abstraction Instructions:**

Indicate if Felt Reinforcement was used at the anastomosis site.

#### Selections:

- Yes
- No

**Required:** 

# No

### Hypogastric ligated/occluded (OAAA)

### **Data Abstraction Instructions:**

Indicate if the hypogastric artery was ligated or occluded during the procedure.

#### Selections:



Required:

Yes

### **Proximal Clamp Position (OAAA)**

### **Data Abstraction Instructions:**

Indicate the position of the proximal clamp during the repair.

### Selections:

- Infrarenal
- Above 1 renal
- Above both renal
- Supraceliac
- Clamp not utilized

**Required:** 

Yes

### Inferior Mesenteric Artery at Completion (OAAA)

### **Data Abstraction Instructions:**

Indicate the status of the IMA at the end of the procedure.

### Selections:

- Occluded
- Ligated
- Re-implanted
- Graft Not Utilized
- Not documented

### Supporting Definitions:

Graft not utilized is not a substitution for "not documented".

The IMA may be occluded by thrombus prior to the start of the procedure (dictated no flow/not visualized on pre-procedure imaging).

The IMA may be sutured/ligated during the procedure due to back-bleeding from the artery.

If the IMA appears to be a major source of flow to the bowels, it may be re-implanted to prevent bowel ischemia.

#### Required: Yes

### **Renal/Visceral Ischemic Time (OAAA)**

### **Data Abstraction Instructions:**

Indicate time of renal/visceral clamp time. Include any aortic reclamp time for hypotension.

#### Selections:

Documented Enter value in minutes

- Not documented
- Clamp not utilized

### Supporting Definitions:

Infrarenal clamp position does not have a value for renal/visceral ischemic time, document 0.

#### Required: Yes Suffix:

minutes

### Intra Operative Graft Revision (OAAA)

### **Data Abstraction Instructions:**

Indicate if graft revision was performed during the procedure.

#### Selections:

- Yes
- No

### Supporting Definitions:

A revision is a part of the procedure that is unplanned after the initial graft is sewn into place and tested for patency.

### Required:

Yes

### Cold Renal Perfusion (OAAA)

### **Data Abstraction Instructions:**

Indicate if cold renal perfusion was performed during the procedure.

### Selections:

- Yes
- No

### Supporting Definitions:

The infusion of cold crystalloids into the renal artery during the AAA repair to prevent post procedure renal failure.

### Required:

Yes

### Mannitol administered during procedure (OAAA)

### **Data Abstraction Instructions:**

Indicate if Mannitol was administered during procedure.

### Selections:

Yes

No

Required:

Yes

### **Closure for Open Exposure (OAAA)**

#### **Data Abstraction Instructions:**

Identify all that apply for incision closure. Include all layers of closure - muscle, subcutaneous, and skin.

### Selections:

- Suture
  - Absorbable
     Permanent
- Staples
- Delayed
- Other

### Required:

Yes

### Endovascular Aneurysm Repair (EVAR)

### **Qualifying Procedures:**

- Endovascular Infrarenal, Juxtarenal, and Suprarenal AAA repair Endovascular Ruptured AAA repair. Even if the patient expires after the sheath was inserted
- · An EVAR was done for the indication of iliac artery aneurysm repair, AND a main body endograft was implanted into the aorta in addition to the iliac endograft

- An EVAR that was converted to an OAAA during the same OR time
- A qualifying EVAR where the sheath was inserted; even if the wire or device did not cross the index lesion
- EVAR Revision: Collect if a new main body was implanted
- Patch angioplasty, open endarterectomy, open thromboembolectomy, or pseudoaneurysm repair as a concomitant procedure to an EVAR. You would enter
  these concomitant procedures under the Locations section of the website.

#### EVAR procedures that do not qualify:

- · An endovascular aneurysm repair in the thoracic, abdominal aorta (above the diaphragm)
- · An EVAR that is done for the indication of chronic iliac occlusive disease, aortic stenosis, or pseudoaneurysm repair
- A qualifying EVAR where the sheath was unable to be inserted EVAR Revision: a limb extension or other endovascular device was added to a main body
  endograft

### Prior Family History of AAA (EVAR)

### **Data Abstraction Instructions:**

Indicate if relative (parent, sibling, aunt, uncle, child) has/had a history of AAA.

#### Selections:

- Yes
- No

### **Required:**

Yes

### Prior Aortic Surgery (EVAR)

#### **Data Abstraction Instructions:**

Enter the year, and type of repair, of the prior procedure (yyyy).

#### Selections:

- Yes
  - Enter year
    - AAA (Infrarenal)SAAA (Suprarenal)
    - Bypass
    - Other (Endarterectomy or Other)

## No

Required: No Minimum: 1900 Maximum: 9999

### Maximum AAA Diameter (EVAR)

#### **Data Abstraction Instructions:**

Enter AAA largest diameter in millimeters.

#### Selections:

- Documented
  - Enter value in mm
- Not documented

### Supporting Definitions:

Use largest diameter. If multiple imaging modality, use most accurate in following hierarchy: CT > MRI > ECHO > Arteriogram. This can also be taken from the EVAR planning sheet.

For AAA repair performed due only to iliac aneurysm size (no abdominal aortic aneurysm), enter Not Documented.

Required:		
Yes		
Suffix:		
mm		
Minimum:		
25		
Maximum:		
200		

### Iliac Aneurysm (EVAR)

#### **Data Abstraction Instructions:**

Indicate if the patient had an iliac aneurysm. Choose if Unilateral, Bilateral or None. If yes, enter value for maximum diameter in millimeters.

### Selections:

Yes

 Unilateral
 Bilateral
 Enter value in mm

No

### Supporting Definitions:

Iliac diameter > 1.5 cm. Maximum diameter of largest iliac artery.

Required: Yes Suffix: mm Minimum: 10 Maximum: 100

### Aneurysm location (EVAR)

### **Data Abstraction Instructions:**

Enter the aneurysm location as described in the pre-imaging or surgical dictation.

### Selections:

- Infrarenal
- Juxtarenal
- Suprarenal
- Not documented

Required:

Yes

### Aneurysm anatomy (EVAR)

### **Data Abstraction Instructions:**

Enter the aneurysm anatomy as described in the pre-imaging or surgical dictation.

### Selections:

- Fusiform
- Saccular
- Not documented

### Required:

Yes

### Contained rupture (EVAR)

### Data Abstraction Instructions:

Indicate if the aneurysm was described as a contained rupture on pre-imaging or in the physician dictation.

### Required:

Yes

### Infrarenal Neck Diameter (EVAR)

### **Data Abstraction Instructions:**

Enter neck diameter in millimeters as determined by CTA or IVUS.

### Selections:

- Yes
   Enter value in mm
- No

### Supporting Definitions:

This measurement will come from a CTA scan, may be documented in radiology or operative note. This can also be taken from the EVAR planning sheet.

Required: Yes Suffix:			
mm Minimum: 5 Maximum: 75			

### Infrarenal Neck Length (EVAR)

### **Data Abstraction Instructions:**

Enter length of infrarenal neck (from renal artery to top of aneurysm) in millimeters as determined by CTA or IVUS.

#### Selections:

- Yes
- Enter value in mmNo

### Supporting Definitions:

This measurement will come from a CTA scan, may be documented in radiology or operative note. This can also be taken from the EVAR planning sheet.

Required: Yes Suffix: mm Minimum: 0 Maximum: 50

### **Ruptured AAA (EVAR)**

### **Data Abstraction Instructions:**

Indicate if the aneurysm was ruptured at the time of the procedure. This includes free ruptures with extravasation of blood/contrast on pre-imaging or in the physician dictation.

#### Selections:

- Yes
- No

### Supporting Definitions:

Contained ruptures do not qualify.

Required: Yes

### Lowest Pre-Intubation Blood Pressure (for ruptured AAA repairs only) (EVAR)

#### **Data Abstraction Instructions:**

Indicate the lowest pre-intubation BP, after arrival in hospital, use systolic pressure.

### Selections:

- Yes
- Enter value in mmHgNot documented

Required: Yes Suffix: mmHg Minimum: 0 Maximum: 200

### Mental Status (for ruptured AAA repairs only) (EVAR)

#### **Data Abstraction Instructions:**

Indicate the patient's mental status on arrival to operating room.

### Selections:

- Yes
  - Normal (alert and oriented)
  - Disoriented to person, place, or time.
  - Unconscious
- Not documented

### Required:

Yes

### Cardiac Arrest (for ruptured AAA repairs only) (EVAR)

### **Data Abstraction Instructions:**

Indicate if the patient was in cardiac arrest on arrival to operating room.

# Yes No

....

Required:

Yes

### Timeframe: Symptoms to Incision (for ruptured AAA repairs only) (EVAR)

### **Data Abstraction Instructions:**

Record the time from the patients first symptoms until the surgical incision in hours.

#### Selections:

- Yes
  - Enter value in hours
  - Not documented

#### Required: Yes Minimum: 0 Maximum: 72

### Timeframe: Admission to Incision (for ruptured AAA repairs only) (EVAR)

#### **Data Abstraction Instructions:**

Record the time from when the patient was admitted to the surgical incision in hours.

#### Selections:

- Yes
- Enter value in hoursNot documented

Required: Yes Minimum: 0 Maximum: 24

### Abdomen Explored (EVAR)

### **Data Abstraction Instructions:**

Indicate if the abdomen was explored by the surgeon to evacuate hematoma, but not to repair rupture.

### Selections:

- Yes
- No

### Supporting Definitions:

Aneurysm was repaired endovascularly, but the abdomen was opened to remove clot prior to leaving the OR.

## Required:

Yes

### Procedure Aborted (EVAR)

### **Data Abstraction Instructions:**

Indicate if the procedure was aborted prior to endograft placement (for any reason).

### Selections:

Yes

No

Required: Yes

### Graft Body Diameter (EVAR)

### **Data Abstraction Instructions:**

Enter the diameter of the main body graft.

To find the main body diameter enter the catalog number in Google and go to the manufacturer's website.

You can also go to the Endovascular Today Device Guide website to look up the distal diameter of the main body. https://evtoday.com/device-guide/us/main-body-grafts

### Selections:

- Documented
  - Enter value in mm
- Not documented
- Graft Not Utilized

### Supporting Definitions:

Body Size is diameter of most proximal portion of graft.

Required: Yes Suffix: mm Minimum: 10 Maximum: 40

### **Right Distal Seal Zone Diameter**

#### **Data Abstraction Instructions:**

Enter the diameter of the most distal portion of iliac treated with any covered stent such as iliac limbs, iliac extensions, contralateral limbs, ipsilateral limbs and bridge devices. Please note: a bridging device is essentially a flared iliac limb.

To find the distal seal zone enter the catalog number in Google and go to the manufacturer's website. Depending on the manufacturer, the distal seal zone diameter may be labeled as: iliac endoprosthesis diameter, distal device diameter, iliac limb diameter or iliac leg diameter.

You can also go to the Endovascular Today Device Guide website to look up the distal diameter of a device. For iliac leg extensions, go to: <u>https://evtoday.com/device-guide/us/search?q=extension</u> For iliac limbs, go to: <u>https://evtoday.com/device-guide/us/iliac-limbs</u>

#### Selections:

- Yes
- Enter diameter \_\_\_\_mm

### No

### Required:

Yes

### Left Distal Seal Zone Diameter

#### **Data Abstraction Instructions:**

Enter the diameter of the most distal portion of iliac treated with any covered stent such as iliac limbs, iliac extensions, contralateral limbs, ipsilateral limbs and bridge devices. Please note: a bridging device is essentially a flared iliac limb.

To find the distal seal zone enter the catalog number in Google and go to the manufacturer's website. Depending on the manufacturer, the distal seal zone diameter may be labeled as: iliac endoprosthesis diameter, distal device diameter, iliac limb diameter or iliac leg diameter.

You can also go to the Endovascular Today Device Guide website to look up the distal diameter of a device. For iliac leg extensions, go to: <u>https://evtoday.com/device-guide/us/search?q=extension</u> For iliac limbs, go to: <u>https://evtoday.com/device-guide/us/iliac-limbs</u>

#### Selections:

- Yes
- Enter diameter \_\_\_\_mm
  No

**Required:** 

Yes

### Graft Type (EVAR)

### Data Abstraction Instructions:

Indicate type of graft used.

- Aneurx
- Excluder
  Talent
- Zenith
- PowerlinkEndurant

- UnifitLow Profile
- Aptus
- AFX

- Endologix
- Nellix
- Ovation TrivascularOther
- Graft Not Utilized

-----

Required: Yes

### **Graft Configuration**

### Data Abstraction Instructions:

Indicate graft configuration.

### Selections:

- Aorto-bi-iliac
- Aorto-uni-iliac RT
- Aorto-uni-iliac LT
- Aorto-aortic
- Fenestrated
- Graft Not Utilized

Required: Yes

### Additional graft components (EVAR)

#### **Data Abstraction Instructions:**

Indicate if additional graft components were implanted other than iliac limbs, iliac extensions, stents or bridging devices. When selecting the option Iliac Branch Device, two additional fields will display. Enter the distal hypogastric (internal iliac) diameter and the distal external iliac diameter. This field is a multi-select field.

- An aortic cuff is an extension device used to provide a seal at the proximal aortic neck.
- Aortic screws are implanted through the aorta and fix the endograft to the aorta. Aortic screws stop the endograft from migrating. May also be call endo anchors.
   An Ilica Bread Device is a biturgated devices that is implanted in the every and they internal ilica arter. (hypersection) An additional ecomponent is
- An Iliac Brach Device is a bifurcated devices that is implanted in the external iliac artery and they internal iliac artery (hypogastric). An additional component is implanted into the hypogastric.

### Selections:

- Aortic cuff
- · Aortic screws
- · Right iliac branch device
  - Distal hypogastric diameter
    - Documented
    - Enter value (mm)
    - Not Documented
  - Distal external iliac diameter
    - Documented
      - Enter value (mm)
  - Not Documented
- Left iliac branch device
  - Distal hypogastric diameter
    - Documented
    - Enter value (mm)Not Documented
  - Distal external iliac diameter
    - Documented
    - Enter value (mm)
    - Not Documented
- Additional main body
- Other

Required:

Yes

### Renal Status

### **Data Abstraction Instructions:**

Indicate renal status and interventions, check all that apply.

- Yes
  - Patent, No Intervention
  - Chronically Occluded
  - Purposely Occluded
  - o De-Branch/Bypass
  - Stent
  - Chimney
  - Fenestrated/scallop
  - Side Branch From Graft
  - Accessory Renal Artery Covered
- No

Not documented

### Supporting Definitions:

- · Patent, No Intervention: No intervention required to maintain renal artery patency
- Chronically Occluded: Renal artery was chronically occluded prior to procedure
- Purposely Occluded: Renal artery was intentionally occluded during the procedure
- De-Branch/Bypass: Additional intra-operative surgery to bypass renal or visceral vessels
- Stent Only: Renal stent placed to maintain renal artery patency
- Chimney: Placement of a bare metal/covered stent that maintains renal artery patency where the graft occludes the orifice of the renal(s)
- Fenestrated/scallop: Hole or orifice in the graft to maintain renal or visceral vessel patency
- · Side Branch From Graft: Custom made grafts with an additional smaller graft off the main body to maintain renal or visceral vessel patency
- Accessory Renal Artery Covered: Multiple arteries to the kidney are present and one (or more) is covered by the graft

Additional covered stents can be (and usually are) added to the fenestrated grafts at the scalloped and side branches to maintain renal patency. The side arm branches may or may not reach beyond the aneurysm by themselves and a covered graft may bridge between the main body and renal artery.

Required: Yes

### Hypogastric Coiled/Plugged

```
Data Abstraction Instructions:
```

Indicate if the patient had coiling of the hypogastric arteries, either pre procedure or during the procedure.

### Selections:

- Yes
  - Coiled/Plugged Pre-op
  - Coiled/Plugged Intra-op
    - UnilateralBilateral
- No

Required:

Yes

### Hypogastric Intentionally Covered

#### **Data Abstraction Instructions:**

Indicate if the Hypogastric was intentionally covered and planned prior to procedure to treat distal aneurysm extent.

#### Selections:

- Yes
  - Unilateral
  - Bilateral
- No
- Graft Not Utilized

**Required:** 

Yes

### Hypogastric Unintentionally Covered

#### **Data Abstraction Instructions:**

Indicate if the Hypogastric was inadvertently covered with extension of graft not necessary to treat distal aneurysm extent.

#### Selections:

- Yes
   Unilateral
  - Bilateral
- No
- Graft Not Utilized

Required:

Yes

### Arterial Injury

#### **Data Abstraction Instructions:**

Indicate if an arterial injury or occlusion occurred, requiring an intervention, and document what intervention was performed.

	Yes		
•			Select option for injury
		0	Femoral
		-	lliac
			Renal
			Aorta
			-
			Aorta

- Multiple
- · Select option for repair
  - Stent/PTA
  - Stent/Graft
  - Open Repair Not documented

No

### Required:

Yes

### Intra-Operative Revision Needed

### **Data Abstraction Instructions:**

Indicate if a graft revision was performed during the procedure.

### Selections:

- Yes
- No

### Supporting Definitions:

A revision may include an unplanned limb extension, the addition of a proximal cuff, or a main body extension.

## **Required:**

Yes

### **Endoleak at Completion**

### **Data Abstraction Instructions:**

Indicate if there was an endoleak at the end of the procedure.

### Selections:

- Yes
  - Attachment site (Type 1) (Proximal or distal attachment site leak)
  - Branch (Type II) (Retrograde filling of sac via lumbars, IMA or accessory renals) 0
  - · Mid Graft (Type III) (filling of sac via leak at component overlap sites or fabric tear)
  - Indeterminate
- No

### Supporting Definitions:

If an endoleak is identified and then repaired and is no longer present at the end of the procedure, mark no.

#### **Required:**

Yes

### **Closure for Groin Access**

### Data Abstraction Instructions:

Identify all that apply for closure. Include all layers of closure - muscle, subcutaneous, and skin.

### Selections:

- percutaneous
  - manual
  - perclose ٥
  - angioseal
  - mynx
  - starclose
  - Exoseal ٥ compression device
- open
  - suture
    - absorbable permanent •
    - . Not documented
  - staples
  - skin glue
  - other

### **Required:**

Yes

### **Additional Planned Procedures**

### **Data Abstraction Instructions:**

Indicate if additional procedures were performed.

- Yes
  - Femoral Endarterectomy
  - Thromboembolectomy
  - Other Arterial Reconstruction

### Required:

No

Yes

### **Conversion to Open**

### **Data Abstraction Instructions:**

Indicate if the procedure had to be converted to an open procedure and identify reason.

#### Selections:

- Yes
  - · Unable to deploy appropriately
- No

### Supporting Definitions:

If the EVAR is converted to an open AAA repair, the open AAA must be entered as an additional procedure within the discharge record.

#### Required: Yes

### **Open Thrombectomy (OT)**

Endoleak

#### **Qualifying Procedures:**

- · Open Thrombectomy of the upper and lower extremities
- Urgent or emergent case
- Indication of Acute Limb Ischemia (ALI)
- · The surgeon made an arteriotomy (an incision) into the artery or graft
- Thrombus was removed manually or with a Fogarty
- The procedure might be documented as a Thrombectomy or an embolectomy if the clot traveled to that location
- Independent case
- The Open Thrombectomy was not performed at the same time as another VS or PVI procedure
- A qualifying open Thrombectomy where the primary incision was made, but the procedure was not completed
- Patch angioplasty, aneurysm, or pseudoaneurysm repair as a concomitant procedure to an open thrombectomy. You would enter these concomitant procedures
  under the Locations section of the website

#### Open Thrombectomy procedures that do not qualify:

- Open Thrombectomy with the Indication of Critical Limb Ischemia (CLI)
- An open Thrombectomy that was an outcome of another procedure within the same hospitalization or discharge. Enter this open Thrombectomy as an outcome
  of the prior procedure (if applicable)
- If the open Thrombectomy is done at the insertion or origin site of an open bypass, you will not enter this as an open Thrombectomy case. It would be considered part of a typical open bypass procedure and preparing the artery for anastomosis and would not be entered in Locations
- For an open Thrombectomy that was performed at the same time as another VS or PVI procedure and in a different location as the VS or PVI procedure, you
- would enter this open Thrombectomy in Locations of the concomitant VS or PVI procedure as an additional procedure performed
- A qualifying open thrombectomy where the procedure was aborted BEFORE the primary incision was made.

### Timeframe: Symptoms to incision (OT)

#### **Data Abstraction Instructions:**

Record the time from symptom onset until surgical incision.

#### Selections:

- Enter value (hours)
- Not Documented

### Required:

Yes

### Timeframe: Presentation to incision (OT)

#### **Data Abstraction Instructions:**

Record the time from arrival at the hospital until surgical incision.

### Selections:

- Enter value (hours)
- Not documented

Required: Yes

### Prior lytic procedure (OT)

### **Data Abstraction Instructions:**

Indicate if the patient had a prior lytic procedure during the current hospitalization.

#### Selections:

- Yes
- No

### **Required:**

Yes

### Vessel location (OT)

#### **Data Abstraction Instructions:**

Indicate vessel location of the procedure. If an open thrombectomy is perfomed on multiple vessels in the same surgery, enter the location of the incision.

#### Selections:

Choose from drop down.

Required: Yes

### Vessel closure (OT)

#### **Data Abstraction Instructions:**

Indicate how the vessel at the site of thrombectomy was closed.

#### Selections:

- Primary
- PatchNot documented
- Not document

### Required:

Yes

### Completion angio (OT)

### **Data Abstraction Instructions:**

Indicate if a completion angio was performed at the end of the procedure and if flow through the thrombectomized vessel was normal or abnormal.

### Selections:

No

- Abnormal

# Required:

Yes

### Concomitant endarterectomy (OT)

### **Data Abstraction Instructions:**

Indicate if an endarterectomy was performed at the same time as the thrombectomy.

### Selections:

- Yes
- No

Required: Yes

### **Data Fields for All Procedure Types**

### Anesthesia Type

#### **Data Abstraction Instructions:**

Indicate the type of anesthesia used.

- Local
- · Epidural
- Regional
- Spinal

- General
- Epidural + General

MAC

**Required:** Yes

### **Antibiotics Pre Procedure**

#### **Data Abstraction Instructions:**

Indicate if an antibiotic was given within one hour of incision and whether it was redosed in the given timeframe.

### Selections:

٠	Yes		
		0	Cefazolin
			<ul> <li>Redosed (Q4 hours)</li> </ul>
			Yes
			No
		0	Clindamycin
			<ul> <li>Redosed (Q6 hours)</li> </ul>
			Yes
			No
		0	On scheduled antibiotic
		٥	Other
	No		

### Supporting Definitions:

No

If Vancomycin is the antibiotic used, it can be administered 2 hours before incision.

On scheduled antibiotic = An inpatient did not receive the pre procedure antibiotic within 1 hour of incision time (2 hours for Vancomycin) because the patient is on a scheduled antibiotic.

#### Required:

Yes

### **Skin Preparation**

### **Data Abstraction Instructions:**

Indicate all skin prep items used.

### Selections:

- Chlorhexidine
- Alcohol
- Iodine
- Chlorhexidine + Iodine
- Chlorhexidine + Alcohol
- Iodine + Alcohol

### **Required:**

Yes

### **Contraindicated to Chlorhexidine & Alcohol Skin Preparation**

### **Data Abstraction Instructions:**

Indicate if there is a contraindication to Chlorhexidine & Alcohol skin prep (for example: allergy, open wound, etc.).

### Selections:

- Yes
- No

**Required:** 

### Yes

### Glucose (peak)

### **Data Abstraction Instructions:**

Indicate the highest intra operative blood glucose.

### Selections:

- Yes
  - Enter value in mg/dL
- Not documented

Required: Yes Suffix:

mg/dL

### Nadir body temperature

### **Data Abstraction Instructions:**

Enter the lowest body temperature that is taken after the procedure start time and before the procedure end time.

### Selections:

- Yes
- Enter value in Celsius · Not documented

Required: Yes Suffix: Celsius Minimum: 30 Maximum: 40

### Crystalloids

#### **Data Abstraction Instructions:**

Indicate the total volume of intravenous crystalloids and/or Plasma-lyte infused intra operatively. Also, document Plasma-lyte as Other IV Hydration under Meds Given During Procedure.

### Selections:

- Yes
  - Enter value in ml
- Not documented

### Supporting Definitions:

Examples of crystalloids include Normal Saline, .45 Normal Saline, D5W, and Lactated Ringer's.

**Required:** Yes Suffix: ml Maximum: 20000 Soft Maximum: 10000

### Estimated Blood Loss (EBL)

#### **Data Abstraction Instructions:**

Indicate the estimated blood loss during the procedure.

### Selections:

- Yes
- Enter EBL in mL · Not documented

**Required:** Yes Suffix: ml Maximum: 5000 Soft Maximum:

### ASA (American Society of Anesthesiologists) Class

### **Data Abstraction Instructions:**

Choose class determination as determined by anesthesia.

### Selections:

1000

- Does not apply
- Class 1 normal/healthy
- Class 1 normalinearity
   Class 2 mild systemic disease Class 3 – severe systemic disease
- Class 4 severe systemic disease that is a constant threat to life
- Class 5 moribund/not expected to survive without operation

### **Supporting Definitions:**

Choose "Does not apply" only if anesthesia is not involved in the patients care during the procedure.

### Required:

### Yes

### **Contrast Types**

### **Data Abstraction Instructions:**

Indicate all types of contrast that was used. If more than one contrast were used, choose all that apply.

#### Selections:

- Nonionic, low-osmolar
- Nonionic, Iso-osmolar
- lonic, hyperosmolar lonic, low-osmolar
- Unknown/Investigational contrast agent
- Gadolinium
- Carbon Dioxide (CO2)
- None

### Supporting Definitions:

#### **Commonly used Contrast Agents**

- Nonionic low-osmolar
- Omnipaque, Isovue, Optiray, Ultravist, Oxilam
- Nonionic Iso-osmolar
- VisipaqueIonic, hyperosmolar
- Hypaque, Conray
- Ionic, low-osmolar
- Hexabrix
- Unknow/Investigational contrast agent
- Gadolinium
- Carbon Dioxide (CO2)

### Required:

#### Yes

### **Total IV Contrast Used**

### **Data Abstraction Instructions:**

Indicate the volume of contrast (ionic & non-ionic) used in milliliters (ml). The volume recorded should be the total volume for the lab visit. This should be the total between the start of procedure and end of procedure.

### Selections:

- Yes
  - Enter value in ml
- Not documented

#### Supporting Definitions:

If half dose contrast was used during the procedure, record only the dose of the contrast given, not the total volume. If CO2 contrast is used, do not include the volume of CO2 used in the total contrast.

**Required:** 

Yes Suffix: ml Minimum: 0 Maximum: 500

### Heparin Administered

### **Data Abstraction Instructions:**

Record if heparin was given. If yes, record the total dose(s)/bolus(es) of unfractionated heparin units that were given during the procedure. If heparin was given and you cannot find documentation of the dose given choose Not documented.

### Selections:

Heparin Administered

- Yes
  - Total Heparin Dosage
  - Documented
     Enter value in units
  - Not documented
  - 0

Do not include heparin drip doses in this value. Include only the bolus doses.

Required: Yes Suffix: units Maximum: 40000

### Peak Intra Procedure Activated Clotting Time (ACT)

#### **Data Abstraction Instructions:**

Indicate the peak intraoperative ACT in seconds.

### Selections:

- Yes
- Enter value in seconds
- Not documented

#### Supporting Definitions:

Activated clotting time (ACT) should be measured after the heparin IV bolus is given. In long cases, as clinically indicated, additional heparin boluses may be given and subsequent ACT measurements may be done. The ACT recorded here must be done during, NOT at the end of the procedure. There must be some part of the intervention procedure performed after the ACT value for it to qualify for peak ACT. Record highest measurement of ACT (peak) in seconds. Enter "Not documented" if peak ACT or clotting measurement was not drawn/document in the patient record.

#### **Required:**

Yes Suffix: seconds Maximum: 600

### End of procedure ACT

### **Data Abstraction Instructions:**

Record the activated clotting time (ACT) at the conclusion of the procedure, during closure or before the sheath is changed out at the end of the case. In the OR cases, you can enter the ACT that is drawn approximately an hour from the end of the procedure.

### Selections:

- Yes
  - Enter value in seconds
- Not documented

Required: Yes Suffix: seconds Minimum: 50 Maximum: 600

### Locations

Each procedure type can have one or many locations. In the case of a PVI procedure, there will always be locations. For a vascular surgery procedure, the locations field should be available only in the case of an "Open Lower Extremity Bypass" where an option is selected in the "Vascular Surgery Procedure Performed" field.

### Vessel Location

### **Data Abstraction Instructions:**

Indicate vessel location of the procedure.

### Selections:

Choose Vessel Location from the drop down list

Required: Yes

### ....

### Lesion Segment Area

- Proximal
- Mid
- Distal
- DiffuseNot documented

### Supporting Definitions:

Identify if the lesion is proximal, mid, distal, or diffuse. If the lesion treated involves more than one segment, check diffuse (e.g. proximal and mid).

#### Required: Yes

### **PVI Procedure Performed**

### **Data Abstraction Instructions:**

Indicate procedure performed. Select all that apply.

Selections:

- Aspirational Atherectomy (JetStream, Pathways) = Asp-Ather
- Mechanical Thrombectomy (Angiojet) M-Throm
- Balloon = BA
- Cryoballoon = Cryo-B
- CTO device = CTO
- Cutting Balloon = CB
- Directional Atherectomy (Fox hollow, SilverHawk) = D-Ather
- Distal Protection Device (balloon) = DPD-B
- Distal Protection Device (filter) = DPD-F
- Drug Coated Balloon = DCB
- Flow-wire = FW
- Infusion Catheter (Benephit) = Inf-Cath
- Intravascular Ultrasound = IVUS
- Laser Atherectomy (Excimer laser) = L-Ather
- Lysis (Note: do not record lysis only procedures). Select this box if lytic agents were used during the procedure in addition to any other device. Do not record
  procedures if only angiojet or fogarty catheter was used.) = LYS
- Not crossed with a device = ND
- Not crossed with a wire = NW
- Other Atherectomy (ClearPath) = Oth-Ather
- Open Endarterectomy
- Open Thrombectomy
- Rotational/Orbital Atherectomy (DiamondBack) = R-Ather
- Re-Entry Catheter (Pioneer, Outback) = Re-Ent-Cath
- Research (whether the procedure was done for research purpose only) = Research
- Scoring Balloon (Angiosculpt) = S-BA
- Stent = STNT
- Thrombus Aspiration (Pronto, Export, Aspire, Diver, Xtract, Fetch, QuickCat) Throm-Asp
- Vascular Embolectomy = Vasc-E (Fogarty)

### Required:

Yes

### **Bypass Graft**

### **Data Abstraction Instructions:**

Indicate if the procedure is performed on an arterial bypass graft.

#### Selections:

- Yes
- No

Required:

Yes

### Graft Type

#### **Data Abstraction Instructions:**

Select the type of bypass graft: synthetic or vein.

### Selections:

- Synthetic
- Vein
- Not Documented

### Required:

### Yes

### **Graft Origin**

#### **Data Abstraction Instructions:**

Select the bypass graft origin using the vessel drop down box.

Select artery name from the drop down list

### **Required:**

Yes

### **Graft Insertion**

### **Data Abstraction Instructions:**

Select the bypass graft insertion using the vessel drop down box.

#### Selections:

· Select artery name from the drop down list

Required:

Yes

### Lesion Length

### **Data Abstraction Instructions:**

Visual estimate of the length of the lesion. If not dictated, use balloon/stent length. For tandem lesions, add lengths together. For diffuse disease use the length of the treated segment.

### Selections:

Enter value in mm

### Supporting Definitions:

In completing this field, first use a dictated value. If no value is dictated, select stent from the PVI Procedure Performed section. If multiple stents are used, add the lengths. If no stent, then use balloon size.

Required: No Suffix: mm Minimum: 0 Maximum: 1000

### Heavy Calcium

### **Data Abstraction Instructions:**

Indicate if moderate to heavy calcium is documented as being present in the lesion.

#### Selections:

- Yes
- No

#### Required: Yes

### **In-stent Restenosis**

### **Data Abstraction Instructions:**

Select Yes if the lesion that is being treated is within a previously place stent. Select diffuse (greater than 1/3 the length of the stent) or focal.

### Selections:

- Yes
- No

### Required:

Yes

### Thrombus

### Data Abstraction Instructions:

Thrombus is suggested by certain angiographic features: haziness, reduced contrast density or contrast persistence, irregular lesion contours or globular filling defects. Mark "yes" if present (before intervention)

### Selections:

- Yes
- No

Required: Yes

### Pre Stenosis % (0-100)

### Data Abstraction Instructions:

For each segment treated, record the preprocedural percent of stenosis. If a range is given, take the highest value. If unavailable, choose not documented.

### Selections:

Yes

- Enter value (0 100)
- Not documented

Required:

Yes Suffix: % Maximum: 100

### Post Stenosis % (0-100)

### **Data Abstraction Instructions:**

For each segment treated, record the postprocedural percent stenosis; if range is given, take the lowest value. If not recorded, choose not documented.

### Selections:

- Yes
- Enter value (0 100)Not documented

Required: Yes Suffix: % Maximum: 100

### **Final Balloon Diameter**

### **Data Abstraction Instructions:**

Indicate the diameter, in millimeters, of the final balloon used to treat this lesion. If not known, mark not documented.

### Selections:

- Yes
- Enter value in mmNot documented

Required:

Yes Suffix: millimeters Minimum: 1.5 Maximum: 30

### Stents

### Stent Name

### **Data Abstraction Instructions:**

Select the name of the stent used. If the name of the stent is not included in this list, please choose other.

### Selections:

Choose Stent name

Required:

Yes

### Stent Diameter

### **Data Abstraction Instructions:**

Enter the diameter of the stent.

### Selections:

• Enter Stent Diameter in mm

**Required:** 

Yes Suffix: mm Minimum: 2 Maximum: 30

### Stent Length

### **Data Abstraction Instructions:**

Enter the length of the stent.

### Selections:

Enter Stent Length in mm

Required:
Yes
Suffix:
mm
Minimum:
1
Maximum:
250

### Vascular Access

### Vascular Access Site(s)

### **Data Abstraction Instructions:**

Indicate location of vascular access.

#### Selections:

· Select artery from the drop down list

Required: Yes

### Vascular Access Type

### **Data Abstraction Instructions:**

Indicate vascular access type.

### Selections:

- Percutaneous
- Surgical Cutdown

### Supporting Definitions:

Percutaneous is vascular access obtained via skin puncture without direct visualization of artery. Surgical cutdown is access via skin incision with direct visualization

### Required:

Yes

### Vessel Accessed

### **Data Abstraction Instructions:**

Indicate if the native artery or bypass graft was accessed for the current procedure.

### Selections:

- Native Artery
- Bypass Graft

#### Required: Yes

### **Access Guidance**

### **Data Abstraction Instructions:**

Indicate if guidance was used for vascular access. If both are utilized, select ultrasound.

- Yes
   Fluoro
- https://bmc2.org/print/book/export/html/78

# UltrasoundNo

### Required:

Yes

### Access Approach

#### **Data Abstraction Instructions:**

Record if access approach was antegrade or retrograde.

#### Selections:

- Antegrade
- Retrograde
- Both

### Supporting Definitions:

This indicates sheath direction at site of insertion. If more than one access was attempted, record the access approach that was used to gain access rather than the failed access approach. If a sheath was utilized in both the retrograde and antegrade direction at the same insertion site, document "both".

#### Required:

Yes

### Sheath Size

### **Data Abstraction Instructions:**

Indicate the largest size of the sheath placed during the procedure.

#### Selections:

· Enter value (French)

#### Supporting Definitions:

Include sheaths placed at the end of the procedure.

Required: Yes Suffix: French Minimum: 3 Maximum: 30

### Sheath Removed

#### **Data Abstraction Instructions:**

Indicate if the sheath was removed by the physician, nurse, technician, or advanced practice professional (NP or PA). In cases of manual removal, indicate the person responsible for holding pressure.

In lysis procedures in which the sheath is left in at the end of the procedure, select yes for "Sheath Removed" and indicate the timeframe/method of eventual removal.

#### Selections:

- Yes
- No

#### Supporting Definitions:

Answering "Yes" to this field triggers the 3 following conditional fields: Vascular Closure Type, Failed Closure, and Sheath Removal Time Post Procedure.

**Required:** 

Yes

### Vascular Closure Type

#### Data Abstraction Instructions:

Indicate the arterial closure methods used regardless of whether or not they provided hemostasis. Note: If more than one vascular closure type per access site was used, select all that were used.

- Manual: no device or a mechanical type was used, e.g. manual pressure by the personnel pulling the sheath.
- Perclose
- Angioseal
- Mynx
- Starclose
- SurgicalExoseal

- https://bmc2.org/print/book/export/html/78
- Compression Device (ie: Femstop, C Clamp, TR Band)
- Boomerang
- Hemostatic PatchFISH
- Vascade

### Supporting Definitions:

Conditional on "Yes" indicated for Sheath Removed

Required: Yes

### **Sheath Removal Time**

### **Data Abstraction Instructions:**

Indicate time between end of procedure and sheath removal.

#### Selections:

- 0-3 hours
- 3-24 hours
- >24 hours

#### Supporting Definitions:

Conditional on "Yes" indicated for Sheath Removed

**Required:** 

Yes

### **Outcomes During Procedure**

### Death/Cause (ODP)

#### **Data Abstraction Instructions:**

Indicate if the patient died in association with this procedure while in the lab/OR. If yes, indicate cause of death.

#### Selections:

- Yes
  - · Cardiovascular (includes: AMI, bleed, stroke, cardiogenic shock)
  - Hemorrhage
  - Multisystem Organ Failure (includes acute lung injury and systematic inflammatory response system)
  - Other (include neurologic, renal, liver, GI, cancer)
  - Unknown cause of death
- No

### Supporting Definitions:

Select no if the patient was alive throughout the procedure.

Required: Yes

### **Dissection (Not Repaired)**

### **Data Abstraction Instructions:**

Indicate if there was a dissection that was clinically significant (causing a decrease in blood flow) or residual blood flow limiting dissection at the intervention/procedure site.

#### Selections:

- Yes
- No

#### Supporting Definitions:

The appearance of contrast material outside the expected luminal dimensions of the target vessel and extending longitudinally beyond the length of the lesion. If the dissection is successfully treated with, e.g., angioplasty or stent, then it should not be considered as a complication. It would be considered a complication if the patient has a dissection identified during a return trip to the lab.

### Required:

Yes

### Myocardial Infarction (MI) (ODP)

### **Data Abstraction Instructions:**

Indicate if the patient had an MI during the vascular procedure, while the patient was still in the lab or operating room.

- Yes
- No

### Supporting Definitions:

#### A myocardial infarction is evidenced by any of the following:

- 1. A rise and fall of cardiac biomarkers (preferably troponin) with at least one of the values in the abnormal range for that laboratory [typically above the 99<sup>th</sup>
  - percentile of the upper reference limit (URL) for normal subjects] together with at least one of the following manifestations of myocardial ischemia:
    - Ischemic symptoms
    - ECG changes indicative of new ischemia (new ST-T changes, new left bundle branch block, or loss of R wave voltage).
    - Development of pathological Q waves in 2 or more contiguous leads in the ECG (or equivalent finding for true posterior MI.
    - Imaging evidence of new loss of viable myocardium or new regional wall motion abnormality.
    - Documentation in the medical record of the diagnosis of acute myocardial infarction based on the cardiac biomarker pattern in the absence of any items enumerated in a-d due to conditions that may mask their appearance (eg., peri-operative infarct when the patient cannot report ischemic symptoms; baseline left bundle branch block or ventricular pacing).
- 2. ECG changes associated with prior myocardial infarction can include the following (with or without prior symptoms):
  - Any Q wave in leads V2-V3 >= 0.02 seconds or QS complex in leads V2 and V3.
    - Q-wave >= 0.03 seconds and >= 0.1 mV deep or QS complex in leads I, II, aVL, aVF, or V4-V6 in any two leads of a contiguous lead grouping (I, aVL, V6; V4-V6; I, III, and aVF).
  - R-wave >= 0.04 seconds in V1-V2 and R/S >= 1 with a concordant positive T-wave in the absence of a conduction defect.
- 3. Imaging evidence of a region with new loss of viable myocardium at rest in the absence of a non-ischemic cause. This can manifest as:
  - Echocardiographic, CT, MR, ventriculographic or nuclear imaging evidence of left ventricular thinning or scarring and a failure to contact appropriately (i.e., hypokinesis, akinesis, or dyskinesis).
  - Fixed (non-reversible) perfusion defects on nuclear radioisotope imaging (e.g., MIBI, thallium).
  - Medical records documentation of prior myocardial infarction.

#### Source:

Joint EXC-ACC-AHA=WHF 2007 Task Force consensus document "Universal Definition of Myocardial Infarction".

#### **Required:**

Yes

### **Cardiac Arrest**

#### **Data Abstraction Instructions:**

Indicate if the patient was in cardiac arrest during the procedure.

#### Selections:

- Yes
- No

#### Supporting Definitions:

Do not indicate if the patient came into the lab in cardiac arrest, as an example ruptured AAA arresting on arrival.

### Required:

Yes

### Embolus (ODP)

#### Data Abstraction Instructions:

Indicate if the patient is identified to have an embolus during the procedure. If yes, indicate if it was treated successfully.

### Selections:

- Yes
  - Successful
     Unsuccessful
- Unsuce
   No

### Supporting Definitions:

An embolus (compromised of atherosclerotic debris and / or blood clot) moves through the blood vessels until it reaches a vessel that is too small to let it pass. When this happens, the blood flow is stopped by the **embolus**. This occurs after exiting the procedure area.

### Required:

Yes

### Thrombus (ODP)

#### **Data Abstraction Instructions:**

Indicate if a blood clot formed during the procedure, within the treated vessel, that limits distal flow.

- Yes
- No

### Supporting Definitions:

Do not include any thrombus that was present at the beginning of the procedure.

### Required: Yes

### Stent/Graft Thrombosis

### **Data Abstraction Instructions:**

Indicate if there was a stent/graft thrombosis where blood clot forms within the stent/graft that limits distal blood flow during the procedure. If yes, indicate if it was treated successfully.

### Selections:

```
    Yes

            Successful
            Unsuccessful

    No
```

### Required:

Yes

### **Vessel Perforation**

### **Data Abstraction Instructions:**

Indicate if there was a vessel perforation during the procedure. If yes, indicate if it was treated successfully.

### Selections:

Yes

- Successful
  - Balloon
  - Covered stent
  - Bare metal stent
  - External compression
  - Reversal of anticoagulationNo treatment
- No treatme
   Unsuccessful
- No

### Supporting Definitions:

A perforation occurs when there is angiographic or clinical evidence of a dissection or intimal tear that extends through the full thickness of the arterial wall, distant from the access site caused by device manipulation. Extravasations of contrast beyond vessel wall is usually seen.

### Required:

Yes

### TIA/Stroke (ODP)

### Data Abstraction Instructions:

Indicate if there was abrupt loss of neurological function with complete return of function within 24 hours or loss of neurological function caused by an ischemic event that is severe enough to leave a persistent deficit for greater than 24 hours. The symptoms should begin while the patient is in the procedure area.

#### Selections:

- Yes
- No

Required: Yes

### Transfusion (ODP)

### Data Abstraction Instructions:

Indicate if the patient received any transfusion for any reason during the procedure. If yes, select the type of transfusion: PRBC, Whole blood, Platelets, FFP, Other (e.g. Cryoprecipitate, Factor VIII infusion). Select all that apply.

- Yes
- Select type of transfusion
  - PRBC, if yes, enter the number of units for PRBC's (enter # of packed red blood cells 1, 2, 3, etc.)
  - Platelets
  - ₀ FFP
  - Other
- No

### **Supporting Definitions:**

NOTE: Return of cell saver product is not captured as a transfusion.

Required:	
Yes	
Minimum:	
1	
Maximum:	
20	

### Vascular Access Complications (ODP)

### **Data Abstraction Instructions:**

Indicate vascular complications at the access site requiring transfusion, prolonged hospital stay, causing a drop in hemoglobin 3.0 gm/dl, or any access site complications requiring surgical repair. Select all that apply.

#### Selections:

- Yes
  - Retroperitoneal hematoma
  - Pseudo-aneurysm
  - Hematoma at access site
  - Bleeding at access site
  - AV fistula
  - Acute thrombosis
  - Surgical repair of the vascular access site
     Other
- No

### Supporting Definitions:

### Retroperitoneal hematoma

Pseudoaneurysm. The occurrence of a disruption and dilation of the arterial wall without identification of the arterial wall layers at the site of the catheter entry demonstrated by arteriography or ultrasound.

Hematoma requiring transfusion. Blood loss at the site of arterial or venous access or due to perforation of a traversed artery or vein requiring transfusion and/or prolonging the hospital stay, and/or causing a drop in hemoglobin > 3.0 gm/dl.

Bleeding at access site= Blood loss associated with decreased Hgb (greater than or equal to 3.0 gm/dl) and/or causes an increased length of hospital stay. Without other obvious source (GI, GU, operative, or hemolysis) that is attributable to intraprocedural blood loss (e.g. during equipment changes) should be considered bleeding at the access site even if no hematoma is palpable or documented on imaging studies.

AV fistula. A connection between the access artery and the accompanying vein that is demonstrated by arteriography or ultrasound and most often characterized by a continuous bruit.

Acute thrombosis. Total obstruction of the artery by thrombus most commonly at the site of access.

Surgical repair of the vascular access site (such as surgical closures, exploration of the arteriotomy site, balloon angioplasty or covered stent (JOMED GraftMaster) placement to deal the arterial tear)

**Required:** 

Yes

### Vascular Surgery Emergent (ODP)

### **Data Abstraction Instructions:**

Indicate if the patient needed to go to the operating room immediately from intervention room or conversion to an unplanned open procedure.

### Selections:

- Yes
  - Artery Rupture
  - Access Site Complication
  - Bleeding
  - Bowel Ischemia
  - Limb Ischemia
  - Thrombosis/Embolus
  - Conversion to Open Procedure
    Other
- No

### Supporting Definitions:

The procedure may include any of the following: dissection of artery requiring surgical repair, embolus or thrombosis not manageable by percutaneous devices, ischemic leg in lab requiring surgery, device removal, and repair of vascular access complications. Emergent surgery must be performed to prevent loss of major organ, tissue/limb or life. Do not include staged procedures.

### Required:

Yes

### Amputation (in Lab/OR)

### **Data Abstraction Instructions:**

Indicate if an amputation is performed at any time during the procedure.

### Selections:

Selections:

- Yes
  - Select type of amputation
    - Left hip disarticulation
    - Left AKA
    - Left BKA
    - Left foot
    - · Left metatarsal
    - Left digit
    - Right hip disarticulation
    - Right AKA
    - Right BKA
    - Right foot
    - · Right metatarsal
    - Right digit
  - No

**Required:** 

No

### Compartment Syndrome (in Lab/OR)

### **Data Abstraction Instructions:**

Indicate if the patient was determined to have compartment syndrome at any time during the procedure.

#### Selections:

- Yes
- ٠ No

### **Supporting Definitions:**

This is defined as compression of nerves and blood vessels within an enclosed space which leads to muscle and nerve damage and problems with blood flow. Include fasciotomy for prophylaxis.

### **Required:**

No

### **Outcomes Post Procedure**

All outcomes from the end of the procedure through discharge are captured here. In discharges with multiple procedures, outcomes should be included on the procedure they follow so the record reads like a book.

### Death/Cause (OPP)

### **Data Abstraction Instructions:**

Indicate if the patient died in association with this hospitalization. If yes, indicate cause of death.

### Selections:

- Yes
  - Cardiovascular (includes: AMI, bleed, stroke, cardiogenic shock) 0
  - Hemorrhage
  - Multi System Organ Failure (includes acute lung injury, and systemic inflammatory response system)
  - Other (include neurologic, renal, liver, GI, cancer) ٥ Unknown cause of death
  - 0
- No

### Supporting Definitions:

Select no if the patient was alive throughout the hospitalization.

**Required:** 

Yes

### Comfort care measures implemented

**Data Abstraction Instructions:** 

Indicate if care was withdrawn or comfort care measures were implemented prior to death. If so, indicate the date.

### Selections:

- Yes
   Enter date
- c ...
- No

Required: Yes

### Stay in ICU

### Data Abstraction Instructions:

Indicate the length of stay in the ICU post procedure. Include any days in the ICU for any reason.

### Selections:

- Yes
  - Enter value in days
- No

### Supporting Definitions:

Indicate value in days, whole numbers. Less than or equal to 24 hours equals one day, greater than 24 hours up to 48 hours equals two days, etc.

Required: Yes Suffix: days Minimum: 1 Maximum:

100

### Vasopressors Post Operatively

#### **Data Abstraction Instructions:**

Document if dopamine ≥ 5 mcg/kg/min, or Neo-Synephrine, Levophed, epinephrine, vasopressin, or other IV vasopressor was administered post procedure.

#### Selections:

- Yes
- No

### Required:

Yes

### **Respiratory**

### **Data Abstraction Instructions:**

Indicate if the patient had any respiratory issues post procedure.

### Selections:

- Ventilator (continued after leaving OR)
- Reintubation (required after initially extubated)
- None

### Supporting Definitions:

If a patient was not intubated for the procedure and requires intubation post procedure, it is not captured as an outcome here. Do not include elective reintubation for additional procedures.

### **Required:**

Yes

### **Myocardial Injury**

### **Data Abstraction Instructions:**

Indicate if the patient suffered any type of myocardial injury post procedure, including a troponin leak, demand ischemia, NSTEMI or STEMI. If so, indicate the date of the first elevated troponin value as well as the peak troponin value. The peak troponin value should be obtained within 30 days of the procedure.

### Selections:

Yes

- Enter date of first occurence post procedure \_\_\_\_
- Enter type of injury:
  - Troponin leak
  - Demand ischemia

- NSTEMI
  - STEMI
  - Not documented

No

### Supporting Definitions:

Utilize progress notes and consults to help in the determination of the type of myocardial injury. If no determination is made, select "Not documented".

If only a single abnormal troponin value was found in absence of other criteria for myocardial injury, then record as No.

Troponin: Troponin rise alone should be reported if there was a rise in cardiac biomarker values [preferably cardiac troponin (cTn)] with at least one value above the 99th percentile upper reference limit (URL) in absence of the qualifying criteria for myocardial infarction or sudden death as listed in the clinical MI definition below. This elevation may be classified as a troponin leak or demand ischemia. Note, "rise" in troponin would imply that the troponin c an be elevated at baseline (either pre-op or post-op baseline), but not rise above whatever the patient's baseline level is. The lack of rising troponin above the baseline number would indicate that there was no additional myocardial injury.

A myocardial infarction is evidenced by any of the following:

- 1. A rise and fall of cardiac biomarkers (preferably troponin) with at least one of the values in the abnormal range for that laboratory [typically above the 99th percentile of the upper reference limit (URL) for normal subjects] together with at least one of the following manifestations of myocardial ischemia: 2. Ischemic symptoms such as angina or acute shortness of breath.
- ECG changes indicative of new ischemia (new ST-T changes, new left bundle branch block, or loss of R wave voltage). 4. Development of pathological Q waves in 2 or more contiguous leads in the ECG (or equivalent finding for true posterior MI.
- 5. Imaging evidence of new loss of viable myocardium or new regional wall motion abnormality.
- Documentation in the medical record of the diagnosis of acute myocardial infarction based on the cardiac biomarker pattern in the absence of any items enumerated in a-d due to conditions that may mask their appearance (e.g., peri-operative infarct when the patient cannot report ischemic symptoms; baseline left bundle branch block or ventricular pacing).
- 7. ECG changes associated with prior myocardial infarction can include the following (with or without prior symptoms):
  - Any Q wave in leads V2-V3 >= 0.02 seconds or QS complex in leads V2 and V3.
  - Q-wave >= 0.03 seconds and >= 0.1 mV deep or QS complex in leads I, II, aVL, aVF, or V4-V6 in any two leads of a contiguous lead grouping (I, aVL, V6; V4-V6; I, III, and aVF).
  - R-wave >= 0.04 seconds in V1-V2 and R/S >= 1 with a concordant positive T-wave in the absence of a conduction defect.
- 8. Imaging evidence of a region with new loss of viable myocardium at rest in the absence of a non-ischemic cause. This can manifest as:
- 9. Echocardiographic, CT, MR, ventriculographic or nuclear imaging evidence of left ventricular thinning or scarring and a failure to contact appropriately (i.e.,
- hypokinesis, akinesis, or dyskinesis).
- 10. Fixed (non-reversible) perfusion defects on nuclear radioisotope imaging (e.g., MIBI, thallium).
- 11. Cardiac death with symptoms suggestive of myocardial ischemia and presumed new ischemic ECG changes or new LBBB, but death occurred before cardiac biomarkers were obtained, or before cardiac biomarker values would be increased.

#### Source

Thygesen K, Alpert JS, White HD, et al. (Circulation 2007). Universal Definition of Myocardial Infarction. ESC/ACCF/AHA/WHF expert consensus document. AHA Journals, 140(13), 2634-53. https://ahajournals.org/journal/circ

#### **Required:** Yes

### Peak post-operative troponin value

#### **Data Abstraction Instructions:**

Indicate the peak value and type of troponin drawn within 30 days post procedure.

#### Selections:

Peak post-operative troponin

Yes

 troponin I Yes

- Enter lab value
- Pick unit of lab value from list na/dl

21			
		No	<ul><li>ng/mL</li><li>ng/L</li><li>pg/mL</li></ul>
	∘ tr	oponin T ∎ Yes	Enter lab value
		•	Pick unit of lab value from list ng/dL ng/mL pa/mL
	∘ tr	■ No roponin I HS ■ Yes	13
		:	Enter lab value Pick unit of lab value from list
	∘ tr	■ No oponin T HS ■ Yes	
		:	Enter lab value Pick unit of lab value from list • ng/dL • ng/L • ng/L • pg/mL

No
 Not Drawn

Required:

Yes

### Dysrhythmia

### **Data Abstraction Instructions:**

Indicate if there was a new rhythm disturbance post procedure, requiring treatment with medications or cardioversion.

#### Selections:

- Yes
- Enter date of first occurrence post procedure
  No

### Required:

Yes

### **Congestive Heart Failure (CHF)**

### **Data Abstraction Instructions:**

Indicate if it was documented that the patient had new onset or exacerbation of CHF post procedure.

### Selections:

- Yes
- Enter date of first occurrence post procedure
   No

### Supporting Definitions:

Heart failure is defined as physician documentation or report of any of the following clinical symptoms of heart failure described as unusual dyspnea on light exertion, recurrent dyspnea occurring in the supine position, fluid retention, or the description of rales, jugular venous distension, pulmonary edema on physical exam, or pulmonary edema on chest x-ray. A low ejection fraction alone, without clinical evidence of heart failure, does not qualify as heart failure.

#### Required: Yes

### TIA/Stroke (OPP)

### Data Abstraction Instructions:

Indicate if there was abrupt loss of neurological function with complete return of function within 24 hours or loss of neurological function caused by an ischemic event that is severe enough to leave a persistent deficit for greater than 24 hours.

### Selections:

- Yes
- Enter date of first occurrence post procedure
   No

### **Required:**

Yes

### Infection/Sepsis

### **Data Abstraction Instructions:**

Positive cultures requiring treatment with antibiotics. Do not include patients that are placed on antibiotics during a hospitalization with no positive cultures.

Pneumonia may be indicated when evidenced on CXR (lobar infiltrate on CXR and/or pure growth of recognized pathogen or 4+ growth of recognized pathogen in presence of mixed growth) and treatment with antibiotics, even without positive culture.

If yes, select all that apply.

### Selections:

- Yes
  - Access site
  - Central Line/IV
  - Blood
  - Graft infection Pulmonary
  - UTI
  - 0
  - Wound site • Unknown
    - Enter date of first occurrence post procedure (date of first positive culture)

### No

#### Required:

Yes

### New Requirement for Dialysis

#### **Data Abstraction Instructions:**

Indicate if the patient had acute or worsening renal failure, post procedure, that led to dialysis during the hospitalization.

#### Selections:

- Yes
  - Enter date of first occurrence post procedure 0 No

# Supporting Definitions:

Indicate only if the dialysis was initiated post procedure.

### **Required:**

Yes

### Transfusion (OPP)

### **Data Abstraction Instructions:**

Indicate if the patient received any transfusion for any reason post procedure. If yes, select the type of transfusion: PRBC, Whole blood, Platelets, FFP, Other (e.g. Cryoprecipitate, Factor VIII infusion). Select all that apply.

### Selections:

- Yes
- Select type of transfusion • PRBC
  - if yes, enter the number of units for PRBC's (enter # of packed red blood cells 1, 2, 3, etc.)
  - Platelets
  - FFP
  - Other
    - Enter date of first occurrence post procedure

No

Required: Yes Minimum: Maximum:

20

### Hemoglobin prior to Transfusion

### **Data Abstraction Instructions:**

Enter the hemoglobin value drawn prior to the first Transfusion of PRBC's post procedure. This is the value on which they made the decision to transfuse.

- Yes ٠
- No
- Not Documented ٠

Required: Yes Suffix: mg/dL Soft Minimum: 2 Soft Maximum: 20

### Symptomatic Prior to Transfusion

#### **Data Abstraction Instructions:**

Select if the patient was symptomatic prior to transfusion of PRBCs. Select all that apply.

#### Selections:

- Yes
  - Angina
  - Hypotension
  - Tachycardia
  - EKG Changes
  - Shortness of Air
  - Bleeding
  - Cancer/Chronic Anemia
- No

### **Required:**

Yes

### Vascular Access Complications (OPP)

### **Data Abstraction Instructions:**

Indicate vascular complications at the access site requiring transfusion, prolonged hospital stay, causing a drop in hemoglobin 3.0 gm/dl, or any access site complications requiring surgical repair. Select all that apply.

# • Yes

- Yes
  - Retroperitoneal hematoma
  - Pseudo-aneurysm
  - Hematoma at access site
  - Bleeding at access site
     AV fistula
  - Av listula
  - Acute thrombosis
  - Surgical repair of the vascular access site
     Other
- No

### Supporting Definitions:

#### Retroperitoneal hematoma

Pseudoaneurysm. The occurrence of a disruption and dilation of the arterial wall without identification of the arterial wall layers at the site of the catheter entry demonstrated by arteriography or ultrasound.

Hematoma requiring transfusion. Blood loss at the site of arterial or venous access or due to perforation of a traversed artery or vein requiring transfusion and/or prolonging the hospital stay, and/or causing a drop in hemoglobin > 3.0 gm/dl.

Bleeding at access site. Blood loss associated with decreased Hgb (greater than or equal to 3.0 gm/dl) and/or causes an increased length of hospital stay. Without other obvious source (GI, GU, operative, or hemolysis) that is attributable to intraprocedural blood loss (e.g. during equipment changes) should be considered bleeding at the access site even if no hematoma is palpable or documented on imaging studies.

AV fistula. A connection between the access artery and the accompanying vein that is demonstrated by arteriography or ultrasound and most often characterized by a continuous bruit.

Acute thrombosis. Total obstruction of the artery by thrombus most commonly at the site of access.

Surgical repair of the vascular access site (such as surgical closures, exploration of the arteriotomy site, balloon angioplasty or covered stent (JOMED GraftMaster) placement to deal the arterial tear)

### Required:

Yes

### **Compartment Syndrome**

### **Data Abstraction Instructions:**

Indicate if the patient was determined to have compartment syndrome at any time post procedure. When entering compartment syndrome as an outcomes postprocedure, do not enter Return to OR for the same outcome. It is implied that the patient went back to the OR to have the compartment syndrome treated.

### Selections:

- Yes
- Enter date of first occurrence post procedure
   No

#### Supporting Definitions:

### https://bmc2.org/print/book/export/html/78

This is defined as compression of nerves and blood vessels within an enclosed space which leads to muscle and nerve damage and problems with blood flow. Include fasciotomy for prophylaxis.

**Required:** 

Yes

### Embolus (OPP)

#### **Data Abstraction Instructions:**

Indicate if the patient is identified to have an embolus post procedure. If yes, indicate if it was treated successfully. When entering an embolus as an outcomes postprocedure, do not enter Return to OR for the same outcome. It is implied that the patient went back to the OR to have the embolus treated.

### Selections:

- Yes
  - Successful
     Unsuccessful
    - Enter date of first occurrence post procedure
- No

#### Supporting Definitions:

An embolus (compromised of atherosclerotic debris and / or blood clot) moves through the blood vessels until it reaches a vessel that is too small to let it pass. When this happens, the blood flow is stopped by the **embolus**. This occurs after exiting the procedure area.

#### **Required:**

Yes

### Thrombus (OPP)

#### **Data Abstraction Instructions:**

Indicate if a blood clot formed, post procedure, within the treated vessel, that limits distal flow. When entering thrombus as an outcomes post-procedure, do not enter Return to OR for the same outcome. It is implied that the patient went back to the OR to have the thrombus treated.

#### Selections:

- Yes
  - Enter date of first occurrence post procedure No

### Supporting Definitions:

Do not include any thrombus that was present at the beginning of the procedure. If the thrombus formed within a stent of graft, select the outcome "Stent/graft thrombosis" and do not select thrombus.

Required: Yes

### Stent/Graft Thrombosis

#### **Data Abstraction Instructions:**

Indicate if there was a stent/graft thrombosis where blood clot forms within the stent/graft that limits distal blood flow. If yes, indicate if it was treated successfully. When entering stent/graft thrombosis as an outcomes post-procedure, do not enter Return to OR for the same outcome. It is implied that the patient went back to the OR to have the stent/graft thrombosis treated.

#### Selections:

- Yes
  - Successful
  - Unsuccessful
    - Enter date of first occurrence post procedure

Required:

Yes

### Amputation

No

### **Data Abstraction Instructions:**

Indicate if an amputation is performed at any time post procedure. When entering amputation as an outcomes post-procedure, do not enter Return to OR for the same outcome. It is implied that the patient went back to the OR to have the amputation performed.

- Yes
  - Select type of amputation
    - Left hip disarticulation
    - Left AKA
    - Left BKALeft foot

- Left metatarsal
- Left diait
- Right hip disarticulation Right AKA 0
- Right BKA
- Right foot 0
- Right metatarsal Right digit 0
- Enter date of first occurrence post procedure 0

```
    No
```

**Required:** 

Yes

### Return to Operating Room (OPP)

#### **Data Abstraction Instructions:**

Indicate if the patient had to return to the Operating Room post procedure for an open procedure, or to fix the original surgery issues.

#### Selections:

- Yes
  - Bleeding Renal Ischemia

  - Endoleak
  - Infection 0 Graft Revision
  - Other
  - Enter date of first occurrence post procedure
- No

### Supporting Definitions:

If a patient returns to the OR for an evactuation of a hematoma post procedure and the hematoma is not in the treated vessel (for example, a hematoma at the incision site) enter Bleeding as the reason for the return to the OR. even though there may not be any active bleeding, bleeding caused the hematoma.

#### **Required:**

Yes

### **Bowel Ischemia**

### **Data Abstraction Instructions:**

Indicate if the patient had bowel ischemia post procedure. When entering bowel ischemia as an outcomes post-procedure, do not enter Return to OR for the same outcome. It is implied that the patient went back to the OR to have the bowel ischemia treated.

### Selections:

- Yes
  - Medical Treatment
  - Surgical Treatment 0
    - Enter date of first occurrence post procedure

# No

**Required:** Yes

### Was the LOS >2 days after EVAR? (OPP)

#### **Data Abstraction Instructions:**

Indicate if the length of stay (LOS) for the Elective EVAR procedure was >2 days, and the reason the patient was in the hospital >2 days. If Yes is entered, select all reasons that apply.

### Selections:

- Yes
  - Hypertension
  - Lack of transportation No caregiver/support at home
  - COPD 0
  - Urinary retention
  - Other
- No

### Supporting Definitions:

Hypertension = Indicate if the patient experienced hypertension for >24 hours post procedure requiring parenteral drug treatment. Hypertension is defined as a systolic blood pressure (SBP) > 160 mmHg and the need for IV antihypertensives, ACE inhibitors, calcium channel blockers, beta blockers, or diuretics to maintain a SBP <160 mmHa.

COPD = Indicate if the patient developed an exacerbation of COPD after procedure through discharge.

Urinary retention = Patient is unable to void (urinate) requiring catheterization within 24 hours postoperatively or >6 hours after the removal of a preoperatively placed Foley catheter.

**Other** = The reason the patient was in the hospital > 2 days is not on the list.

Required: Yes