

Location: _____

General

Elective Outpatient, Observation, or Admission

- ☐ **Elective outpatient procedure: Discharge following routine recovery** Continuous, Scheduling/ADT, Routine
- ☐ **Outpatient observation services under general supervision** Once, Scheduling/ADT, Routine

Admitting Physician:

Attending Provider:

Patient Condition:

Bed request comments:

- ☐ **Outpatient in a bed - extended recovery** Once, Scheduling/ADT, Routine

Admitting Physician:

Bed request comments:

- ☐ **Admit to Inpatient** Once, 1, Scheduling/ADT, Routine

Admitting Physician:

Level of Care:

Patient Condition:

Bed request comments:

Certification: I certify that based on my best clinical judgment and the patient's condition as documented in the HP and progress notes, I expect that the patient will need hospital services for two or more midnights.

Elective Outpatient, Observation, or Admission

- ☐ **Elective outpatient procedure: Discharge following routine recovery** Continuous, Scheduling/ADT, Routine
- ☐ **Outpatient observation services under general supervision** Once, Scheduling/ADT, Routine

Admitting Physician:

Attending Provider:

Patient Condition:

Bed request comments:

- ☐ **Outpatient in a bed - extended recovery** Once, Scheduling/ADT, Routine

Admitting Physician:

Bed request comments:

- ☐ **Admit to Inpatient** Once, 1, Scheduling/ADT, Routine

Admitting Physician:

Level of Care:

Patient Condition:

Bed request comments:

Certification: I certify that based on my best clinical judgment and the patient's condition as documented in the HP and progress notes, I expect that the patient will need hospital services for two or more midnights.

Isolation

- ☐ **Airborne isolation status**
- ☒ **Airborne isolation status** Continuous, Routine
- ☐ **Mycobacterium tuberculosis by PCR - If you suspect Tuberculosis, please order this test for rapid diagnostics.**
Once, Routine
- ☐ **Contact isolation status** Continuous, Post-op, Routine
- ☐ **Droplet isolation status** Continuous, Post-op, Routine
- ☐ **Enteric isolation status** Continuous, Post-op, Routine

Precautions

- ☐ **Aspiration precautions** Continuous, Post-op, Routine
- ☐ **Fall precautions** Continuous, Post-op, Routine
- Increased observation level needed:
- ☐ **Latex precautions** Continuous, Post-op, Routine
- ☐ **Seizure precautions** Continuous, Post-op, Routine
- Increased observation level needed:

Nursing - Post Procedure

Femoral - Sheath Removal

- ☐ **Closure Devices**

Sign: _____ Printed Name: _____ Date/Time: _____

☒ **The physician must be notified for any signs of complications.** Until discontinued, Post-op, Routine, for abnormal vital signs, uncontrolled pain, absence of pulses/limb discoloration, bleeding, hematoma formation, or signs of complications.

☒ **Activity (Required)**

☒ **Patient was treated with a closure device.** Until discontinued, Post-op, Routine, Bedrest required minimum of *** hours. Keep affected leg straight.

☒ **Patient Education Prior to Sheath Removal and Hospital Discharge**

☒ **Patient education prior to post-sheath removal** Once, 1, Occurrences, S, Post-op, Routine, Provide patient post-sheath removal instructions to include reports of warmth, moistness, swelling, numbness or pain at insertion site.
Patient/Family: ☐ Patient
Education for: ☐ Other (specify) ☐ Activity
Specify: Patient education prior to post sheath removal.

☒ **Patient education prior to discharge** Prior to discharge, S, Post-op, Routine, Provide discharge instruction on emergent physician contact/symptom reporting due to bleeding/hematoma/swelling/pain/tenderness/numbness/tingling, Activity and Limitations and site care. Activity including Limiting movement in affected arm 6 hrs post procedure and keep wrist straight, refrain from lifting or pushing with the affected arm for 48 hrs., and site care.
Patient/Family: ☐ Patient
Education for: ☐ Other (specify) ☐ Activity ☐ Discharge ☐ Smoking cessation counseling
Specify: Patient education prior to discharge.

☒ **Post-Sheath Removal**

☒ **Vital signs after sheath removal** Every 15 min, -1, Post-op, Routine, Assess/document vital signs Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.

☒ **Assess post-sheath cath site** Every 15 min, -1, Post-op, Routine, Assess site for signs and symptoms of a hematoma or other vascular compromise distal to site Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.

☒ **Site care** Once, Post-op, Routine, Ensure complete hemostasis at catheter site, palpate for hematoma, apply appropriate dressing. At a minimum, cover site with 2X2 gauze and transparent dressing.
Site: ☐ catheter site

☐ **Assess for pulse distal to assess site post-sheath removal** Every 15 min, -1, Post-op, Routine, Assess and document Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q 4 hours x4 unless otherwise ordered by physician.
Pulses to assess: ☐ Distal
Side:

☐ **Neurological assessment after sheath removal** Every 15 min, -1, Post-op, Routine, Assess/document neurological assessment Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.
Assessment to Perform:

☐ **Manual Pressure**

☒ **The physician must be notified prior to sheath removal of a systolic blood if pressure >160mmHg.** Until discontinued, Post-op, Routine, prior to sheath removal if systolic blood pressure is >160mmHg.

☒ **Remove sheath** Once, 1, Occurrences, Post-op, Routine, when ACT less than 160 or within physician specified parameters. Sheath may be removed 2 hours after discontinuation of Angiomax (Bivalirudin) infusion unless otherwise specified by physician order.

☒ **The physician must be notified for any signs of complications.** Until discontinued, Post-op, Routine, for abnormal vital signs, uncontrolled pain, absence of pulses/limb discoloration, bleeding, hematoma formation, or signs of complications.

☒ **Activity Post Sheath Removal-Femoral Approach (Required)**

☒ **Bed rest times following Procedure using femoral artery access are: (Must Select One) (Required)**

☐ **Patient was treated with a 4 French catheter. Minimum 15 minutes of pressure at site/Bedrest required minimum of 2 hours.** Until discontinued, Post-op, Routine, Patient may bend unaffected leg. Use urinal or bedpan as needed.

☐ **Patient was treated with a 5 French catheter. Minimum 15 minutes of pressure at site/Bedrest required minimum of 3 hours.** Until discontinued, Post-op, Routine, Patient may bend unaffected leg. Use urinal or bedpan as needed.

☐ **Patient was treated with a 6 French catheter. Minimum 20 minutes for PCI/15 minutes of pressure at site for Diagnostic/Bedrest required minimum of 4 hours.** Until discontinued, Post-op, Routine, Patient may bend unaffected leg. Use urinal or bedpan as needed.

Sign: _____ Printed Name: _____ Date/Time: _____

☐ **Patient was treated with a 7 French or greater catheter. Minimum 25 minutes of pressure at site/Bedrest required minimum of *** hours.** Until discontinued, Post-op, Routine, Bedrest required minimum of *** hours. Keep affected leg straight. Patient may bend unaffected leg. Use urinal or bedpan as needed.

☒ **Patient Education Prior to Sheath Removal and Hospital Discharge**

☒ **Patient education prior to post-sheath removal** Once, 1, Occurrences, S, Post-op, Routine, Provide patient post-sheath removal instructions to include reports of warmth, moistness, swelling, numbness or pain at insertion site.
Patient/Family: ☐ Patient

Education for: ☐ Other (specify) ☐ Activity

Specify: Patient education prior to post sheath removal.

☒ **Patient education prior to discharge** Prior to discharge, S, Post-op, Routine, Provide discharge instruction on emergent physician contact/symptom reporting due to bleeding/hematoma/swelling/pain/tenderness/numbness/tingling, Activity and Limitations and site care. Activity including Limiting movement in affected arm 6 hrs post procedure and keep wrist straight, refrain from lifting or pushing with the affected arm for 48 hrs., and site care.

Patient/Family: ☐ Patient

Education for: ☐ Other (specify) ☐ Activity ☐ Discharge ☐ Smoking cessation counseling

Specify: Patient education prior to discharge.

☒ **Pre-Sheath Removal**

☒ **Vital signs prior to sheath removal** Every 15 min, Post-op, Routine, Vital signs prior to sheath removal - Obtain base line vital signs, include verified ACT results of less than 160 or within parameters ordered by physician, unless otherwise ordered by the physician. For Temp, check every 4 hours.

☒ **Assist patient to void** Once, 1, Occurrences, Post-op, Routine, Assist patient to void prior to sheath removal.

☒ **Assess pre-sheath cath site** Once, 1, Occurrences, Post-op, Routine, Assess for signs and symptoms of hematoma or other vascular compromise distal to site on arrival unless otherwise ordered by the physician. If hematoma is present, mark on skin surface and complete hematoma documentation.

☐ **Patient transferred with sheaths left in place** Until discontinued, Post-op, Routine, Patient transferred with Sheaths left in place.

☐ **Apply hemostatic patch after assessment for hematoma, distal pulses.** Until discontinued, Post-op, Routine, Apply pressure proximal to site, place patch over site, slowly remove sheath, allow blood to moisten patch. Apply direct pressure to site/proximal pressure for ½ allotted time. Slowly release proximal pressure, continue direct pressure over the site for a minimum of 20 minutes for PCI/10 minutes for diagnostic cath.

☐ **Antegrade sheaths present** Until discontinued, Post-op, Routine, Antegrade sheath must be pulled by Physicians or appropriately trained staff in the Cath Lab setting.

☒ **Post-Sheath Removal**

☒ **Vital signs after sheath removal** Every 15 min, -1, Post-op, Routine, Assess/document vital signs Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.

☒ **Assess post-sheath cath site** Every 15 min, -1, Post-op, Routine, Assess site for signs and symptoms of a hematoma or other vascular compromise distal to site Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.

☒ **Site care** Once, Post-op, Routine, Ensure complete hemostasis at catheter site, palpate for hematoma, apply appropriate dressing. At a minimum, cover site with 2X2 gauze and transparent dressing.

Site: ☐ catheter site

☐ **Assess for pulse distal to assess site post-sheath removal** Every 15 min, -1, Post-op, Routine, Assess and document Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q 4 hours x4 unless otherwise ordered by physician.

Pulses to assess: ☐ Distal

Side:

☐ **Neurological assessment after sheath removal** Every 15 min, -1, Post-op, Routine, Assess/document neurological assessment Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician. Assessment to Perform:

☐ **Compression Systems**

☐ **C-clamp (Required)**

☒ **The physician must be notified prior to sheath removal of a systolic blood if pressure >160mmHg.** Until discontinued, Post-op, Routine, prior to sheath removal of a systolic blood if pressure >160mmHg.

☒ **Remove sheath** Once, 1, Occurrences, Post-op, Routine, when ACT less than 160 or within physician specified parameters. Sheath may be removed 2 hours after discontinuation of Angiomax (Bivalirudin) infusion unless otherwise specified by physician order.

Sign: _____ Printed Name: _____ Date/Time: _____

☒ **The physician must be notified for any signs of complications.** Until discontinued, Post-op, Routine, for abnormal vital signs, uncontrolled pain, absence of pulses/limb discoloration, bleeding, hematoma formation, or signs of complications.

☒ **Activity Post Sheath Removal-Femoral Approach (Required)**

☒ **Bed rest times following Procedure using femoral artery access are: (Must Select One) (Required)**

- ☐ **Patient was treated with a 4 French catheter. Minimum 15 minutes of pressure at site/Bedrest required minimum of 2 hours.** Until discontinued, Post-op, Routine, Patient may bend unaffected leg. Use urinal or bedpan as needed.
- ☐ **Patient was treated with a 5 French catheter. Minimum 15 minutes of pressure at site/Bedrest required minimum of 3 hours.** Until discontinued, Post-op, Routine, Patient may bend unaffected leg. Use urinal or bedpan as needed.
- ☐ **Patient was treated with a 6 French catheter. Minimum 20 minutes for PCI/15 minutes of pressure at site for Diagnostic/Bedrest required minimum of 4 hours.** Until discontinued, Post-op, Routine, Patient may bend unaffected leg. Use urinal or bedpan as needed.
- ☐ **Patient was treated with a 7 French or greater catheter. Minimum 25 minutes of pressure at site/Bedrest required minimum of *** hours.** Until discontinued, Post-op, Routine, Bedrest required minimum of *** hours. Keep affected leg straight. Patient may bend unaffected leg. Use urinal or bedpan as needed.

☒ **Patient Education Prior to Sheath Removal and Hospital Discharge**

☒ **Patient education prior to post-sheath removal** Once, 1, Occurrences, S, Post-op, Routine, Provide patient post-sheath removal instructions to include reports of warmth, moistness, swelling, numbness or pain at insertion site.

Patient/Family: ☐ Patient

Education for: ☐ Other (specify) ☐ Activity

Specify: Patient education prior to post sheath removal.

☒ **Patient education prior to discharge** Prior to discharge, S, Post-op, Routine, Provide discharge instruction on emergent physician contact/symptom reporting due to bleeding/hematoma/swelling/pain/tenderness/numbness/tingling, Activity and Limitations and site care. Activity including Limiting movement in affected arm 6 hrs post procedure and keep wrist straight, refrain from lifting or pushing with the affected arm for 48 hrs., and site care.

Patient/Family: ☐ Patient

Education for: ☐ Other (specify) ☐ Activity ☐ Discharge ☐ Smoking cessation counseling

Specify: Patient education prior to discharge.

☒ **Pre-Sheath Removal**

☒ **Vital signs prior to sheath removal** Every 15 min, Post-op, Routine, Vital signs prior to sheath removal - Obtain base line vital signs, include verified ACT results of less than 160 or within parameters ordered by physician, unless otherwise ordered by the physician. For Temp, check every 4 hours.

☒ **Assist patient to void** Once, 1, Occurrences, Post-op, Routine, Assist patient to void prior to sheath removal.

☒ **Assess pre-sheath cath site** Once, 1, Occurrences, Post-op, Routine, Assess for signs and symptoms of hematoma or other vascular compromise distal to site on arrival unless otherwise ordered by the physician. If hematoma is present, mark on skin surface and complete hematoma documentation.

☐ **Patient transferred with sheaths left in place** Until discontinued, Post-op, Routine, Patient transferred with Sheaths left in place.

☐ **Apply hemostatic patch after assessment for hematoma, distal pulses.** Until discontinued, Post-op, Routine, Apply pressure proximal to site, place patch over site, slowly remove sheath, allow blood to moisten patch. Apply direct pressure to site/proximal pressure for ½ allotted time. Slowly release proximal pressure, continue direct pressure over the site for a minimum of 20 minutes for PCI/10 minutes for diagnostic cath.

☐ **Antegrade sheaths present** Until discontinued, Post-op, Routine, Antegrade sheath must be pulled by Physicians or appropriately trained staff in the Cath Lab setting.

☒ **Post-Sheath Removal**

Sign: _____ Printed Name: _____ Date/Time: _____

- ☒ **Vital signs after sheath removal** Every 15 min, -1, Post-op, Routine, Assess/document vital signs Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.
- ☒ **Assess post-sheath cath site** Every 15 min, -1, Post-op, Routine, Assess site for signs and symptoms of a hematoma or other vascular compromise distal to site Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.
- ☒ **Site care** Once, Post-op, Routine, Ensure complete hemostasis at catheter site, palpate for hematoma, apply appropriate dressing. At a minimum, cover site with 2X2 gauze and transparent dressing.
Site: ☐ catheter site
- ☐ **Assess for pulse distal to assess site post-sheath removal** Every 15 min, -1, Post-op, Routine, Assess and document Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q 4 hours x4 unless otherwise ordered by physician.
Pulses to assess: ☐ Distal
Side:
- ☐ **Neurological assessment after sheath removal** Every 15 min, -1, Post-op, Routine, Assess/document neurological assessment Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.
Assessment to Perform:

☐ **Femostop**

- ☒ **The physician must be notified prior to sheath removal of a systolic blood if pressure >160mmHg.** Until discontinued, Post-op, Routine, prior to sheath removal of a systolic blood if pressure >160mmHg.
- ☒ **Remove sheath** Once, 1, Occurrences, Post-op, Routine, when ACT less than 160 or within physician specified parameters. Sheath may be removed 2 hours after discontinuation of Angiomax (Bivalirudin) infusion unless otherwise specified by physician order.
- ☒ **The physician must be notified for any signs of complications.** Until discontinued, Post-op, Routine, capillary refill > 3 seconds, cyanosis, numbness and/or pain in affected extremity, bleeding, hematoma formation, or signs of complication.
- ☒ **Follow Femostop manufacturer's guidelines in package insert.** Until discontinued, Post-op, Routine
- ☒ **Activity Post Sheath Removal-Femoral Approach** (Required)
 - ☒ **Bed rest times following Procedure using femoral artery access are: (Must Select One)** (Required)
 - ☐ **Patient was treated with a 4 French catheter. Minimum 15 minutes of pressure at site/Bedrest required minimum of 2 hours.** Until discontinued, Post-op, Routine, Patient may bend unaffected leg. Use urinal or bedpan as needed.
 - ☐ **Patient was treated with a 5 French catheter. Minimum 15 minutes of pressure at site/Bedrest required minimum of 3 hours.** Until discontinued, Post-op, Routine, Patient may bend unaffected leg. Use urinal or bedpan as needed.
 - ☐ **Patient was treated with a 6 French catheter. Minimum 20 minutes for PCI/15 minutes of pressure at site for Diagnostic/Bedrest required minimum of 4 hours.** Until discontinued, Post-op, Routine, Patient may bend unaffected leg. Use urinal or bedpan as needed.
 - ☐ **Patient was treated with a 7 French or greater catheter. Minimum 25 minutes of pressure at site/Bedrest required minimum of *** hours.** Until discontinued, Post-op, Routine, Bedrest required minimum of *** hours. Keep affected leg straight. Patient may bend unaffected leg. Use urinal or bedpan as needed.
- ☒ **Patient Education Prior to Sheath Removal and Hospital Discharge**
 - ☒ **Patient education prior to post-sheath removal** Once, 1, Occurrences, S, Post-op, Routine, Provide patient post-sheath removal instructions to include reports of warmth, moistness, swelling, numbness or pain at insertion site.
Patient/Family: ☐ Patient
Education for: ☐ Other (specify) ☐ Activity
Specify: Patient education prior to post sheath removal.

Sign: _____ Printed Name: _____ Date/Time: _____

- ☒ **Patient education prior to discharge** Prior to discharge, S, Post-op, Routine, Provide discharge instruction on emergent physician contact/symptom reporting due to bleeding/hematoma/swelling/pain/tenderness/numbness/tingling, Activity and Limitations and site care. Activity including Limiting movement in affected arm 6 hrs post procedure and keep wrist straight , refrain from lifting or pushing with the affected arm for 48 hrs., and site care.
Patient/Family: ☐ Patient
Education for: ☐ Other (specify) ☐ Activity ☐ Discharge ☐ Smoking cessation counseling
Specify: Patient education prior to discharge.

☒ **Pre-Sheath Removal**

- ☒ **Vital signs prior to sheath removal** Every 15 min, Post-op, Routine, Vital signs prior to sheath removal - Obtain base line vital signs, include verified ACT results of less than 160 or within parameters ordered by physician, unless otherwise ordered by the physician. For Temp, check every 4 hours.
- ☒ **Assist patient to void** Once, 1, Occurrences, Post-op, Routine, Assist patient to void prior to sheath removal.
- ☒ **Assess pre-sheath cath site** Once, 1, Occurrences, Post-op, Routine, Assess for signs and symptoms of hematoma or other vascular compromise distal to site on arrival unless otherwise ordered by the physician. If hematoma is present, mark on skin surface and complete hematoma documentation.
- ☐ **Patient transferred with sheaths left in place** Until discontinued, Post-op, Routine, Patient transferred with Sheaths left in place.
- ☐ **Apply hemostatic patch after assessment for hematoma, distal pulses.** Until discontinued, Post-op, Routine, Apply pressure proximal to site, place patch over site, slowly remove sheath, allow blood to moisten patch. Apply direct pressure to site/proximal pressure for ½ allotted time. Slowly release proximal pressure, continue direct pressure over the site for a minimum of 20 minutes for PCI/10 minutes for diagnostic cath.
- ☐ **Antegrade sheaths present** Until discontinued, Post-op, Routine, Antegrade sheath must be pulled by Physicians or appropriately trained staff in the Cath Lab setting.

☒ **Post-Sheath Removal**

- ☒ **Vital signs after sheath removal** Every 15 min, -1, Post-op, Routine, Assess/document vital signs Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.
- ☒ **Assess post-sheath cath site** Every 15 min, -1, Post-op, Routine, Assess site for signs and symptoms of a hematoma or other vascular compromise distal to site Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.
- ☒ **Site care** Once, Post-op, Routine, Ensure complete hemostasis at catheter site, palpate for hematoma, apply appropriate dressing. At a minimum, cover site with 2X2 gauze and transparent dressing.
Site: ☐ catheter site
- ☐ **Assess for pulse distal to assess site post-sheath removal** Every 15 min, -1, Post-op, Routine, Assess and document Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q 4 hours x4 unless otherwise ordered by physician.
Pulses to assess: ☐ Distal
Side:
- ☐ **Neurological assessment after sheath removal** Every 15 min, -1, Post-op, Routine, Assess/document neurological assessment Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.
Assessment to Perform:

Radial - Sheath Removal

- ☐ **Radial Compression Device** (Required)
- ☒ **The physician must be notified prior to sheath removal of a systolic blood if pressure >160mmHg.** Until discontinued, Post-op, Routine, prior to sheath removal if systolic blood pressure is >160mmHg.
- ☒ **Remove sheath** Once, 1, Occurrences, Post-op, Routine, when ACT less than 160 or within physician specified parameters. Sheath may be removed 2 hours after discontinuation of Angiomax (Bivalirudin) infusion unless otherwise specified by physician order.
- ☒ **The physician must be notified for any signs of complications.** Until discontinued, Post-op, Routine, for abnormal vital signs, uncontrolled pain, absence of pulses/limb discoloration, bleeding, hematoma formation, or signs of complications.

Sign: _____ Printed Name: _____ Date/Time: _____

- ☒ **Place/Maintain Radial Compression Device** Continuous, Post-op, Routine, Follow manufacturer insert/instructions for use, physician orders, or Progressive Cuff Deflation instruction specific to Diagnostic or Interventional Procedure performed.
Radial Band
Side: Bilateral
Select Sleeve(s):
- ☒ **Progressive cuff deflation** (Required)
- ☐ **Diagnostic Procedures only** (Required)
- ☒ **30-60 minutes after Radial Compression Device applied** Until discontinued, Post-op, Routine, Begin deflating 1-3cc of air from cuff after 30-60 minutes from application. If no bleeding occurs from site, deflate 1-3cc of air from the Radial Compression Device every 5-15 minutes until all air is completely removed. If bleeding occurs when 1-3cc of air is removed, re-inflate with 1-3cc of air. Wait 15 minutes, then restart releasing 1-3cc of air every 5-15 minutes until all air is completely removed. If site remains free of bleeding/hematoma after 5-15 minutes, remove TR band, apply dressing.
- ☒ **Monitor access site and extremity distal to puncture wound** Until discontinued, Post-op, Routine, every 15 minutes until Radial Compression Device is removed.
- ☒ **Assess for absence of ulnar pulse, capillary refill greater than 3 seconds, cyanosis, numbness and/or pain in affected extremity.** Until discontinued, Post-op, Routine, If any of these are present, notify the procedural Cardiologist.
- ☐ **Interventional Procedures only** (Required)
- ☒ **2 hours after Radial Compression Device applied deflate 3cc** Until discontinued, Post-op, Routine, if no bleeding at site, deflate 1-3cc every 10-15 minutes until all air removed from cuff. If bleeding occurs when 1-3cc of air is removed, re-inflate with 1-3cc of air. Wait 30 minutes then restart releasing 1-3cc of air every 10-15 minutes until all air has been removed. If site remains free of bleeding/hematoma after 5-15 minutes, remove TR band, apply dressing.
- ☒ **Evaluate access site for bleeding as follows:** Until discontinued, Post-op, Routine, every 15 minutes x 4; every 30 minutes x2; and every hour x2.
- ☒ **Patient Education Prior to Sheath Removal and Hospital Discharge**
- ☒ **Patient education prior to post-sheath removal** Once, 1, Occurrences, S, Post-op, Routine, Provide patient post-sheath removal instructions to include reports of warmth, moistness, swelling, numbness or pain at insertion site.
Patient/Family: ☐ Patient
Education for: ☐ Other (specify) ☐ Activity
Specify: Patient education prior to post sheath removal.
- ☒ **Patient education prior to discharge** Prior to discharge, S, Post-op, Routine, Provide discharge instruction on emergent physician contact/symptom reporting due to bleeding/hematoma/swelling/pain/tenderness/numbness/tingling, Activity and Limitations and site care. Activity including Limiting movement in affected arm 6 hrs post procedure and keep wrist straight, refrain from lifting or pushing with the affected arm for 48 hrs., and site care.
Patient/Family: ☐ Patient
Education for: ☐ Other (specify) ☐ Activity ☐ Discharge ☐ Smoking cessation counseling
Specify: Patient education prior to discharge.
- ☒ **Pre-Sheath Removal**
- ☒ **Vital signs prior to sheath removal** Every 15 min, Post-op, Routine, Vital signs prior to sheath removal - Obtain base line vital signs, include verified ACT results of less than 160 or within parameters ordered by physician, unless otherwise ordered by the physician. For Temp, check every 4 hours.
- ☒ **Assist patient to void** Once, 1, Occurrences, Post-op, Routine, Assist patient to void prior to sheath removal.
- ☒ **Assess pre-sheath cath site** Once, 1, Occurrences, Post-op, Routine, Assess for signs and symptoms of hematoma or other vascular compromise distal to site on arrival unless otherwise ordered by the physician. If hematoma is present, mark on skin surface and complete hematoma documentation.
- ☐ **Patient transferred with sheaths left in place** Until discontinued, Post-op, Routine, Patient transferred with Sheaths left in place.
- ☐ **Apply hemostatic patch after assessment for hematoma, distal pulses.** Until discontinued, Post-op, Routine, Apply pressure proximal to site, place patch over site, slowly remove sheath, allow blood to moisten patch. Apply direct pressure to site/proximal pressure for ½ allotted time. Slowly release proximal pressure, continue direct pressure over the site for a minimum of 20 minutes for PCI/10 minutes for diagnostic cath.

Sign: _____ Printed Name: _____ Date/Time: _____

☐ **Antegrade sheaths present** Until discontinued, Post-op, Routine, Antegrade sheath must be pulled by Physicians or appropriately trained staff in the Cath Lab setting.

☒ **Radial Approach (Required)**

☒ **Vital signs after sheath removal** Every 15 min, -1, Post-op, Routine, Assess/document vital signs Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.

☒ **Peripheral vascular assessment - Monitor access site** Every 15 min, Post-op, Routine, Monitor access site, extremity distal to puncture every 15 min until Radial approach cath band removed.

☒ **Notify physician of bleeding and/or loss of pulses.** Until discontinued, Post-op, Routine, Notify physician of bleeding and/or loss of pulses.

☒ **Site care** Once, Post-op, Routine, Ensure complete hemostasis at catheter site, palpate for hematoma, apply appropriate dressing. At a minimum, cover site with 2X2 gauze and transparent dressing.
Site: ☐ catheter site

☒ **No blood pressure readings, lab draws, or IV access** Until discontinued, Post-op, Routine, No blood pressure readings, lab draws, or IV access in the affected arm for 24 hours.

☒ **Limit movement in affected arm 6 hrs post procedure** Until discontinued, Post-op, Routine, keep wrist straight, refrain from lifting or pushing with the affected arm for 48 hrs. If needed, place wrist on arm board to restrict movement.

☒ **Patient may ambulate 30 minutes after arrival in recovery area.** Until discontinued, S, Post-op, Routine
Specify: ☐ Other activity (specify)
Other: Patient may ambulate 30 minutes after arrival in recovery area.

☐ **Assess for pulse distal to access site post-sheath removal** Every 15 min, -1, Post-op, Routine, Assess and document Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q 4 hours x4 unless otherwise ordered by physician.
Pulses to assess: ☐ Distal
Side:

☐ **Neurological assessment after sheath removal** Every 15 min, -1, Post-op, Routine, Assess/document neurological assessment Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.
Assessment to Perform:

☐ **Manual Pressure - without Radial Compression Device**

☒ **The physician must be notified prior to sheath removal of a systolic blood if pressure >160mmHg.** Until discontinued, Post-op, Routine, prior to sheath removal of a systolic blood if pressure >160mmHg.

☒ **Remove sheath** Once, 1, Occurrences, Post-op, Routine, when ACT less than 160 or within physician specified parameters. Sheath may be removed 2 hours after discontinuation of Angiomax (Bivalirudin) infusion unless otherwise specified by physician order.

☒ **Notify physician - for abnormal vital signs, uncontrolled pain, absence of pulses/limb discoloration, bleeding, hematoma formation, or signs of complications.** Until discontinued, Post-op, Routine, for abnormal vital signs, uncontrolled pain, absence of pulses/limb discoloration, bleeding, hematoma formation, or signs of complications.

☒ **Patient Education Prior to Sheath Removal and Hospital Discharge**

☒ **Patient education prior to post-sheath removal** Once, 1, Occurrences, S, Post-op, Routine, Provide patient post-sheath removal instructions to include reports of warmth, moistness, swelling, numbness or pain at insertion site.
Patient/Family: ☐ Patient
Education for: ☐ Other (specify) ☐ Activity
Specify: Patient education prior to post sheath removal.

☒ **Patient education prior to discharge** Prior to discharge, S, Post-op, Routine, Provide discharge instruction on emergent physician contact/symptom reporting due to bleeding/hematoma/swelling/pain/tenderness/numbness/tingling, Activity and Limitations and site care. Activity including Limiting movement in affected arm 6 hrs post procedure and keep wrist straight, refrain from lifting or pushing with the affected arm for 48 hrs., and site care.
Patient/Family: ☐ Patient
Education for: ☐ Other (specify) ☐ Activity ☐ Discharge ☐ Smoking cessation counseling
Specify: Patient education prior to discharge.

☒ **Pre-Sheath Removal**

☒ **Vital signs prior to sheath removal** Every 15 min, Post-op, Routine, Vital signs prior to sheath removal - Obtain base line vital signs, include verified ACT results of less than 160 or within parameters ordered by physician, unless otherwise ordered by the physician. For Temp, check every 4 hours.

☒ **Assist patient to void** Once, 1, Occurrences, Post-op, Routine, Assist patient to void prior to sheath removal.

Sign: _____ Printed Name: _____ Date/Time: _____

☒ **Assess pre-sheath cath site** Once, 1, Occurrences, Post-op, Routine, Assess for signs and symptoms of hematoma or other vascular compromise distal to site on arrival unless otherwise ordered by the physician. If hematoma is present, mark on skin surface and complete hematoma documentation.

☐ **Patient transferred with sheaths left in place** Until discontinued, Post-op, Routine, Patient transferred with Sheaths left in place.

☐ **Apply hemostatic patch after assessment for hematoma, distal pulses.** Until discontinued, Post-op, Routine, Apply pressure proximal to site, place patch over site, slowly remove sheath, allow blood to moisten patch. Apply direct pressure to site/proximal pressure for ½ allotted time. Slowly release proximal pressure, continue direct pressure over the site for a minimum of 20 minutes for PCI/10 minutes for diagnostic cath.

☐ **Antegrade sheaths present** Until discontinued, Post-op, Routine, Antegrade sheath must be pulled by Physicians or appropriately trained staff in the Cath Lab setting.

☒ **Post-Sheath Removal**

☒ **Vital signs after sheath removal** Every 15 min, -1, Post-op, Routine, Assess/document vital signs Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.

☒ **Notify physician of bleeding and/or loss of pulses.** Until discontinued, Post-op, Routine, Notify physician of bleeding and/or loss of pulses.

☒ **Site care** Once, Post-op, Routine, Ensure complete hemostasis at catheter site, palpate for hematoma, apply appropriate dressing. At a minimum, cover site with 2X2 gauze and transparent dressing.
Site: ○ catheter site

☒ **No blood pressure readings, lab draws, or IV access** Until discontinued, Post-op, Routine, No blood pressure readings, lab draws, or IV access in the affected arm for 24 hours.

☒ **Limit movement in affected arm 6 hrs post procedure** Until discontinued, Post-op, Routine, keep wrist straight, refrain from lifting or pushing with the affected arm for 48 hrs. If needed, place wrist on arm board to restrict movement.

☒ **Patient may ambulate 30 minutes after arrival in recovery area.** Until discontinued, Post-op, Routine
Specify: ○ Other activity (specify)

Other: Patient may ambulate 30 minutes after arrival in recovery area.

☐ **Assess for pulse distal to assess site post-sheath removal** Every 15 min, -1, Post-op, Routine, Assess and document Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q 4 hours x4 unless otherwise ordered by physician.

Pulses to assess: ○ Distal

Side:

☐ **Neurological assessment after sheath removal** Every 15 min, -1, Post-op, Routine, Assess/document neurological assessment Q 15 min x4, Q 30 min x4, Q 1 hour x4, and Q4 x4 unless otherwise ordered by the physician.
Assessment to Perform:

Pre-sheath(s) Removal Diet

☐ **Diet Clear Liquids** Diet effective now, Post-op, Routine, Until sheath(s) removed.

Diet(s): ○ Clear Liquids

Advance Diet as Tolerated? ○ No

Cultural/Special:

Other Options:

IDDSI Liquid Consistency:

Fluid Restriction:

Foods to Avoid:

Foods to Avoid:

Diet - Post-sheath(s) Removal

☒ **Diet - Clear Liquids** Diet effective now, Post-op, Routine

Diet(s): ○ Clear Liquids

Advance Diet as Tolerated? ○ Yes

Target Diet: Heart Healthy

Cultural/Special:

Other Options:

IDDSI Liquid Consistency:

Fluid Restriction:

Foods to Avoid:

Foods to Avoid:

Sign: _____ Printed Name: _____ Date/Time: _____

☐ **Diet - Heart Healthy** Diet effective now, Post-op, Routine

Diet(s): ☐ Heart Healthy

Cultural/Special:

Other Options:

Advance Diet as Tolerated?

IDDSI Liquid Consistency:

Fluid Restriction:

Foods to Avoid:

Foods to Avoid:

☐ **Diet - 1800 Kcal/202 gm Carbohydrate** Diet effective now, Post-op, Routine

Diet(s): ☐ Other Diabetic/Cal

Diabetic/Calorie: 1800 Kcal/202 gm Carbohydrate

Cultural/Special:

Other Options:

Advance Diet as Tolerated?

IDDSI Liquid Consistency:

Fluid Restriction:

Foods to Avoid:

Foods to Avoid:

Diet - Post Sheath(s) Removal HMSJ

☐ **Diet - Regular** Diet effective now, Post-op, Routine

Diet(s): ☐ Regular

Cultural/Special:

Other Options:

Advance Diet as Tolerated?

IDDSI Liquid Consistency:

Fluid Restriction:

Foods to Avoid:

Foods to Avoid:

☐ **Diet - 1800 Carb Control Diabetic** Diet effective now, Post-op, Routine

Diet(s): ☐ Other Diabetic/Cal

Diabetic/Calorie: 1800 Kcal/202 gm Carbohydrate

Cultural/Special:

Other Options:

Advance Diet as Tolerated?

IDDSI Liquid Consistency:

Fluid Restriction:

Foods to Avoid:

Foods to Avoid:

☐ **Diet - Heart Healthy** Diet effective now, Post-op, Routine

Diet(s): ☐ Heart Healthy

Cultural/Special:

Other Options:

Advance Diet as Tolerated?

IDDSI Liquid Consistency:

Fluid Restriction:

Foods to Avoid:

Foods to Avoid:

☐ **Diet - Finger Foods** Diet effective now, Post-op, Routine

Diet(s): ☐ Additional Instructions

Additional Instructions: Finger Foods

Cultural/Special:

Other Options:

Advance Diet as Tolerated?

IDDSI Liquid Consistency:

Fluid Restriction:

Foods to Avoid:

Foods to Avoid:

Telemetry and IV

☐ **Telemetry**

☒ **Telemetry monitoring** Continuous, 48, Hours, Routine

Order: Place in Centralized Telemetry Monitor: EKG Monitoring Only (Telemetry Box)

Reason for telemetry:

Can be off of Telemetry for baths? Yes

Can be off for transport and tests? Yes

☒ **Telemetry additional setup information** Continuous, 48, Hours, Routine

High Heart Rate (BPM): 130.000

Low Heart Rate(BPM): 50.000

High PVC's (per minute): 10.000

☐ **Saline lock IV** Continuous, Post-op, Routine

☐ **Maintain IV access** Until discontinued, Post-op, Routine

☐ **Discontinue IV** Once, Post-op, Routine

Hydration Protocol-Prevention of Contrast Induced Nephropathy

IV Fluids

☐ **sodium chloride 0.9 % infusion** 150 mL/hr, intravenous, continuous, 10, Hours, Post-op

☐ **Post-Procedure Hydration**

☒ **Inpatient**

☐ **sodium chloride 0.9% infusion** 125 mL/hr, continuous

☐ **sodium chloride 0.9% infusion** .9 , continuous

Medications

Mild Pain (Pain Score 1-3)

☐ **acetaminophen (TYLENOL) tablet** 650 mg, oral, every 4 hours PRN, Post-op, mild pain (score 1-3)

Maximum of 3 grams of acetaminophen per day from all sources. (Cirrhosis patients maximum: 2 grams per day from all sources).

Moderate Pain (Pain Score 4-6)

☐ **acetaminophen-codeine (TYLENOL #3) 300-30 mg tablet** 1 tablet, oral, every 4 hours PRN, Post-op, moderate pain (score 4-6)

The use of codeine-containing products is contraindicated in patients LESS THAN 12 years of age. Is this patient OVER 12 years of age? Y/N:

Allowance for Patient Preference:

Give if patient is able to tolerate oral medication

☐ **HYDROcodone-acetaminophen (NORCO) 5-325 mg tablet** 1 tablet, oral, every 4 hours PRN, Post-op, moderate pain (score 4-6)

Allowance for Patient Preference:

Give if patient is able to tolerate oral medication

Give if patient can receive oral tablet/capsule.

Severe Pain (Pain Score 7-10)

☐ **morphine 2 mg/mL injection** 2 mg, intravenous, every 2 hour PRN, Post-op, severe pain (score 7-10)

Use if patient is unable to swallow or faster onset is needed

☐ **fentaNYL (SUBLIMAZE) injection** 25 mcg, intravenous, every 2 hour PRN, Post-op, severe pain (score 7-10)

Use if patient is unable to swallow or faster onset is needed

Beta-Blockers

☐ **metoprolol tartrate (LOPRESSOR) tablet** 25 mg, oral, 2 times daily, Post-op

BP & HR HOLD parameters for this order:

Contact Physician if:

☐ **metoprolol succinate XL (TOPROL-XL) 24 hr tablet** 25 mg, oral, daily, Post-op

BP & HR HOLD parameters for this order:

Contact Physician if:

Do not crush or chew.

☐ **carvedilol (COREG) tablet** 3.125 mg, oral, 2 times daily, Post-op

BP & HR HOLD parameters for this order:

Contact Physician if:

Nitrates

☐ **nitroglycerin infusion** intravenous, continuous, Post-op

Sign: _____ Printed Name: _____ Date/Time: _____

☐ **isosorbide mononitrate (ISMO, MONOKET) tablet** 20 mg, oral, 2 times daily at 0900, 1600, Post-op

BP HOLD parameters for this order:

Contact Physician if:

Post-Op

☐ **isosorbide mononitrate (IMDUR) 24 hr tablet** 30 , oral, daily, Post-op

BP HOLD parameters for this order:

Contact Physician if:

Post-Op

Do not crush or chew.

☐ **nitroglycerin (NITRODUR) 24 hr patch** transdermal, daily, 12, Hours, Post-op

Post-Op

☐ **nitroglycerin (NITROSTAT) 2% ointment** 1 inch, Topical, every 6 hours scheduled, Post-op

Post-Op, Apply to chest wall

☐ **nitroglycerin (NITROSTAT) SL tablet** 0.4 mg, sublingual, every 5 min PRN, 3, Occurrences, Post-op, chest pain

Post-Op. Call provider after third dose.

Antiplatelet Agents - ONE MUST BE SELECTED (Required)

☐ **Loading Dose Followed By Maintenance**

☐ **clopidogrel (PLAVIX) 300 mg Loading Dose followed by 75 mg Maintenance Dose and aspirin EC 81 mg tablet**

☒ **clopidogrel (PLAVIX) 300 mg Loading Dose followed by 75 mg Maintenance Dose and aspirin EC 81 mg tablet**

☒ **clopidogrel (PLAVIX) Loading and Maintenance doses**

☒ **Loading Dose - clopidogrel (PLAVIX) tablet** 300 mg, oral, once, 1, Occurrences, Post-op
Loading Dose

☒ **Maintenance Dose - clopidogrel (PLAVIX) tablet** 75 mg, oral, daily, S+1, Post-op
Maintenance Dose

☐ **ticagrelor (BRILINTA) 180 mg Loading Dose followed by 90 mg Maintenance Dose and aspirin EC 81 mg tablet**

☒ **ticagrelor (BRILINTA) Oral Loading and Maintenance Doses**

☒ **ticagrelor (BRILINTA) tablet** 180 mg, oral, once, 1, Occurrences, Pre-op

Does the patient have active or a history of pathological bleeding (e.g., peptic ulcer or intracranial hemorrhage)?
Is the patient receiving maintenance aspirin dose greater than 100 mg/day?

☒ **ticagrelor (BRILINTA) tablet** 90 mg, oral, 2 times daily, Pre-op

Does the patient have active or a history of pathological bleeding (e.g., peptic ulcer or intracranial hemorrhage)?
Is the patient receiving maintenance aspirin dose greater than 100 mg/day?

☒ **aspirin (ECOTRIN) enteric coated tablet** 81 mg, oral, daily, S+1, Post-op

☐ **prasugrel (EFFIENT) 60 mg Loading Dose followed by 10 mg Maintenance Dose and aspirin EC 81 mg tablet**
(Required)

☒ **prasugrel (EFFIENT) Loading and Maintenance Doses**

Maintenance Dose Instructions:

Lower the dose to 5 mg for high risk patients (age GREATER than or EQUAL to 75 OR weight LESS than 60 kg)

☒ **prasugrel (EFFIENT) tablet** 60 mg, oral, once, 1, Occurrences, Pre-op

Does this patient have a history of transient ischemic attack (TIA) or stroke?
Is the patient's age 75 years or older?

Is the patient's weight less than 60 kilograms?

☒ **prasugrel (EFFIENT) tablet** 10 mg, oral, daily, Pre-op

Does this patient have a history of transient ischemic attack (TIA) or stroke?
Is the patient's age 75 years or older?

Is the patient's weight less than 60 kilograms?

☒ **aspirin chewable tablet** 81 mg, oral, once, 1, Occurrences, S+1, Post-op

☒ **Pharmacy Consult to educate patient on prasugrel (EFFIENT) (Required)**

Sign: _____ Printed Name: _____ Date/Time: _____

☒ **Pharmacy Consult to educate patient on prasugrel (EFFIENT)** Once, 1, Occurrences, STAT

Which drug do you need help dosing? ☐ prasugrel (EFFIENT)

Contact Number:

☐ **Maintenance Doses Only**

☐ **clopidogrel (PLAVIX) 75 mg Maintenance Dose and aspirin EC 81 mg tablet - Start Tomorrow**

☒ **clopidogrel (PLAVIX) tablet** 75 mg, oral, daily, S+1, Post-op

☒ **aspirin (ECOTRIN) enteric coated tablet** 81 mg, oral, daily, S+1, Post-op

☐ **ticagrelor (BRILINTA) 90 mg Maintenance Dose and aspirin EC 81 mg tablet - Start 12 Hours from Now**

☒ **ticagrelor (BRILINTA) tablet** 90 mg, oral, 2 times daily, Post-op

Does the patient have active or a history of pathological bleeding (e.g., peptic ulcer or intracranial hemorrhage)?

Is the patient receiving maintenance aspirin dose greater than 100 mg/day?

☒ **aspirin (ECOTRIN) enteric coated tablet** 81 mg, oral, daily, S+1, Post-op

☐ **prasugrel (EFFIENT) 10 mg Maintenance Dose and aspirin EC 81 mg tablet - Start Tomorrow**

☒ **prasugrel (EFFIENT) tablet** (Required)

☒ **prasugrel (EFFIENT) tablet** 5 , oral, daily

Does this patient have a history of transient ischemic attack (TIA) or stroke?

Is the patient's age 75 years or older?

Is the patient's weight less than 60 kilograms?

☒ **aspirin (ECOTRIN) enteric coated tablet** 81 mg, oral, daily, S+1, Post-op

☐ **Anti-Platelet Contraindication** Until discontinued, Post-op, Routine

Reason for "No" order:

☐ **Patient already on antiplatelet therapy** Until discontinued, Routine

Antihyperlipidemic Agents - ONE MUST BE SELECTED (Required)

☐ **Moderate Intensity**

☐ **atorvastatin (LIPITOR) tablet - Moderate Intensity** 10 mg, oral, nightly, Post-op

☐ **atorvastatin (LIPITOR) tablet - Moderate Intensity** 20 mg, oral, nightly, Post-op

☐ **rosuvastatin (CRESTOR) tablet - Moderate Intensity** 10 mg, oral, nightly, Post-op

☐ **High Intensity**

☐ **atorvastatin (LIPITOR) tablet - High Intensity** 40 mg, oral, nightly, Post-op

☐ **atorvastatin (LIPITOR) tablet - High Intensity** 80 mg, oral, nightly, Post-op

☐ **rosuvastatin (CRESTOR) tablet - High Intensity** 10 mg, oral, nightly, Post-op

☐ **The patient is currently on a statin** Once, Routine

☐ **The patient is not on a statin due to contraindication.** Once, Routine

The patient is not on a statin due to: ☐ Other

Other: Contraindication

☐ **Discharge medication prescription - evolocumab (REPATHA) subcutaneous pen or wearable injector**

☐ **evolocumab (Repatha SureClick) 140 mg/mL pen injector injection** 140 mg, subcutaneous, every 14 days, 2 mL, 0

☐ **evolocumab (Repatha Pushtronex) 420 mg/3.5 mL wearable injector** 420 mg, subcutaneous (via wearable injector), every 30 days, 3.5 mL, 0

GPIIb/IIIa Inhibitors

☐ **eptifibatide (INTEGRILIN) 0.75 mg/mL infusion** 0.75 , intravenous, continuous, Post-op

☐ **bivalirudin (ANGIOMAX) 5 mg/mL in sodium chloride 0.9 % 50 mL infusion** intravenous, continuous, Post-op

Do not administer other IV medications with or in the same IV line

ACE/ARB Inhibitors

☐ **enalaprilat (VASOTEC) injection** 0.625 mg, intravenous, every 6 hours, Post-op

BP HOLD parameters for this order:

Contact Physician if:

Sign: _____ Printed Name: _____ Date/Time: _____

☐ **enalapril (VASOTEC) tablet** 40 mg, oral, daily, Post-op

BP HOLD parameters for this order:

Contact Physician if:

☐ **captopril (CAPOTEN) tablet** 25 mg, oral, 3 times daily, Post-op

BP HOLD parameters for this order:

Contact Physician if:

☐ **lisinopril (PRINIVIL,ZESTRIL) tablet** 5 mg, oral, daily, Post-op

BP HOLD parameters for this order:

Contact Physician if:

☐ **valsartan (DIOVAN) tablet** 160 mg, oral, 2 times daily, Post-op

BP HOLD parameters for this order:

Contact Physician if:

☐ **losartan (COZAAR) tablet** 50 mg, oral, daily, Post-op

BP HOLD parameters for this order:

Contact Physician if:

Anti-Anginal

☐ **ranolazine (RANEXA) 12 hr tablet** 500 , oral, 2 times daily, Post-op

For Sheath(s) Pull Only

☐ **atropine injection** 0.5 mg, intravenous, once PRN, Post-op, for heart rate LESS than 55 beats per minute

☐ **diazepam (VALIUM) injection** 1 mg, intravenous, once PRN, Post-op, sedation

Indication(s): Sedation

Indication:

☐ **MIDAZolam (VERSED) injection** 1 mg, intravenous, once PRN, Post-op, sedation

Indication(s): ◦ Sedation

☐ **fentaNYL (SUBLIMAZE) injection** 25 mcg, intravenous, once PRN, Post-op, sheath pull

☐ **morPHINE injection** 1 mg, intravenous, once PRN, Post-op, sheath pull

Antiemetics

☒ **ondansetron (ZOFRAN) IV or Oral (Required)**

☒ **ondansetron ODT (ZOFRAN-ODT) disintegrating tablet** 4 mg, oral, every 8 hours PRN, nausea vomiting

Give if patient is able to tolerate oral medication.

May cause QTc prolongation.

☒ **ondansetron (ZOFRAN) 4 mg/2 mL injection** 4 mg, intravenous, every 8 hours PRN, nausea vomiting

Give if patient is UNable to tolerate oral medication OR if a faster onset of action is required.

May cause QTc prolongation.

☐ **promethazine (PHENERGAN) IV or Oral or Rectal**

☒ **promethazine (PHENERGAN) injection** 12.5 mg, intravenous, every 6 hours PRN, nausea vomiting

Give if ondansetron (ZOFRAN) is ineffective and patient is UNable to tolerate oral or rectal medication OR if a faster onset of action is required.

☒ **promethazine (PHENERGAN) tablet** 12.5 mg, oral, every 6 hours PRN, nausea vomiting

Give if ondansetron (ZOFRAN) is ineffective and patient is able to tolerate oral medication.

☒ **promethazine (PHENERGAN) suppository** 12.5 mg, rectal, every 6 hours PRN, nausea vomiting

Give if ondansetron (ZOFRAN) is ineffective and patient is UNable to tolerate oral medication.

Anxiolytics

☐ **LORazepam (ATIVAN) tablet** 0.5 mg, oral, every 4 hours PRN, Post-op, anxiety

Indication(s): ◦ Anxiety

☐ **ALPRAZolam (XANAX) tablet** 0.25 mg, oral, every 8 hours PRN, Post-op, anxiety

Indication(s): ◦ Anxiety

Insomnia: For Patients LESS than 70 years old

☐ **zolpidem (AMBIEN) or ramelteon (ROZEREM) tablet** nightly PRN sleep

Sign: _____ Printed Name: _____ Date/Time: _____

- ☐ **zolpidem (AMBIEN) tablet** 5 mg, oral, nightly PRN, sleep
- ☐ **ramelteon (ROZEREM) tablet** 8 mg, oral, nightly PRN, sleep

Insomnia: For Patients GREATER than 70 years old

- ☐ **ramelteon (ROZEREM) tablet** 8 mg, oral, nightly PRN, Post-op, sleep

HM IP CARDIAC CATHETERIZATION POST PROCEDURE - OTHER

- ☐ **docusate sodium (COLACE) capsule** 100 mg, oral, 2 times daily PRN, Post-op, constipation
- ☐ **magnesium hydroxide suspension** 30 mL, 4 times daily PRN, Post-op, indigestion

VTE

DVT Risk and Prophylaxis Tool (Required)

- ☐ **Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis** Once, PACU & Post-op, Routine
No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication.
Therapy for the following:

- ☐ **LOW Risk of VTE (Required)**

☒ **Low Risk (Required)**

☒ **Low risk of VTE** Once, Routine

Low risk: ☐ Due to low risk, no VTE prophylaxis is needed. Will encourage early ambulation ☐ Due to low risk, no VTE prophylaxis is needed. Will encourage early ambulation

- ☐ **MODERATE Risk of VTE - Surgical (Required)**

☒ **Moderate Risk (Required)**

☒ **Moderate risk of VTE** Once, Routine

☒ **Moderate Risk Pharmacological Prophylaxis - Surgical Patient (Required)**

- ☐ **Contraindications exist for pharmacologic prophylaxis - Order Sequential compression device**

☒ **Contraindications exist for pharmacologic prophylaxis** Once, Routine

No pharmacologic VTE prophylaxis due to the following contraindication(s):

☒ **Place/Maintain sequential compression device continuous** Continuous, Routine

Side: Bilateral

Select Sleeve(s):

- ☐ **Contraindications exist for pharmacologic prophylaxis AND mechanical prophylaxis**

☒ **Contraindications exist for pharmacologic prophylaxis** Once, Routine

No pharmacologic VTE prophylaxis due to the following contraindication(s):

☒ **Contraindications exist for mechanical prophylaxis** Once, Routine

No mechanical VTE prophylaxis due to the following contraindication(s):

- ☐ **Enoxaparin (LOVENOX) for Prophylactic Anticoagulation (Required)**

Patient renal status: @CRCL@

For patients with CrCl GREATER than or EQUAL to 30mL/min, enoxaparin orders will apply the following recommended doses by weight:

Weight	Dose
LESS THAN 100kg	enoxaparin 40mg daily
100 to 139kg	enoxaparin 30mg every 12 hours
GREATER THAN or EQUAL to 140kg	enoxaparin 40mg every 12 hours

- ☐ **ENOXAPARIN 30 MG DAILY**

Sign: _____ Printed Name: _____ Date/Time: _____

- ☒ **enoxaparin (LOVENOX) injection** 30 mg, subcutaneous, daily at 1700, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

- ☐ **ENOXAPARIN SQ DAILY**

- ☒ **enoxaparin (LOVENOX) injection** subcutaneous, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

- ☐ **fondaparinux (ARIXTRA) injection** 2.5 mg, subcutaneous, daily, S+1

If the patient does not have a history of or suspected case of Heparin-Induced Thrombocytopenia (HIT) do NOT order this medication. Contraindicated in patients LESS than 50kg, prior to surgery/invasive procedure, or CrCl LESS than 30 mL/min.

- ☐ **heparin**

High Risk Bleeding Characteristics
Age \geq 75
Weight < 50 kg
Unstable Hgb
Renal impairment
Plt count < 100 K/uL
Dual antiplatelet therapy
Active cancer
Cirrhosis/hepatic failure
Prior intra-cranial hemorrhage
Prior ischemic stroke
History of bleeding event requiring admission and/or transfusion
Chronic use of NSAIDs/steroids
Active GI ulcer

- ☐ **High Bleed Risk**

Every 12 hour frequency is appropriate for most high bleeding risk patients. However, some high bleeding risk patients also have high clotting risk in which every 8 hour frequency may be clinically appropriate.

Please weight the risks/benefits of bleeding and clotting when selecting the dosing frequency.

- ☐ **HEParin (porcine) injection - Q12 Hours** 5000 Units, every 12 hours scheduled

- ☐ **HEParin (porcine) injection - Q8 Hours** 5000 Units, every 8 hours scheduled

- ☐ **Not high bleed risk**

- ☐ **Wt > 100 kg** 7500 Units, subcutaneous, every 8 hours scheduled

- ☐ **Wt LESS than or equal to 100 kg** 5000 Units, subcutaneous, every 8 hours scheduled

- ☐ **warfarin (COUMADIN)**

- ☐ **WITHOUT pharmacy consult** 1 , oral, daily at 1700

Indication:

Dose Selection Guidance:

- ☐ **Medications**

- ☒ **Pharmacy consult to manage warfarin (COUMADIN)** Until discontinued, Routine

Indication:

- ☐ **warfarin (COUMADIN) tablet** 1 , oral

Indication:

Dose Selection Guidance:

- ☒ **Mechanical Prophylaxis (Required)**

Sign: _____ Printed Name: _____ Date/Time: _____

☐ **Contraindications exist for mechanical prophylaxis** Once, Routine
No mechanical VTE prophylaxis due to the following contraindication(s):

☒ **Place/Maintain sequential compression device continuous** Continuous, Routine
Side: Bilateral
Select Sleeve(s):

☐ **MODERATE Risk of VTE - Non-Surgical** (Required)

☒ **Moderate Risk Pharmacological Prophylaxis - Non-Surgical Patient** (Required)

☒ **Moderate Risk** (Required)

☒ **Moderate risk of VTE** Once, Routine

☒ **Moderate Risk Pharmacological Prophylaxis - Non-Surgical Patient** (Required)

☐ **Contraindications exist for pharmacologic prophylaxis - Order Sequential compression device**

☒ **Contraindications exist for pharmacologic prophylaxis** Once, Routine
No pharmacologic VTE prophylaxis due to the following contraindication(s):

☒ **Place/Maintain sequential compression device continuous** Continuous, Routine
Side: Bilateral
Select Sleeve(s):

☐ **Contraindications exist for pharmacologic prophylaxis AND mechanical prophylaxis**

☒ **Contraindications exist for pharmacologic prophylaxis** Once, Routine
No pharmacologic VTE prophylaxis due to the following contraindication(s):

☒ **Contraindications exist for mechanical prophylaxis** Once, Routine
No mechanical VTE prophylaxis due to the following contraindication(s):

☐ **Enoxaparin for Prophylactic Anticoagulation Nonsurgical** (Required)

Patient renal status: @CRCL@

For patients with CrCl GREATER than or EQUAL to 30mL/min, enoxaparin orders will apply the following recommended doses by weight:

Weight	Dose
LESS THAN 100kg	enoxaparin 40mg daily
100 to 139kg	enoxaparin 30mg every 12 hours
GREATER THAN or EQUAL to 140kg	enoxaparin 40mg every 12 hours

☐ **ENOXAPARIN 30 MG DAILY**

☒ **enoxaparin (LOVENOX) injection** 30 mg, subcutaneous, daily at 1700, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **ENOXAPARIN SQ DAILY**

☒ **enoxaparin (LOVENOX) injection** subcutaneous, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

Sign: _____ Printed Name: _____ Date/Time: _____

- ☐ **fondaparinux (ARIXTRA) injection** 2.5 mg, subcutaneous, daily

If the patient does not have a history of or suspected case of Heparin-Induced Thrombocytopenia (HIT), do NOT order this medication. Contraindicated in patients LESS than 50kg, prior to surgery/invasive procedure, or CrCl LESS than 30 mL/min

- ☐ **heparin**

High Risk Bleeding Characteristics
Age \geq 75
Weight < 50 kg
Unstable Hgb
Renal impairment
Plt count < 100 K/uL
Dual antiplatelet therapy
Active cancer
Cirrhosis/hepatic failure
Prior intra-cranial hemorrhage
Prior ischemic stroke
History of bleeding event requiring admission and/or transfusion
Chronic use of NSAIDs/steroids
Active GI ulcer

- ☐ **High Bleed Risk**

Every 12 hour frequency is appropriate for most high bleeding risk patients. However, some high bleeding risk patients also have high clotting risk in which every 8 hour frequency may be clinically appropriate.

Please weight the risks/benefits of bleeding and clotting when selecting the dosing frequency.

- ☐ **HEParin (porcine) injection - Q12 Hours** 5000 Units, every 12 hours scheduled

- ☐ **HEParin (porcine) injection - Q8 Hours** 5000 Units, every 8 hours scheduled

- ☐ **Not high bleed risk**

- ☐ **Wt > 100 kg** 7500 Units, subcutaneous, every 8 hours scheduled

- ☐ **Wt LESS than or equal to 100 kg** 5000 Units, subcutaneous, every 8 hours scheduled

- ☐ **warfarin (COUMADIN)**

- ☐ **WITHOUT pharmacy consult** 1 , oral, daily at 1700

Indication:

Dose Selection Guidance:

- ☐ **Medications**

- ☒ **Pharmacy consult to manage warfarin (COUMADIN)** Until discontinued, Routine

Indication:

- ☐ **warfarin (COUMADIN) tablet** 1 , oral

Indication:

Dose Selection Guidance:

- ☒ **Mechanical Prophylaxis** (Required)

- ☐ **Contraindications exist for mechanical prophylaxis** Once, Routine

No mechanical VTE prophylaxis due to the following contraindication(s):

- ☒ **Place/Maintain sequential compression device continuous** Continuous, Routine

Side: Bilateral

Select Sleeve(s):

- ☐ **HIGH Risk of VTE - Surgical** (Required)

- ☒ **High Risk** (Required)

- ☒ **High risk of VTE** Once, Routine

Sign: _____ Printed Name: _____ Date/Time: _____

☒ **High Risk Pharmacological Prophylaxis - Surgical Patient (Required)**

☐ **Contraindications exist for pharmacologic prophylaxis** Once, Routine
No pharmacologic VTE prophylaxis due to the following contraindication(s):

☐ **Enoxaparin (LOVENOX) for Prophylactic Anticoagulation (Required)**

Patient renal status: @CRCL@

For patients with CrCl GREATER than or EQUAL to 30mL/min, enoxaparin orders will apply the following recommended doses by weight:

Weight	Dose
LESS THAN 100kg	enoxaparin 40mg daily
100 to 139kg	enoxaparin 30mg every 12 hours
GREATER THAN or EQUAL to 140kg	enoxaparin 40mg every 12 hours

☐ **ENOXAPARIN 30 MG DAILY**

☒ **enoxaparin (LOVENOX) injection** 30 mg, subcutaneous, daily at 1700, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **ENOXAPARIN SQ DAILY**

☒ **enoxaparin (LOVENOX) injection** subcutaneous, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **fondaparinux (ARIXTRA) injection** 2.5 mg, subcutaneous, daily, S+1

If the patient does not have a history or suspected case of Heparin-Induced Thrombocytopenia (HIT) do NOT order this medication. Contraindicated in patients LESS than 50kg, prior to surgery/invasive procedure, or CrCl LESS than 30 mL/min.

☐ **heparin**

High Risk Bleeding Characteristics

Age \geq 75

Weight < 50 kg

Unstable Hgb

Renal impairment

Plt count < 100 K/uL

Dual antiplatelet therapy

Active cancer

Cirrhosis/hepatic failure

Prior intra-cranial hemorrhage

Prior ischemic stroke

History of bleeding event requiring admission and/or transfusion

Chronic use of NSAIDs/steroids

Active GI ulcer

☐ **High Bleed Risk**

Every 12 hour frequency is appropriate for most high bleeding risk patients. However, some high bleeding risk patients also have high clotting risk in which every 8 hour frequency may be clinically appropriate.

Sign: _____ Printed Name: _____ Date/Time: _____

Please weight the risks/benefits of bleeding and clotting when selecting the dosing frequency.

- ☐ **HEParin (porcine) injection - Q12 Hours** 5000 Units, every 12 hours scheduled
- ☐ **HEParin (porcine) injection - Q8 Hours** 5000 Units, every 8 hours scheduled
- ☐ **Not high bleed risk**
- ☐ **Wt > 100 kg** 7500 Units, subcutaneous, every 8 hours scheduled
- ☐ **Wt LESS than or equal to 100 kg** 5000 Units, subcutaneous, every 8 hours scheduled
- ☐ **warfarin (COUMADIN)**
- ☐ **WITHOUT pharmacy consult** 1 , oral, daily at 1700
Indication:
Dose Selection Guidance:
- ☐ **Medications**
- ☒ **Pharmacy consult to manage warfarin (COUMADIN)** Until discontinued, Routine
Indication:
☐ **warfarin (COUMADIN) tablet** 1 , oral
Indication:
Dose Selection Guidance:

☒ **Mechanical Prophylaxis (Required)**

- ☐ **Contraindications exist for mechanical prophylaxis** Once, Routine
No mechanical VTE prophylaxis due to the following contraindication(s):
- ☒ **Place/Maintain sequential compression device continuous** Continuous, Routine
Side: Bilateral
Select Sleeve(s):

☐ **HIGH Risk of VTE - Non-Surgical (Required)**

☒ **High Risk (Required)**

- ☒ **High risk of VTE** Once, Routine

☒ **High Risk Pharmacological Prophylaxis - Non-Surgical Patient (Required)**

- ☐ **Contraindications exist for pharmacologic prophylaxis** Once, Routine
No pharmacologic VTE prophylaxis due to the following contraindication(s):
- ☐ **Enoxaparin for Prophylactic Anticoagulation Nonsurgical (Required)**
Patient renal status: @CRCL@

For patients with CrCl GREATER than or EQUAL to 30mL/min, enoxaparin orders will apply the following recommended doses by weight:

Weight	Dose
LESS THAN 100kg	enoxaparin 40mg daily
100 to 139kg	enoxaparin 30mg every 12 hours
GREATER THAN or EQUAL to 140kg	enoxaparin 40mg every 12 hours

☐ **ENOXAPARIN 30 MG DAILY**

- ☒ **enoxaparin (LOVENOX) injection** 30 mg, subcutaneous, daily at 1700, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

Sign: _____ Printed Name: _____ Date/Time: _____

☐ **ENOXAPARIN SQ DAILY**☒ **enoxaparin (LOVENOX) injection** subcutaneous, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **fondaparinux (ARIXTRA) injection** 2.5 mg, subcutaneous, daily

If the patient does not have a history of or suspected case of Heparin-Induced Thrombocytopenia (HIT) do NOT order this medication. Contraindicated in patients LESS than 50kg, prior to surgery/invasive procedure, or CrCl LESS than 30 mL/min.

☐ **heparin****High Risk Bleeding Characteristics**Age \geq 75

Weight < 50 kg

Unstable Hgb

Renal impairment

Plt count < 100 K/uL

Dual antiplatelet therapy

Active cancer

Cirrhosis/hepatic failure

Prior intra-cranial hemorrhage

Prior ischemic stroke

History of bleeding event requiring admission and/or transfusion

Chronic use of NSAIDs/steroids

Active GI ulcer

☐ **High Bleed Risk**

Every 12 hour frequency is appropriate for most high bleeding risk patients. However, some high bleeding risk patients also have high clotting risk in which every 8 hour frequency may be clinically appropriate.

Please weight the risks/benefits of bleeding and clotting when selecting the dosing frequency.

☐ **HEParin (porcine) injection - Q12 Hours** 5000 Units, every 12 hours scheduled☐ **HEParin (porcine) injection - Q8 Hours** 5000 Units, every 8 hours scheduled☐ **Not high bleed risk**☐ **Wt > 100 kg** 7500 Units, subcutaneous, every 8 hours scheduled☐ **Wt LESS than or equal to 100 kg** 5000 Units, subcutaneous, every 8 hours scheduled☐ **warfarin (COUMADIN)**☐ **WITHOUT pharmacy consult** 1 , oral, daily at 1700

Indication:

Dose Selection Guidance:

☐ **Medications**☒ **Pharmacy consult to manage warfarin (COUMADIN)** Until discontinued, Routine

Indication:

☐ **warfarin (COUMADIN) tablet** 1 , oral

Indication:

Dose Selection Guidance:

☒ **Mechanical Prophylaxis (Required)**☐ **Contraindications exist for mechanical prophylaxis** Once, Routine
No mechanical VTE prophylaxis due to the following contraindication(s):☒ **Place/Maintain sequential compression device continuous** Continuous, Routine

Side: Bilateral

Select Sleeve(s):

Sign: _____ Printed Name: _____ Date/Time: _____

☐ **HIGH Risk of VTE - Surgical (Hip/Knee) (Required)**

☒ **High Risk (Required)**

☒ **High risk of VTE** Once, Routine

☒ **High Risk Pharmacological Prophylaxis - Hip or Knee (Arthroplasty) Surgical Patient (Required)**

☐ **Contraindications exist for pharmacologic prophylaxis** Once, Routine

No pharmacologic VTE prophylaxis due to the following contraindication(s):

☐ **aspirin chewable tablet** 162 mg, daily, S+1

☐ **aspirin (ECOTRIN) enteric coated tablet** 162 mg, daily, S+1

☐ **Apixaban and Pharmacy Consult (Required)**

☒ **apixaban (ELIQUIS) tablet** 2.5 mg, 2 times daily, S+1

Indications: ☐ VTE prophylaxis

☒ **Pharmacy consult to monitor apixaban (ELIQUIS) therapy** Until discontinued, STAT

Indications: VTE prophylaxis

☐ **Enoxaparin (LOVENOX) for Prophylactic Anticoagulation (Required)**

Patient renal status: @CRCL@

For patients with CrCl GREATER than or EQUAL to 30mL/min, enoxaparin orders will apply the following recommended doses by weight:

Weight	Dose
LESS THAN 100kg	enoxaparin 40mg daily
100 to 139kg	enoxaparin 30mg every 12 hours
GREATER THAN or EQUAL to 140kg	enoxaparin 40mg every 12 hours

☐ **ENOXAPARIN 30 MG DAILY**

☒ **enoxaparin (LOVENOX) injection** 30 mg, subcutaneous, daily at 1700, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **ENOXAPARIN SQ DAILY**

☒ **enoxaparin (LOVENOX) injection** subcutaneous, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **fondaparinux (ARIXTRA) injection** 2.5 mg, subcutaneous, daily, S+1

If the patient does not have a history or suspected case of Heparin-Induced Thrombocytopenia (HIT) do NOT order this medication. Contraindicated in patients LESS than 50kg, prior to surgery/invasive procedure, or CrCl LESS than 30 mL/min

☐ **heparin**

Sign: _____ Printed Name: _____ Date/Time: _____

High Risk Bleeding Characteristics
Age \geq 75
Weight < 50 kg
Unstable Hgb
Renal impairment
Plt count < 100 K/uL
Dual antiplatelet therapy
Active cancer
Cirrhosis/hepatic failure
Prior intra-cranial hemorrhage
Prior ischemic stroke
History of bleeding event requiring admission and/or transfusion
Chronic use of NSAIDs/steroids
Active GI ulcer

☐ **High Bleed Risk**

Every 12 hour frequency is appropriate for most high bleeding risk patients. However, some high bleeding risk patients also have high clotting risk in which every 8 hour frequency may be clinically appropriate.

Please weight the risks/benefits of bleeding and clotting when selecting the dosing frequency.

☐ **HEParin (porcine) injection - Q12 Hours** 5000 Units, every 12 hours scheduled

☐ **HEParin (porcine) injection - Q8 Hours** 5000 Units, every 8 hours scheduled

☐ **Not high bleed risk**

☐ **Wt > 100 kg** 7500 Units, subcutaneous, every 8 hours scheduled

☐ **Wt LESS than or equal to 100 kg** 5000 Units, subcutaneous, every 8 hours scheduled

☐ **Rivaroxaban and Pharmacy Consult (Required)**

☒ **rivaroxaban (XARELTO) tablet for hip or knee arthroplasty planned during this admission** 10 mg, daily at 0600 (TIME CRITICAL)

Indications: ☐ VTE prophylaxis

For Xarelto 15 mg and 20 mg, give with food or follow administration with enteral feeding to increase medication absorption. Do not administer via post-pyloric routes.

☒ **Pharmacy consult to monitor rivaroxaban (XARELTO) therapy** Until discontinued, STAT

Indications: VTE prophylaxis

☐ **warfarin (COUMADIN)**

☐ **WITHOUT pharmacy consult** 1 , oral, daily at 1700

Indication:

Dose Selection Guidance:

☐ **Medications**

☒ **Pharmacy consult to manage warfarin (COUMADIN)** Until discontinued, Routine

Indication:

☐ **warfarin (COUMADIN) tablet** 1 , oral

Indication:

Dose Selection Guidance:

☒ **Mechanical Prophylaxis (Required)**

☐ **Contraindications exist for mechanical prophylaxis** Once, Routine
No mechanical VTE prophylaxis due to the following contraindication(s):

☒ **Place/Maintain sequential compression device continuous** Continuous, Routine

Side: Bilateral

Select Sleeve(s):

DVT Risk and Prophylaxis Tool

Sign: _____ Printed Name: _____ Date/Time: _____

☐ **Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis** Once, PACU & Post-op, Routine
No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication.
Therapy for the following:

☐ **LOW Risk of VTE** (Required)

☒ **Low Risk** (Required)

☒ **Low risk of VTE** Once, Routine

Low risk: ☐ Due to low risk, no VTE prophylaxis is needed. Will encourage early ambulation ☐ Due to low risk, no VTE prophylaxis is needed. Will encourage early ambulation

☐ **MODERATE Risk of VTE - Surgical** (Required)

☒ **Moderate Risk** (Required)

☒ **Moderate risk of VTE** Once, Routine

☒ **Moderate Risk Pharmacological Prophylaxis - Surgical Patient** (Required)

☐ **Contraindications exist for pharmacologic prophylaxis - Order Sequential compression device**

☒ **Contraindications exist for pharmacologic prophylaxis** Once, Routine

No pharmacologic VTE prophylaxis due to the following contraindication(s):

☒ **Place/Maintain sequential compression device continuous** Continuous, Routine

Side: Bilateral

Select Sleeve(s):

☐ **Contraindications exist for pharmacologic prophylaxis AND mechanical prophylaxis**

☒ **Contraindications exist for pharmacologic prophylaxis** Once, Routine

No pharmacologic VTE prophylaxis due to the following contraindication(s):

☒ **Contraindications exist for mechanical prophylaxis** Once, Routine

No mechanical VTE prophylaxis due to the following contraindication(s):

☐ **Enoxaparin (LOVENOX) for Prophylactic Anticoagulation** (Required)

Patient renal status: @CRCL@

For patients with CrCl GREATER than or EQUAL to 30mL/min, enoxaparin orders will apply the following recommended doses by weight:

Weight	Dose
LESS THAN 100kg	enoxaparin 40mg daily
100 to 139kg	enoxaparin 30mg every 12 hours
GREATER THAN or EQUAL to 140kg	enoxaparin 40mg every 12 hours

☐ **ENOXAPARIN 30 MG DAILY**

☒ **enoxaparin (LOVENOX) injection** 30 mg, subcutaneous, daily at 1700, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **ENOXAPARIN SQ DAILY**

☒ **enoxaparin (LOVENOX) injection** subcutaneous, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

Sign: _____ Printed Name: _____ Date/Time: _____

- ☐ **fondaparinux (ARIXTRA) injection** 2.5 mg, subcutaneous, daily, S+1

If the patient does not have a history of or suspected case of Heparin-Induced Thrombocytopenia (HIT) do NOT order this medication. Contraindicated in patients LESS than 50kg, prior to surgery/invasive procedure, or CrCl LESS than 30 mL/min.

- ☐ **heparin**

High Risk Bleeding Characteristics
Age \geq 75
Weight < 50 kg
Unstable Hgb
Renal impairment
Plt count < 100 K/uL
Dual antiplatelet therapy
Active cancer
Cirrhosis/hepatic failure
Prior intra-cranial hemorrhage
Prior ischemic stroke
History of bleeding event requiring admission and/or transfusion
Chronic use of NSAIDs/steroids
Active GI ulcer

- ☐ **High Bleed Risk**

Every 12 hour frequency is appropriate for most high bleeding risk patients. However, some high bleeding risk patients also have high clotting risk in which every 8 hour frequency may be clinically appropriate.

Please weight the risks/benefits of bleeding and clotting when selecting the dosing frequency.

- ☐ **HEParin (porcine) injection - Q12 Hours** 5000 Units, every 12 hours scheduled

- ☐ **HEParin (porcine) injection - Q8 Hours** 5000 Units, every 8 hours scheduled

- ☐ **Not high bleed risk**

- ☐ **Wt > 100 kg** 7500 Units, subcutaneous, every 8 hours scheduled

- ☐ **Wt LESS than or equal to 100 kg** 5000 Units, subcutaneous, every 8 hours scheduled

- ☐ **warfarin (COUMADIN)**

- ☐ **WITHOUT pharmacy consult** 1 , oral, daily at 1700

Indication:

Dose Selection Guidance:

- ☐ **Medications**

- ☒ **Pharmacy consult to manage warfarin (COUMADIN)** Until discontinued, Routine

Indication:

- ☐ **warfarin (COUMADIN) tablet** 1 , oral

Indication:

Dose Selection Guidance:

- ☒ **Mechanical Prophylaxis (Required)**

- ☐ **Contraindications exist for mechanical prophylaxis** Once, Routine

No mechanical VTE prophylaxis due to the following contraindication(s):

- ☒ **Place/Maintain sequential compression device continuous** Continuous, Routine

Side: Bilateral

Select Sleeve(s):

- ☐ **MODERATE Risk of VTE - Non-Surgical (Required)**

- ☒ **Moderate Risk Pharmacological Prophylaxis - Non-Surgical Patient (Required)**

- ☒ **Moderate Risk (Required)**

- ☒ **Moderate risk of VTE** Once, Routine

Sign: _____ Printed Name: _____ Date/Time: _____

☒ **Moderate Risk Pharmacological Prophylaxis - Non-Surgical Patient** (Required)

☐ **Contraindications exist for pharmacologic prophylaxis - Order Sequential compression device**

☒ **Contraindications exist for pharmacologic prophylaxis** Once, Routine

No pharmacologic VTE prophylaxis due to the following contraindication(s):

☒ **Place/Maintain sequential compression device continuous** Continuous, Routine

Side: Bilateral

Select Sleeve(s):

☐ **Contraindications exist for pharmacologic prophylaxis AND mechanical prophylaxis**

☒ **Contraindications exist for pharmacologic prophylaxis** Once, Routine

No pharmacologic VTE prophylaxis due to the following contraindication(s):

☒ **Contraindications exist for mechanical prophylaxis** Once, Routine

No mechanical VTE prophylaxis due to the following contraindication(s):

☐ **Enoxaparin for Prophylactic Anticoagulation Nonsurgical** (Required)

Patient renal status: @CRCL@

For patients with CrCl GREATER than or EQUAL to 30mL/min, enoxaparin orders will apply the following recommended doses by weight:

Weight	Dose
LESS THAN 100kg	enoxaparin 40mg daily
100 to 139kg	enoxaparin 30mg every 12 hours
GREATER THAN or EQUAL to 140kg	enoxaparin 40mg every 12 hours

☐ **ENOXAPARIN 30 MG DAILY**

☒ **enoxaparin (LOVENOX) injection** 30 mg, subcutaneous, daily at 1700, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **ENOXAPARIN SQ DAILY**

☒ **enoxaparin (LOVENOX) injection** subcutaneous, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **fondaparinux (ARIXTRA) injection** 2.5 mg, subcutaneous, daily

If the patient does not have a history of or suspected case of Heparin-Induced Thrombocytopenia (HIT), do NOT order this medication. Contraindicated in patients LESS than 50kg, prior to surgery/invasive procedure, or CrCl LESS than 30 mL/min

☐ **heparin**

Sign: _____ Printed Name: _____ Date/Time: _____

High Risk Bleeding Characteristics
Age \geq 75
Weight < 50 kg
Unstable Hgb
Renal impairment
Plt count < 100 K/uL
Dual antiplatelet therapy
Active cancer
Cirrhosis/hepatic failure
Prior intra-cranial hemorrhage
Prior ischemic stroke
History of bleeding event requiring admission and/or transfusion
Chronic use of NSAIDs/steroids
Active GI ulcer

☐ **High Bleed Risk**

Every 12 hour frequency is appropriate for most high bleeding risk patients. However, some high bleeding risk patients also have high clotting risk in which every 8 hour frequency may be clinically appropriate.

Please weight the risks/benefits of bleeding and clotting when selecting the dosing frequency.

☐ **HEParin (porcine) injection - Q12 Hours** 5000 Units, every 12 hours scheduled

☐ **HEParin (porcine) injection - Q8 Hours** 5000 Units, every 8 hours scheduled

☐ **Not high bleed risk**

☐ **Wt > 100 kg** 7500 Units, subcutaneous, every 8 hours scheduled

☐ **Wt LESS than or equal to 100 kg** 5000 Units, subcutaneous, every 8 hours scheduled

☐ **warfarin (COUMADIN)**

☐ **WITHOUT pharmacy consult** 1 , oral, daily at 1700

Indication:

Dose Selection Guidance:

☐ **Medications**

☒ **Pharmacy consult to manage warfarin (COUMADIN)** Until discontinued, Routine

Indication:

☐ **warfarin (COUMADIN) tablet** 1 , oral

Indication:

Dose Selection Guidance:

☒ **Mechanical Prophylaxis (Required)**

☐ **Contraindications exist for mechanical prophylaxis** Once, Routine
No mechanical VTE prophylaxis due to the following contraindication(s):

☒ **Place/Maintain sequential compression device continuous** Continuous, Routine

Side: Bilateral

Select Sleeve(s):

☐ **HIGH Risk of VTE - Surgical (Required)**

☒ **High Risk (Required)**

☒ **High risk of VTE** Once, Routine

☒ **High Risk Pharmacological Prophylaxis - Surgical Patient (Required)**

☐ **Contraindications exist for pharmacologic prophylaxis** Once, Routine
No pharmacologic VTE prophylaxis due to the following contraindication(s):

☐ **Enoxaparin (LOVENOX) for Prophylactic Anticoagulation** (Required)

Patient renal status: @CRCL@

Sign: _____ Printed Name: _____ Date/Time: _____

For patients with CrCl GREATER than or EQUAL to 30mL/min, enoxaparin orders will apply the following recommended doses by weight:

Weight	Dose
LESS THAN 100kg	enoxaparin 40mg daily
100 to 139kg	enoxaparin 30mg every 12 hours
GREATER THAN or EQUAL to 140kg	enoxaparin 40mg every 12 hours

☐ **ENOXAPARIN 30 MG DAILY**

☒ **enoxaparin (LOVENOX) injection** 30 mg, subcutaneous, daily at 1700, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **ENOXAPARIN SQ DAILY**

☒ **enoxaparin (LOVENOX) injection** subcutaneous, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **fondaparinux (ARIXTRA) injection** 2.5 mg, subcutaneous, daily, S+1

If the patient does not have a history or suspected case of Heparin-Induced Thrombocytopenia (HIT) do NOT order this medication. Contraindicated in patients LESS than 50kg, prior to surgery/invasive procedure, or CrCl LESS than 30 mL/min.

☐ **heparin**

High Risk Bleeding Characteristics
Age \geq 75
Weight < 50 kg
Unstable Hgb
Renal impairment
Plt count < 100 K/uL
Dual antiplatelet therapy
Active cancer
Cirrhosis/hepatic failure
Prior intra-cranial hemorrhage
Prior ischemic stroke
History of bleeding event requiring admission and/or transfusion
Chronic use of NSAIDs/steroids
Active GI ulcer

☐ **High Bleed Risk**

Every 12 hour frequency is appropriate for most high bleeding risk patients. However, some high bleeding risk patients also have high clotting risk in which every 8 hour frequency may be clinically appropriate.

Please weight the risks/benefits of bleeding and clotting when selecting the dosing frequency.

☐ **HEParin (porcine) injection - Q12 Hours** 5000 Units, every 12 hours scheduled

☐ **HEParin (porcine) injection - Q8 Hours** 5000 Units, every 8 hours scheduled

☐ **Not high bleed risk**

Sign: _____ Printed Name: _____ Date/Time: _____

- ☐ **Wt > 100 kg** 7500 Units, subcutaneous, every 8 hours scheduled
- ☐ **Wt LESS than or equal to 100 kg** 5000 Units, subcutaneous, every 8 hours scheduled

☐ **warfarin (COUMADIN)**

- ☐ **WITHOUT pharmacy consult** 1, oral, daily at 1700

Indication:

Dose Selection Guidance:

- ☐ **Medications**

- ☒ **Pharmacy consult to manage warfarin (COUMADIN)** Until discontinued, Routine

Indication:

- ☐ **warfarin (COUMADIN) tablet** 1, oral

Indication:

Dose Selection Guidance:

☒ **Mechanical Prophylaxis (Required)**

- ☐ **Contraindications exist for mechanical prophylaxis** Once, Routine

No mechanical VTE prophylaxis due to the following contraindication(s):

- ☒ **Place/Maintain sequential compression device continuous** Continuous, Routine

Side: Bilateral

Select Sleeve(s):

☐ **HIGH Risk of VTE - Non-Surgical (Required)**

☒ **High Risk (Required)**

- ☒ **High risk of VTE** Once, Routine

☒ **High Risk Pharmacological Prophylaxis - Non-Surgical Patient (Required)**

- ☐ **Contraindications exist for pharmacologic prophylaxis** Once, Routine

No pharmacologic VTE prophylaxis due to the following contraindication(s):

- ☐ **Enoxaparin for Prophylactic Anticoagulation Nonsurgical (Required)**

Patient renal status: @CRCL@

For patients with CrCl GREATER than or EQUAL to 30mL/min, enoxaparin orders will apply the following recommended doses by weight:

Weight	Dose
LESS THAN 100kg	enoxaparin 40mg daily
100 to 139kg	enoxaparin 30mg every 12 hours
GREATER THAN or EQUAL to 140kg	enoxaparin 40mg every 12 hours

☐ **ENOXAPARIN 30 MG DAILY**

- ☒ **enoxaparin (LOVENOX) injection** 30 mg, subcutaneous, daily at 1700, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **ENOXAPARIN SQ DAILY**

- ☒ **enoxaparin (LOVENOX) injection** subcutaneous, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

Sign: _____ Printed Name: _____ Date/Time: _____

- ☐ **fondaparinux (ARIXTRA) injection** 2.5 mg, subcutaneous, daily

If the patient does not have a history of or suspected case of Heparin-Induced Thrombocytopenia (HIT) do NOT order this medication. Contraindicated in patients LESS than 50kg, prior to surgery/invasive procedure, or CrCl LESS than 30 mL/min.

- ☐ **heparin**

High Risk Bleeding Characteristics
Age \geq 75
Weight < 50 kg
Unstable Hgb
Renal impairment
Plt count < 100 K/uL
Dual antiplatelet therapy
Active cancer
Cirrhosis/hepatic failure
Prior intra-cranial hemorrhage
Prior ischemic stroke
History of bleeding event requiring admission and/or transfusion
Chronic use of NSAIDs/steroids
Active GI ulcer

- ☐ **High Bleed Risk**

Every 12 hour frequency is appropriate for most high bleeding risk patients. However, some high bleeding risk patients also have high clotting risk in which every 8 hour frequency may be clinically appropriate.

Please weight the risks/benefits of bleeding and clotting when selecting the dosing frequency.

- ☐ **HEParin (porcine) injection - Q12 Hours** 5000 Units, every 12 hours scheduled

- ☐ **HEParin (porcine) injection - Q8 Hours** 5000 Units, every 8 hours scheduled

- ☐ **Not high bleed risk**

- ☐ **Wt > 100 kg** 7500 Units, subcutaneous, every 8 hours scheduled

- ☐ **Wt LESS than or equal to 100 kg** 5000 Units, subcutaneous, every 8 hours scheduled

- ☐ **warfarin (COUMADIN)**

- ☐ **WITHOUT pharmacy consult** 1 , oral, daily at 1700

Indication:

Dose Selection Guidance:

- ☐ **Medications**

- ☒ **Pharmacy consult to manage warfarin (COUMADIN)** Until discontinued, Routine

Indication:

- ☐ **warfarin (COUMADIN) tablet** 1 , oral

Indication:

Dose Selection Guidance:

- ☒ **Mechanical Prophylaxis (Required)**

- ☐ **Contraindications exist for mechanical prophylaxis** Once, Routine

No mechanical VTE prophylaxis due to the following contraindication(s):

- ☒ **Place/Maintain sequential compression device continuous** Continuous, Routine

Side: Bilateral

Select Sleeve(s):

- ☐ **HIGH Risk of VTE - Surgical (Hip/Knee) (Required)**

- ☒ **High Risk (Required)**

- ☒ **High risk of VTE** Once, Routine

- ☒ **High Risk Pharmacological Prophylaxis - Hip or Knee (Arthroplasty) Surgical Patient (Required)**

Sign: _____ Printed Name: _____ Date/Time: _____

☐ **Contraindications exist for pharmacologic prophylaxis** Once, Routine
No pharmacologic VTE prophylaxis due to the following contraindication(s):

- ☐ **aspirin chewable tablet** 162 mg, daily, S+1
- ☐ **aspirin (ECOTRIN) enteric coated tablet** 162 mg, daily, S+1
- ☐ **Apixaban and Pharmacy Consult** (Required)

☒ **apixaban (ELIQUIS) tablet** 2.5 mg, 2 times daily, S+1

Indications: ○ VTE prophylaxis

☒ **Pharmacy consult to monitor apixaban (ELIQUIS) therapy** Until discontinued, STAT

Indications: VTE prophylaxis

☐ **Enoxaparin (LOVENOX) for Prophylactic Anticoagulation** (Required)

Patient renal status: @CRCL@

For patients with CrCl GREATER than or EQUAL to 30mL/min, enoxaparin orders will apply the following recommended doses by weight:

Weight	Dose
LESS THAN 100kg	enoxaparin 40mg daily
100 to 139kg	enoxaparin 30mg every 12 hours
GREATER THAN or EQUAL to 140kg	enoxaparin 40mg every 12 hours

☐ **ENOXAPARIN 30 MG DAILY**

☒ **enoxaparin (LOVENOX) injection** 30 mg, subcutaneous, daily at 1700, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **ENOXAPARIN SQ DAILY**

☒ **enoxaparin (LOVENOX) injection** subcutaneous, S+1

Indication(s):

Administer by deep subcutaneous injection into the left and right anterolateral or posterolateral abdominal wall. Alternate injection site with each administration.

☐ **fondaparinux (ARIXTRA) injection** 2.5 mg, subcutaneous, daily, S+1

If the patient does not have a history or suspected case of Heparin-Induced Thrombocytopenia (HIT) do NOT order this medication. Contraindicated in patients LESS than 50kg, prior to surgery/invasive procedure, or CrCl LESS than 30 mL/min

☐ **heparin**

Sign: _____ Printed Name: _____ Date/Time: _____

High Risk Bleeding Characteristics
Age \geq 75
Weight < 50 kg
Unstable Hgb
Renal impairment
Plt count < 100 K/uL
Dual antiplatelet therapy
Active cancer
Cirrhosis/hepatic failure
Prior intra-cranial hemorrhage
Prior ischemic stroke
History of bleeding event requiring admission and/or transfusion
Chronic use of NSAIDs/steroids
Active GI ulcer

☐ **High Bleed Risk**

Every 12 hour frequency is appropriate for most high bleeding risk patients. However, some high bleeding risk patients also have high clotting risk in which every 8 hour frequency may be clinically appropriate.

Please weight the risks/benefits of bleeding and clotting when selecting the dosing frequency.

☐ **HEParin (porcine) injection - Q12 Hours** 5000 Units, every 12 hours scheduled

☐ **HEParin (porcine) injection - Q8 Hours** 5000 Units, every 8 hours scheduled

☐ **Not high bleed risk**

☐ **Wt > 100 kg** 7500 Units, subcutaneous, every 8 hours scheduled

☐ **Wt LESS than or equal to 100 kg** 5000 Units, subcutaneous, every 8 hours scheduled

☐ **Rivaroxaban and Pharmacy Consult (Required)**

☒ **rivaroxaban (XARELTO) tablet for hip or knee arthroplasty planned during this admission** 10 mg, daily at 0600 (TIME CRITICAL)

Indications: ☐ VTE prophylaxis

For Xarelto 15 mg and 20 mg, give with food or follow administration with enteral feeding to increase medication absorption. Do not administer via post-pyloric routes.

☒ **Pharmacy consult to monitor rivaroxaban (XARELTO) therapy** Until discontinued, STAT

Indications: VTE prophylaxis

☐ **warfarin (COUMADIN)**

☐ **WITHOUT pharmacy consult** 1 , oral, daily at 1700

Indication:

Dose Selection Guidance:

☐ **Medications**

☒ **Pharmacy consult to manage warfarin (COUMADIN)** Until discontinued, Routine

Indication:

☐ **warfarin (COUMADIN) tablet** 1 , oral

Indication:

Dose Selection Guidance:

☒ **Mechanical Prophylaxis (Required)**

☐ **Contraindications exist for mechanical prophylaxis** Once, Routine
No mechanical VTE prophylaxis due to the following contraindication(s):

☒ **Place/Maintain sequential compression device continuous** Continuous, Routine

Side: Bilateral

Select Sleeve(s):

Labs

Labs in 4 hours

Sign: _____ Printed Name: _____ Date/Time: _____

- ☒ **Creatinine level** Once, 1, Occurrences, Post-op, Routine, Blood, 3, In 4 Hours
- ☐ **Basic metabolic panel** Once, 1, Occurrences, Post-op, Routine, Blood, 3, In 4 Hours
- ☐ **CBC with differential** Once, 1, Occurrences, Post-op, Routine, Blood, 3, In 4 Hours
- ☐ **Prothrombin time with INR** Once, Post-op, Routine, Blood, 3, In 4 Hours
- ☐ **Troponin T** Now then every 3 hours, 3, Occurrences, Post-op, Routine, Blood, 3, In 4 Hours.

Labs Tomorrow

- ☐ **Basic metabolic panel** AM draw, 1, Occurrences, Post-op, Routine, Blood, 3
- ☐ **CBC with differential** AM draw, 1, Occurrences, Post-op, Routine, Blood, 3
- ☐ **Prothrombin time with INR** AM draw, 1, Occurrences, Post-op, Routine, Blood, 3
- ☐ **Troponin T** AM draw, 1, Occurrences, Post-op, Routine, Blood, 3
- ☐ **Lipid panel** AM draw, 1, Occurrences, Post-op, Routine, Blood, 3

Other Studies**ECG**

- ☒ **ECG Pre/Post Op (PRN)** Conditional Frequency, 6, Occurrences, S, S+7, Post-op, Routine, 6
Clinical Indications: ○ Chest Pain
Interpreting Physician:
- ☒ **ECG Pre/Post Op (in AM)** Once, Post-op, Routine, 6, In AM, ordering cardiologist to interpret EKG
Clinical Indications: ○ Post-Op Surgery
Interpreting Physician:
- ☐ **ECG Pre/Post Op (STAT)** Once, Post-op, STAT, 6, Ordering cardiologist to interpret EKG
Clinical Indications: ○ Post-Op Surgery
Interpreting Physician:
- ☐ **ECG 12 lead** Every 4 hours, 2, Occurrences, Post-op, Routine, 6
Clinical Indications:
Interpreting Physician:

Echo

- ☐ **Transthoracic Echocardiogram Complete, (w Contrast, Strain and 3D if needed)** 1 time imaging, Post-op, Routine
Does this study require a chemo toxicity strain protocol?
Does this exam need a strain protocol?
Call back number for Critical Findings:
Where should test be performed?
Does this exam need a bubble study?
Preferred interpreting Cardiologist or group:
If this patient has had an echocardiogram ordered/performed within the past 120 hours as indicated by repeat Echo orders report on the left. Please contact the Echo department at 713-441-2222 to discuss the reason for a repeat exam with a cardiologist.

For STAT order, select appropriate STAT Indication. Please enter the cell phone number for the ordering physician so the echo attending can communicate the results of the stat test promptly. If the phone number is not entered, we will not be able to perform the test as stat. Please note that nursing unit phone number or NP phone number do not meet this request'

Other Indications should be ordered for TODAY or Routine.

For Discharge or Observation patient, please choose TODAY as Priority.

Consults**Cardiac Rehabilitation Phase I HMM HMWB****Please unselect if patient does not meet requirements for Cardiac Rehab Phase I**

- ☒ **Consult to Cardiac Rehab Phase 1** Once, Post-op, Routine
Clinical Indications: ○ PCI
Patient's Phone Number:

Referral to Cardiac Rehabilitation Phase II (Required)

Please unselect if patient does not meet requirements for Referral to Cardiac Rehab Phase II and select the order: "The patient will not be referred to cardiac rehab due to:" (a reason is required on this order).

Sign: _____ Printed Name: _____ Date/Time: _____

☒ **Referral to Cardiac Rehab Phase 2** Once, Scheduling/ADT

I am referring my patient to outpatient Cardiac Rehabilitation for: ☐ Initial, Phase II (36 Sessions) prescription for Cardiac Rehabilitation.

Medical justification required: s/p MI (last 12 months)

Patient's Phone Number:

Physicians:

Please attach the following information to the referral, if available. This will assist us with patient care, Insurance reimbursement and patient outcomes.

1. Hospital discharge summary, H&P or office note summarizing patient status.
2. Resting 12 lead EKG.
3. Lipid Profile and other lab reports.
4. Recent graded exercise test (within 3 months).
5. Hearth catheterization report.
6. Echocardiogram report.
7. Current Medication List.

Cardiac Rehabilitation Phase II: is the early outpatient phase of Cardiac Rehabilitation and uses exercise training and lifestyle changes to optimize your physical, psychological and social functioning.

Cardiac Rehabilitation benefits may include:

1. Personalized and monitored exercise program proven to increase life expectancy by five years
2. Nutritional counseling
3. Medication review
4. Reduce fear, anxiety and stress
5. Improve your confidence, well being, stamina and strength so that you can return to your usual activities

Houston Methodist Cardiac Rehabilitation Locations:

1. Houston Methodist DeBakey Heart & Vascular Center, Outpatient Center 16th floor, 6445 Main St., Houston, TX 77030 713.441.5575
2. Houston Methodist Baytown Hospital 4201 Garth Road Plaza 1 Suite 290, Baytown TX 77521 281-420-8878
3. Houston Methodist Clear Lake Hospital, MOB 4, 18123 Upper Bay Dr., Suite 110, Houston, TX 77058 281.523.2121
4. Houston Methodist Continuing Care Hospital 701 S. Fry Rd. Suite 215, Katy, TX 77450 832.522.2273
5. Houston Methodist The Woodlands Hospital 7990 State Highway 242, The Woodlands, TX 77385 936.270.3571
6. Houston Methodist Willowbrook Hospital 13802 Centerfield Drive, Suite 200, Houston, TX 77375 281.737.8742
7. Houston Methodist Sugar Land 16605 SW Freeway, Suite 210, Sugar Land, TX 77479 346.874.2050 Fax: 346.874.2051
8. Houston Methodist Cypress 24518 Northwest Fwy. Medical Office Building 2, Suite 105 Cypress, TX 77429 Phone: 346.356.4444 Fax: 713.799.9635

☐ **The patient will not be referred to cardiac rehab due to:** Once, Scheduling/ADT, Routine

The patient will not be referred to cardiac rehab due to:

Sign: _____ Printed Name: _____ Date/Time: _____