### General

**Common Present on Admission Diagnosis** 

[] Acidosis	Post-op
[] Acute Post-Hemorrhagic Anemia	Post-op
[] Acute Renal Failure	Post-op
[] Acute Respiratory Failure	Post-op
[] Acute Thromboembolism of Deep Veins of Lower Extremities	Post-op
[] Anemia	Post-op
[] Bacteremia	Post-op
[] Bipolar disorder, unspecified	Post-op
[] Cardiac Arrest	Post-op
[] Cardiac Dysrhythmia	Post-op
[] Cardiogenic Shock	Post-op
[] Decubitus Ulcer	Post-op
[] Dementia in Conditions Classified Elsewhere	Post-op
[] Disorder of Liver	Post-op
[] Electrolyte and Fluid Disorder	Post-op
[] Intestinal Infection due to Clostridium Difficile	Post-op
[] Methicillin Resistant Staphylococcus Aureus Infection	Post-op
Obstructive Chronic Bronchitis with Exacerbation	Post-op
[] Other Alteration of Consciousness	Post-op
[] Other and Unspecified Coagulation Defects	Post-op
[] Other Pulmonary Embolism and Infarction	Post-op
[] Phlebitis and Thrombophlebitis	Post-op
[] Protein-calorie Malnutrition	Post-op
[] Psychosis, unspecified psychosis type	Post-op
[] Schizophrenia Disorder	Post-op
[] Sepsis	Post-op
[] Septic Shock	Post-op
[] Septicemia	Post-op
[] Type II or Unspecified Type Diabetes Mellitus with	Post-op
Mention of Complication, Not Stated as Uncontrolled	
[] Urinary Tract Infection, Site Not Specified	Post-op
Elective Outpatient, Observation, or Admission (Single	e Response)
() Elective outpatient procedure: Discharge following	Routine, Continuous, PACU & Post-op
routine recovery	
() Outpatient observation services under general	Admitting Physician:
supervision	Patient Condition:
	Bed request comments:
	PACU & Post-op
() Outpatient in a bed - extended recovery	Admitting Physician:
	Bed request comments:
	PACU & Post-op
() Admit to inpatient	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.
	PACU & Post-op
Admission or Observation (Single Response)	

Admission or Observation (Single Response)

Admitting Physician: Level of Care:
Patient Condition:
Bed request comments:
Certification: I certify that based on my best clinical judgmen
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.
PACU & Post-op
Admitting Physician:
Patient Condition:
Bed request comments:
PACU & Post-op
Admitting Physician:
Bed request comments:
PACU & Post-op
Level of Care:
Bed request comments:
Scheduling/ADT
Routine, Until discontinued, Starting S, Scheduling/ADT
Routine, ontil discontinued, Starting S, Scheduling/ADT
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.
PACU & Post-op
Level of Care:
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Level of Care: Bed request comments: Scheduling/ADT Routine, Until discontinued, Starting S, Scheduling/ADT Level of Care: Bed request comments: Scheduling/ADT Routine, Until discontinued, Starting S, Scheduling/ADT Routine, Until discontinued, Starting S, Scheduling/ADT Code Status decision reached by: Post-op d the patient/surrogate require the use of an interpreter? id the patient/surrogate require the use of an interpreter? oes patient have decision-making capacity? ost-op riority:
Level of Care: Bed request comments: Scheduling/ADT Routine, Until discontinued, Starting S, Scheduling/ADT Level of Care: Bed request comments: Scheduling/ADT Routine, Until discontinued, Starting S, Scheduling/ADT Routine, Until discontinued, Starting S, Scheduling/ADT Code Status decision reached by: Post-op id the patient/surrogate require the use of an interpreter? id the patient/surrogate require the use of an interpreter? oes patient have decision-making capacity? ost-op riority: eason for Consult?
Level of Care: Bed request comments: Scheduling/ADT Routine, Until discontinued, Starting S, Scheduling/ADT Level of Care: Bed request comments: Scheduling/ADT Routine, Until discontinued, Starting S, Scheduling/ADT Routine, Until discontinued, Starting S, Scheduling/ADT Code Status decision reached by: Post-op id the patient/surrogate require the use of an interpreter? id the patient/surrogate require the use of an interpreter? oes patient have decision-making capacity? ost-op riority: eason for Consult? rder?
Level of Care: Bed request comments: Scheduling/ADT Routine, Until discontinued, Starting S, Scheduling/ADT Level of Care: Bed request comments: Scheduling/ADT Routine, Until discontinued, Starting S, Scheduling/ADT Routine, Until discontinued, Starting S, Scheduling/ADT Code Status decision reached by: Post-op id the patient/surrogate require the use of an interpreter? id the patient/surrogate require the use of an interpreter? oes patient have decision-making capacity? ost-op riority: eason for Consult?

[] Consult to Social Work	Reason for Consult: Post-op
() Modified Code	Did the patient/surrogate require the use of an interpreter? Did the patient/surrogate require the use of an interpreter? Does patient have decision-making capacity? Modified Code restrictions: Post-op
() Treatment Restrictions ((For use when a patient is in a cardiopulmonary arrest))	I understand that if the patient is NOT in a cardiopulmonary arrest, the selected treatments will NOT be provided. I understand that all other unselected medically indicated treatments will be provided. Treatment Restriction decision reached by: Specify Treatment Restrictions: Post-op
Isolation	
[] Airborne isolation status	
[] Airborne isolation status	Details
<ul> <li>[] Mycobacterium tuberculosis by PCR - If you suspect Tuberculosis, please order this test for rapid diagnostics.</li> </ul>	Once, Sputum, Post-op
[] Contact isolation status	Details
[] Droplet isolation status	Details
[] Enteric isolation status	Details
Precautions	
[] Aspiration precautions	Post-op
[X] Fall precautions	Increased observation level needed: Post-op
[] Latex precautions	Post-op
[] Seizure precautions	Increased observation level needed: Post-op
Nursing	
Vital Signs	
[] Vital signs - T/P/R/BP	Routine, Every 15 min For 999 Occurrences Times 4, then every 30 minutes times 4, then every hour times 4, then every 4 hours times 24 hours, then every 8 hours if stable., PACU & Post-op
Activity	
[] Ambulate patient	Routine, 3 times daily Specify: with assistive device Device: Day of surgery, Post-op
[] Bed rest	Routine, Until discontinued, Starting S, Post-op
[] Up with assistance	Routine, As needed Specify: Up with assistance Post-op
[] Weight bearing	·
[] Weight bearing LLE	Routine, Until discontinued, Starting S Weight Bearing Status: Extremity: Post-op
[] Weight bearing RLE	Routine, Until discontinued, Starting S Weight Bearing Status: Extremity: Post-op

[] Consult to PT eval and treat	Reasons for referral to Physical Therapy (mark all applicable): Are there any restrictions for positioning or mobility? Please provide safe ranges for HR, BP, O2 saturation( if values are very abnormal): Weight Bearing Status:
[] Consult to PT eval and treat	Reasons for referral to Physical Therapy (mark all applicable): Are there any restrictions for positioning or mobility? Please provide safe ranges for HR, BP, O2 saturation( if values are very abnormal): Weight Bearing Status:
[] Elevate Operative Extremity	Routine, Until discontinued, Starting S Position: Additional instructions: elevate the operative extremity Place 2-3 pillows under affected extremity., Post-op
Equipment	
[] Obtain supply / device:	Routine, Once Obtain: Post-op
[] Postop shoe	Routine, Once Left/Right: Sizes: Gender Size: Special Instructions: Post-op
[] Cast shoe	Routine, Once Left/Right: Sizes: Gender Size: Special Instructions: Post-op
[] Cam Walker boot	Routine, Once Left/Right: Sizes: Gender Size: Special Instructions: Post-op
Nursing Assessments	
[X] Peripheral vascular assessment	Routine, Per unit protocol Until discharge., Post-op
Nursing Interventions	
[X] Pulse oximetry	Routine, With vitals For 24 Hours Current FIO2 or Room Air: And while on PCA., Post-op
[X] Incentive spirometry	Routine, Every hour For 10 Occurrences Respiratory to instruct., Post-op
[] Intake and output	Routine, Every shift For 48 Hours, Post-op
[] Insert and Maintain Foley	
[] Insert Foley catheter	Routine, Once Type: Size: Urinometer needed: If unable to void., Post-op
[] Foley Catheter Care	Routine, Until discontinued, Starting S For 48 Hours Orders: to gravity Post-op
[] Remove Foley catheter	Routine, Once Post-op day ***, Post-op

[X] Place antiembolic stockings	Routine, Once Calf sleeve to non operative leg while in bed., PACU &
	Post-op
[X] Apply ice pack	Routine, Until discontinued, Starting S
	Afftected area:
	Waking hours only?
	Nurse to schedule?
	Special Instructions:
	To affected extremity., Post-op
[] Wound Care Orders + Medication	
[] Wound care orders	Routine, Daily, Starting S+1
	Location:
	Site: Irrigate wound?
	Apply:
	Dressing Type:
	Post-op
[] wound care medications	
Please select the appropriate medicatio	n for wound care
[] mupirocin (BACTROBAN) 2 % ointme	nt Topical, 3 times daily
[] collagenase (SANTYL) ointment	Topical, daily
[] silver sulfadiazine (SILVADENE, SSD)	) 1 % Topical, daily
cream	
[] zinc oxide 20 % ointment	Topical
[] white petrolatum-mineral oil (EUCERII	N) Topical, PRN, dry skin
topical cream	
[] silver gel	topical (top)
[] gel dressing gel	topical (top)
[] Verify patient has DME equipment	Routine, Once For 1 Occurrences, Post-op
[] Nursing communication	Routine, Until discontinued, Starting S, Post-op
[] Care Order Instruction	Routine, Until discontinued, Starting S
	Dispense patient instructions, Post-op
Notify	
	Douting Once For 1 Occurrences, Contact Anosthesis for all
[] Notify Physician (Anesthesia)	Routine, Once For 1 Occurrences, Contact Anesthesia for all issues with Nerve Block., Post-op
[] Notify Hospitalist/Internist of patient's location	
Diet	
L1 Dist Clear liquida, advance as telerated t	a Regular Dist offective new Starting S
[] Diet - Clear liquids, advance as tolerated t	o Regular Diet effective now, Starting S Diet(s): Clear Liquids
	Advance Diet as Tolerated? Yes
	Target Diet: Regular
	Advance target diet criteria:
	IDDSI Liquid Consistency:
	Fluid Restriction:
	Foods to Avoid:
	Please assess bowel sounds between progressions., Post-op
[] Diet - Clear liquids, advance as tolerated t	
1800 Carb Control	Diet(s): Clear Liquids
	Advance Diet as Tolerated? Yes Target Diet: Diabetic 1800 Carb Control
	Advance target diet criteria:
	IDDSI Liquid Consistency:
	Fluid Restriction:
	Foods to Avoid:
	Please assess bowel sounds between progressions., Post-op

<ul> <li>[] Diet - Clear liquids, advance as tolerated to Heart Healthy</li> <li>[] Diet - Clear liquids, advance as tolerated to Renal (80GM Pro, 2-3GM Na, 2-3GM K)</li> </ul>	Diet effective now, Starting S Diet(s): Clear Liquids Advance Diet as Tolerated? Yes Target Diet: Heart Healthy Advance target diet criteria: IDDSI Liquid Consistency: Fluid Restriction: Foods to Avoid: Please assess bowel sounds between progressions., Post-op Diet effective now, Starting S Diet(c): Clear Liquids
(80GM PT0, 2-3GM Na, 2-3GM K)	Diet(s): Clear Liquids Advance Diet as Tolerated? Yes Target Diet: Renal (80GM Pro, 2-3GM Na, 2-3GM K) Advance target diet criteria: IDDSI Liquid Consistency: Fluid Restriction: Foods to Avoid: Please assess bowel sounds between progressions., Post-op
IV	
IV Fluids (Single Response)	
() sodium chloride 0.9 % infusion	75 mL/hr, intravenous, continuous, Post-op
() lactated Ringer's infusion	75 mL/hr, intravenous, continuous, Post-op
Medications	
IV Antibiotics: For Patients LESS than or EQUAL to 120 kg	I
<ul> <li>[] cefazolin (ANCEF) IV - For Patients LESS than or EQUAL to 120 kg</li> </ul>	2 g, intravenous, once, For 1 Doses, Post-op For patients LESS than or EQUAL to 120 kg Reason for Therapy: Surgical Prophylaxis
[] FOR MRSA CONCERN OR SEVERE PENICILLIN ALLERGY - vancomycin (VANCOCIN) IV	15 mg/kg, intravenous, once, For 1 Doses, Post-op Reason for Therapy: Surgical Prophylaxis
IV Antibiotics: For Patients GREATER than 120 kg	
[] cefazolin (ANCEF) IV - For Patients GREATER than 120 kg	3 g, intravenous, once, For 1 Doses, Post-op Reason for Therapy: Surgical Prophylaxis
[] FOR MRSA CONCERN OR SEVERE PENICILLIN ALLERGY - vancomycin (VANCOCIN) IV	15 mg/kg, intravenous, once, For 1 Doses, Post-op Reason for Therapy: Surgical Prophylaxis
Pain Medications Check Prescription Drug Monitoring Program. Prior to initiation of opioid therapy, it is recommended to che assess patient's opioid tolerance status. A summarized vers NaRx Score on the patient's Storyboard. You may access th (https://texas.pmpaware.net/login) Texas PMP	ion of the PMP report may be accessed by clicking on the
Pain Management Guide	
Opioid PCA Conversion to Oral Opioid Regimen	
	orphine in patients with renal dysfunction is not recommended.
Due to risk of accumulation of toxic metabolite, the use of m An alternative opioid should be utilized, if possible.	

[] acetaminophen (TYLENOL) tablet	500 mg, oral, every 6 hours scheduled Use if patient can tolerate oral tablet.
[] acetaminophen (TYLENOL) liquid	500 mg, oral, every 6 hours scheduled
() acetaminophen (TYLENOL) 650 mg tablet or	
[] acetaminophen (TYLENOL) tablet	650 mg, oral, every 6 hours scheduled
	Use if patient can tolerate oral tablet.
[] acetaminophen (TYLENOL) liquid	650 mg, oral, every 6 hours scheduled
() NSAIDS: For Patients LESS than 65 years o	
Response)	
() ibuprofen (ADVIL, MOTRIN) tablet or oral s	uspension "Or" Linked Panel
[] ibuprofen (ADVIL, MOTRIN) tablet	600 mg, oral, every 6 hours scheduled
	Not recommended for patients with eGFR LESS than 30 mL/min or acute kidney injury. Give if patient is able to tolerate oral medication.
[] ibuprofen (MOTRIN) 100 mg/5 mL	600 mg, oral, every 6 hours scheduled
suspension	Not recommended for patients with eGFR LESS than 30 mL/min or
	acute kidney injury.
() naproxen (NAPROSYN) tablet	250 mg, oral, 2 times daily
() celecoxib (CeleBREX) capsule	100 mg, oral, 2 times daily
() ketorolac (TORADOL) injection	30 mg, intravenous, every 6 hours scheduled
(,() ,	For patients LESS THAN 65 years old. Not recommended for patients
	with eGFR LESS than 30 mL/min or acute kidney injury.
() NSAIDS: For Patients GREATER than or EC	
years old (Single Response)	· · · · · ·
() ibuprofen (ADVIL, MOTRIN) tablet or oral s	uspension "Or" Linked Panel
[] ibuprofen (ADVIL, MOTRIN) tablet	600 mg, oral, every 6 hours scheduled, Post-op
	Not recommended for patients with eGFR LESS than 30 mL/min or acute kidney injury. Give if patient is able to tolerate oral medication.
[] ibuprofen (MOTRIN) 100 mg/5 mL	600 mg, oral, every 6 hours scheduled, Post-op
suspension	Not recommended for patients with eGFR LESS than 30 mL/min or
	acute kidney injury.
	Use if patient cannot swallow tablet.
() naproxen (NAPROSYN) tablet	250 mg, oral, 2 times daily, Post-op
	Not recommended for patients with eGFR LESS than 30 mL/min or acute kidney injury.
() celecoxib (CeleBREX) capsule	100 mg, oral, 2 times daily, Post-op
	For age GREATER than or EQUAL to 65 yo and patients LESS than
	50kg. Not recommended for patients with eGFR LESS than 30 mL/min
	or acute kidney injury.
() ketorolac (TORADOL) injection	15 mg, intravenous, every 6 hours scheduled, Post-op
PRN Pain Medications	
response. Adjust dose for renal/liver function ar	sent and patient unable to reliably communicate needs. Monitor closely for nd age. Do not order both scheduled and PRN NSAIDs/APAP PO and short acting IV simultaneously. Oral option and IV options to be
] PRN Oral Medications for Mild Pain (Pain Sc For Patients LESS than 65 years old (Single	
	Response)
For Patients LESS than 65 years old (Single	Response) Ds/APAP simultaneously.
For Patients LESS than 65 years old (Single Do not order both scheduled and PRN NSAI () acetaminophen (TYLENOL) tablet OR oral OR rectal suppository	Response)         Ds/APAP simultaneously.         suspension       "Or" Linked Panel
For Patients LESS than 65 years old (Single Do not order both scheduled and PRN NSAII () acetaminophen (TYLENOL) tablet OR oral OR rectal suppository Maximum of 4 grams of acetaminophen per	Response)         Ds/APAP simultaneously.         suspension       "Or" Linked Panel
For Patients LESS than 65 years old (Single Do not order both scheduled and PRN NSAII () acetaminophen (TYLENOL) tablet OR oral OR rectal suppository Maximum of 4 grams of acetaminophen per sources)	Response)         Ds/APAP simultaneously.         suspension       "Or" Linked Panel         r day from all sources.       (Cirrhosis patients maximum: 2 grams per day from         650 mg, oral, every 6 hours PRN, mild pain (score 1-3)
For Patients LESS than 65 years old (Single Do not order both scheduled and PRN NSAII         () acetaminophen (TYLENOL) tablet OR oral SOR rectal suppository         Maximum of 4 grams of acetaminophen per sources)         [] acetaminophen (TYLENOL) tablet	Response)         Ds/APAP simultaneously.         suspension       "Or" Linked Panel         r day from all sources.       (Cirrhosis patients maximum: 2 grams per day from         650 mg, oral, every 6 hours PRN, mild pain (score 1-3)         Give if patient able to swallow tablet.         650 mg, oral, every 6 hours PRN, mild pain (score 1-3)

[] ibuprofen (ADVIL, MOTRIN) tablet	600 mg, oral, every 6 hours PRN, mild pain (score 1-3), Post-op Give if patient is able to tolerate oral medication.
[] ibuprofen (MOTRIN) 100 mg/5 mL suspension	600 mg, oral, every 6 hours PRN, mild pain (score 1-3), Post-op Use if patient cannot swallow tablet.
() naproxen (NAPROSYN) tablet	250 mg, oral, every 8 hours PRN, mild pain (score 1-3), Post-op Not recommended for patients with eGFR LESS than 30 mL/min or acute kidney injury.
() celecoxib (CeleBREX) capsule	100 mg, oral, 2 times daily PRN, mild pain (score 1-3), Post-op Not recommended for patients with eGFR LESS than 30 mL/min or acute kidney injury.
() ketorolac (TORADOL) injection	15 mg, intravenous, every 6 hours PRN, mild pain (score 1-3) Give if patient unable to swallow tablet.
[] PRN Oral Medications for Mild Pain (Pain Sco For Patients GREATER than or EQUAL to 65 (Single Response)	years old
response. Adjust dose for renal/liver function a	esent and patient unable to reliably communicate needs. Monitor closely for and age. Do not order both scheduled and PRN NSAIDs/APAP PO and short acting IV simultaneously. Oral option and IV options to be
() acetaminophen (TYLENOL) tablet OR oral su	uspension "Or" Linked Panel
	day from all sources. (Cirrhosis patients maximum: 2 grams per day from all
[] acetaminophen (TYLENOL) tablet	650 mg, oral, every 6 hours PRN, mild pain (score 1-3) Use if patient can tolerate oral tablet.
[] acetaminophen (TYLENOL)suspension	650 mg, oral, every 6 hours PRN, mild pain (score 1-3) Use if patient cannot tolerate oral tablet.
<ol> <li>PRN Oral Medications for Moderate Pain (Pain 4-6): For Patients LESS than 65 years old (Sir Response)</li> </ol>	ngle
() acetaminophen-codeine (TYLENOL #3) table	
<ul> <li>acetaminophen-codeine (TYLENOL WITH CODEINE #3) 300-30 mg per tablet</li> </ul>	1 tablet, oral, every 6 hours PRN, moderate pain (score 4-6), Post-op Maximum of 4 grams of acetaminophen per day from all sources. (Cirrhosis patients maximum: 2 grams per day from all sources). Give if patient is able to tolerate oral medication. 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:
[] acetaminophen-codeine 300 mg-30 mg /12.5 mL solution	<ul> <li>12.5 mL, oral, every 6 hours PRN, moderate pain (score 4-6), Post-op Maximum of 4 grams of acetaminophen per day from all sources. (Cirrhosis patients maximum: 2 grams per day from all sources) Use if patient cannot swallow tablet.</li> <li>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:</li> </ul>
() HYDROcodone-acetaminophen 5/325 (NOR OR elixir	
	day from all sources. (Cirrhosis patients maximum: 2 grams per day from all
<ul> <li>[] HYDROcodone-acetaminophen (NORCO)</li> <li>5-325 mg per tablet</li> </ul>	1 tablet, oral, every 6 hours PRN, moderate pain (score 4-6), Post-op Give if patient able to swallow tablet.
[] HYDROcodone-acetaminophen (HYCET) 2.5-108.3 mg/5 mL solution	10 mL, oral, every 6 hours PRN, moderate pain (score 4-6), Post-op Give if patient unable to swallow tablet.
() oxyCODONE (ROXICODONE) immediate release tablet	5 mg, oral, every 6 hours PRN, moderate pain (score 4-6), Post-op Tablets may be crushed. Give if patient able to swallow tablet
() traMADoL (ULTRAM) tablet	50 mg, oral, every 6 hours PRN, moderate pain (score 4-6) Max daily dose 200 mg/day in patients with CrCl < 30 ml/min. Give if patient able to swallow tablet.

old (Single Response)	OD alivir "Or" Linkad Danal
<ul> <li>acetaminophen-codeine (TYLENOL #3) tablet</li> <li>acetaminophen-codeine (TYLENOL WITH CODEINE #3) 300-30 mg per tablet</li> </ul>	<ul> <li>OR elixir "Or" Linked Panel</li> <li>1 tablet, oral, every 6 hours PRN, moderate pain (score 4-6), Post-op Maximum of 4 grams of acetaminophen per day from all sources. (Cirrhosis patients maximum: 2 grams per day from all sources). Give if patient is able to tolerate oral medication. The use of codeine-containing products is contraindicated in patients LESS THAN 12 years of age. Is this patient OVER 12 years of age?</li> </ul>
	Y/N:
[] acetaminophen-codeine 300 mg-30 mg /12.5 mL solution	<ul> <li>12.5 mL, oral, every 6 hours PRN, moderate pain (score 4-6), Post-op Maximum of 4 grams of acetaminophen per day from all sources.</li> <li>(Cirrhosis patients maximum: 2 grams per day from all sources) Use if patient cannot swallow tablet.</li> <li>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:</li> </ul>
<ul> <li>HYDROcodone-acetaminophen 5/325 (NORC OR elixir</li> </ul>	CO) tablet "Or" Linked Panel
	ay from all sources. (Cirrhosis patients maximum: 2 grams per day from all
<ul> <li>[] HYDROcodone-acetaminophen (NORCO)</li> <li>5-325 mg per tablet</li> </ul>	1 tablet, oral, every 6 hours PRN, moderate pain (score 4-6), Post-op Give if patient able to swallow tablet.
<ul> <li>[] HYDROcodone-acetaminophen (HYCET) 2.5-108.3 mg/5 mL solution</li> </ul>	10 mL, oral, every 6 hours PRN, moderate pain (score 4-6), Post-op Give if patient unable to swallow tablet.
() oxyCODONE (ROXICODONE) immediate release tablet	2.5 mg, oral, every 6 hours PRN, moderate pain (score 4-6), Post-op Tablets may be crushed. Give if patient able to swallow tablet
() traMADoL (ULTRAM) tablet	25 mg, oral, every 6 hours PRN, moderate pain (score 4-6) Max daily dose 300mg in patients age GREATER THAN 75 years or 200 mg/day in patients with CrCl < 30 ml/min. Give if patient able to swallow tablet.
] PRN IV Medications for Moderate Pain (Pain So For Patients LESS than 65 years old if unable to Oral Pain Medication. (Single Response)	o tolerate
Due to risk of toxicity, the use of morphine prod recommended. An alternative opioid should be	lucts in patients with renal dysfunction, particularly in ESRD, is not utilized.
() morPHINE injection	2 mg, intravenous, every 3 hours PRN, moderate pain (score 4-6), Post-op Give if patient is NPO, unable to swallow oral medication, or if pain
	unrelieved 60 minutes after giving oral pain medications.
() hydromorPHONE (DILAUDID) injection	unrelieved 60 minutes after giving oral pain medications. 0.5 mg, intravenous, every 3 hours PRN, moderate pain (score 4-6), Post-op Give if patient is NPO, unable to swallow oral medication, or if pain unrelieved 60 minutes after giving oral pain medications
<ul> <li>() ketorolac (TORADOL) IV (Single Response)</li> <li>Do NOT use in patients with eGFR LESS than</li> </ul>	0.5 mg, intravenous, every 3 hours PRN, moderate pain (score 4-6), Post-op Give if patient is NPO, unable to swallow oral medication, or if pain unrelieved 60 minutes after giving oral pain medications
() ketorolac (TORADOL) IV (Single Response) Do NOT use in patients with eGFR LESS than WARNING: Use is contraindicated for treatme	<ul> <li>0.5 mg, intravenous, every 3 hours PRN, moderate pain (score 4-6), Post-op</li> <li>Give if patient is NPO, unable to swallow oral medication, or if pain unrelieved 60 minutes after giving oral pain medications</li> <li>a 30 mL/min.</li> <li>a 30 mg, intravenous, every 6 hours PRN, moderate pain (score 4-6), Post-op</li> <li>Use if patient is NPO, unable to swallow oral medication, or if pain unrelieved 60 minutes after giving oral pain medication.</li> </ul>

	ucts in patients with renal dysfunction, particularly in ESRD, is not utilized. (adjust dose for renal/liver function and age)
() morPHINE injection	2 mg, intravenous, every 3 hours PRN, moderate pain (score 4-6), Post-op Give if patient is NPO, unable to swallow oral medication, or if pain unrelieved 60 minutes after giving oral pain medications.
() hydromorPHONE (DILAUDID) injection	<ul> <li>0.2 mg, intravenous, every 3 hours PRN, moderate pain (score 4-6),</li> <li>Post-op</li> <li>Give if patient is NPO, unable to swallow oral medication, or if pain unrelieved 60 minutes after giving oral pain medications</li> </ul>
<ol> <li>PRN Oral Medications for Severe Pain (Pain Se 7-10): For Patients LESS than 65 years old (Sir Response)</li> </ol>	ngle
Due to risk of toxicity, the use of morphine prod recommended. An alternative opioid should be	ucts in patients with renal dysfunction, particularly in ESRD, is not utilized.
() HYDROcodone-acetaminophen 10/325 (NOR OR elixir	CO) tablet "Or" Linked Panel
Maximum of 4 grams of acetaminophen per da sources)	ay from all sources. (Cirrhosis patients maximum: 2 grams per day from all
<ul> <li>[] HYDROcodone-acetaminophen (NORCO) 10-325 mg per tablet</li> </ul>	1 tablet, oral, every 6 hours PRN, severe pain (score 7-10), Post-op Give if patient able to swallow tablet.
<ul> <li>[] HYDROcodone-acetaminophen (HYCET) 2.5-108.3 mg/5 mL solution</li> </ul>	20 mL, oral, every 6 hours PRN, severe pain (score 7-10), Post-op Give if patient unable to swallow tablet.
() morPHINE immediate-release tablet	15 mg, oral, every 6 hours PRN, severe pain (score 7-10), Post-op Tablets may be crushed. Give if patient able to swallow tablet
<ul> <li>() oxyCODONE (ROXICODONE) immediate release tablet</li> </ul>	10 mg, oral, every 6 hours PRN, severe pain (score 7-10), Post-op Tablets may be crushed. Give if patient able to swallow tablet
<ol> <li>PRN Oral Medications for Severe Pain (Pain Se 7-10): For Patients GREATER than or EQUAL years old (Single Response)</li> </ol>	
	ucts in patients with renal dysfunction, particularly in ESRD, is not utilized.
() oxyCODONE (ROXICODONE) immediate release tablet	5 mg, oral, every 6 hours PRN, severe pain (score 7-10), Post-op Oral tablets may be crushed. Give if patient able to swallow tablet
() morPHINE immediate-release tablet	7.5 mg, oral, every 6 hours PRN, severe pain (score 7-10), Post-op Oral tablets may be crushed. Give if patient able to swallow tablets.
() HYDROcodone-acetaminophen 7.5/325 (NOF OR elixir	RCO) tablet "Or" Linked Panel
Maximum of 4 grams of acetaminophen per da sources)	ay from all sources. (Cirrhosis patients maximum: 2 grams per day from all
<ul><li>[] HYDROcodone-acetaminophen (NORCO) 7.5-325 mg per tablet</li></ul>	1 tablet, oral, every 6 hours PRN, severe pain (score 7-10), Post-op Maximum of 4 grams of acetaminophen per day from all sources. (Cirrhosis patients maximum: 2 grams per day from all sources). Give if patient able to swallow tablet.
<ul> <li>[] HYDROcodone-acetaminophen (HYCET) 2.5-108.3 mg/5 mL solution</li> </ul>	10 mL, oral, every 6 hours PRN, severe pain (score 7-10), Post-op Give if patient unable to swallow tablet.
() HYDROcodone-acetaminophen 10/325 (NOR OR elixir	CO) tablet "Or" Linked Panel
Maximum of 4 grams of acetaminophen per da sources)	ay from all sources. (Cirrhosis patients maximum: 2 grams per day from all
<ul> <li>[] HYDROcodone-acetaminophen (NORCO) 10-325 mg per tablet</li> </ul>	1 tablet, oral, every 6 hours PRN, severe pain (score 7-10), Post-op Give if patient able to swallow tablet.
<ul> <li>[] HYDROcodone-acetaminophen (HYCET)</li> <li>2.5-108.3 mg/5 mL solution</li> </ul>	20 mL, oral, every 6 hours PRN, severe pain (score 7-10), Post-op Give if patient unable to swallow tablet.

() traMADoL (ULTRAM) tablet	50 mg, oral, every 6 hours PRN, severe pain (score 7-10) Max daily dose 300mg in patients age GREATER THAN 75 years or 200 mg/day in patients with CrCl < 30 ml/min. Give if patient able to swallow tablet.
[] PRN IV Medications for Severe Pain (Pain Sc For Patients LESS than 65 years old if unable Oral Pain Medication. (Single Response)	ore 7-10):
	ducts in patients with renal dysfunction, particularly in ESRD, is not e utilized.
() fentaNYL (SUBLIMAZE) injection	25 mcg, intravenous, every 3 hours PRN, severe pain (score 7-10), Post-op Give if patient is NPO, unable to swallow oral medication, or if pain 60
	minutes after giving oral pain medications.
() morPHINE injection	4 mg, intravenous, every 4 hours PRN, severe pain (score 7-10), Post-op
	Give if patient is NPO, unable to swallow oral medication, or if pain unrelieved 60 minutes after giving oral pain medications.
() hydromorPHONE (DILAUDID) injection	0.8 mg, intravenous, every 3 hours PRN, severe pain (score 7-10),
	Post-op
	Give if patient is NPO, unable to swallow oral medication, or if pain 60 minutes after giving oral pain medications.
<ol> <li>PRN IV Medications for Severe Pain (Pain Sc For Patients GREATER than or EQUAL to 65 unable to tolerate Oral Pain Medication. (Sing Response)</li> </ol>	years old if
Due to risk of toxicity, the use of morphine pro recommended. An alternative opioid should b	ducts in patients with renal dysfunction, particularly in ESRD, is not e utilized.
() fentaNYL (SUBLIMAZE) injection	12.5 mcg, intravenous, every 3 hours PRN, severe pain (score 7-10), Post-op Give if patient is NPO, unable to swallow oral medication, or if pain 60
() morPHINE injection	minutes after giving oral pain medications. 2 mg, intravenous, every 4 hours PRN, severe pain (score 7-10),
	Post-op
	Give if patient is NPO, unable to swallow oral medication, or if pain unrelieved 60 minutes after giving oral pain medications.
() hydromorPHONE (DILAUDID) injection	0.5 mg, intravenous, every 4 hours PRN, severe pain (score 7-10), Post-op
	Give if patient is NPO, unable to swallow oral medication, or if pain 60 minutes after giving oral pain medications.
CA Medications - Not HMSJ (Single Response)	
fentaNYL PCA (SUBLIMAZE) 1500 mcg/30 mL · Nursing PCA Orders	+
<ul> <li>[] fentaNYL PCA (SUBLIMAZE) 1500 mcg/30 m Response)</li> </ul>	
() fentaNYL (SUBLIMAZE) 1500 mcg/30 mL PCA solution for Opioid Naive	Nurse Loading Dose: Not Ordered PCA Dose: 10 mcg Lockout: 10 Minutes Four Hour Dose Limit: 150 mcg intravenous, continuous
	For breakthrough pain: Administer only if respiratory rate 12 per minute or more and POSS level of 2 or less. RN may bolus *** every *** hours as needed. If pain persists, may increase PCA demand dose by *** mcg ONCE. If more than 2 bolus doses in 12 hours or if pain persists after increase in demand dose, call ordering prescriber.
	Adjust doses for age, renal function or other factors.

[] PCA Documentation	Routine, Every 12 hours At the beginning (or end of each shift), prior to clearing PCA pump data, the following must be documented: doses delivered, number of attempts, total amount of medication infused (in mg or mcg), and volume remaining in syringe (residual volume).
[] Patient education Pain pump	Routine, Once, Starting S For 1 Occurrences Patient/Family: Education for: Pain pump Provide patient education on appropriate use of PCA including no PCA by proxy. Only the patient may press the dosing button.
[] Pasero Opioid-induced Sedation Scale	Routine, Every 6 hours, Starting S Assess POSS while patient has an active PCA order. Contact provider if score 3 or 4.
[] Notify Physician	<ul> <li>Routine, Until discontinued, Starting S, - PCA pump infusion discontinued for any reason</li> <li>Inadequate analgesia</li> <li>Prior to administration of any other narcotics, antiemetics, or sedatives other than those ordered by the prescriber responsible for IV PCA therapy</li> <li>PCA pump discontinued by any service other than the prescriber responsible for IV PCA therapy</li> </ul>
[] Stop the PCA pump and call ordering physician and/or CERT team for any of the following:	Routine, Until discontinued, Starting S, - Respiratory rate 10 per minute or less         -       Severe and/or recent confusion or disorientation         -       POSS sedation level 4: Somnolent and difficult to arouse         -       Sustained hypotension (SBP less than 90)         -       Excessive nausea or vomiting         -       Urinary retention
[] IV Fluids for provision of PCA Therapy (Single Response)	
() sodium chloride 0.9 % infusion	30 mL/hr, intravenous, continuous
() dextrose 5% infusion	30 mL/hr, intravenous, continuous
) morPHINE PCA 30 mg/30 mL + Nursing PCA Ord	
[] morPHINE PCA 30 mg/30 mL (Single Response	
() morPHINE PCA 30 mg/30 mL in sodium	Nurse Loading Dose: Not Ordered PCA Dose: 1 mg Lockout
chloride 0.9% for Opioid Naive	Interval: 10 Minutes MAX (Four hour dose limit): 20 mg intravenous, continuous For breakthrough pain: Administer only if respiratory rate 12 per minute or more and POSS level of 2 or less. RN may bolus *** every *** hours as needed. If pain persists, may increase PCA demand dose by *** mg ONCE. If more than 2 bolus doses in 12 hours or if pain persists after increase in demand dose, call ordering prescriber.
	Adjust doses for age, renal function or other factors.
[] Nursing PCA Orders	
[] Vital signs - T/P/R/BP	<ul> <li>Routine, Per unit protocol</li> <li>Initially and every 30 minutes for 1 hour after PCA started, bolus administration or dose change; then</li> <li>Every hour x 2 starting second hour after PCA started, bolus administered or dose change; then</li> <li>Every 4 hours until PCA therapy is discontinued.</li> <li>Immediately following PCA administration tubing change</li> </ul>
[] PCA Documentation	Routine, Every 12 hours At the beginning (or end of each shift), prior to clearing PCA pump data, the following must be documented: doses delivered, number of attempts, total amount of medication infused (in mg or mcg), and volume remaining in syringe (residual volume).
[] Patient education Pain pump	Routine, Once, Starting S For 1 Occurrences Patient/Family: Education for: Pain pump Provide patient education on appropriate use of PCA including no PCA by proxy. Only the patient may press the dosing button.

[] Pasero Opioid-induced Sedation Scale	Routine, Every 6 hours, Starting S Assess POSS while patient has an active PCA order. Contact provider if
	score 3 or 4.
[] Notify Physician	Routine, Until discontinued, Starting S, - PCA pump infusion discontinued for any reason
	- Inadequate analgesia
	<ul> <li>Prior to administration of any other narcotics, antiemetics, or</li> </ul>
	sedatives other than those ordered by the prescriber responsible for IV PCA therapy
	- PCA pump discontinued by any service other than the prescribe responsible for IV PCA therapy
<ul> <li>Stop the PCA pump and call ordering physician and/or CERT team for any of the</li> </ul>	Routine, Until discontinued, Starting S, - Respiratory rate 10 per minute or less
following:	- Severe and/or recent confusion or disorientation
.ee	- POSS sedation level 4: Somnolent and difficult to arouse
	- Sustained hypotension (SBP less than 90)
	- Excessive nausea or vomiting
	- Urinary retention
[] IV Fluids for provision of PCA Therapy (Single Response)	
() sodium chloride 0.9 % infusion	30 mL/hr, intravenous, continuous
() dextrose 5% infusion	30 mL/hr, intravenous, continuous
CA Medications - HMSJ Only (Single Response)	
) fentaNYL PCA (SUBLIMAZE) 1500 mcg/30 mL + Nursing PCA Orders	
[] fentaNYL PCA (SUBLIMAZE) 1500 mcg/30 mL Response)	. (Single
() fentaNYL (SUBLIMAZE) 1500 mcg/30 mL	Nurse Loading Dose: Not Ordered PCA Dose: 10
PCA solution for Opioid Naive	mcg Lockout: 10 Minutes Four Hour Dose Limit: 150 mcg intravenous, continuous
	For breakthrough pain: Administer only if respiratory rate 12 per minute
	or more and POSS level of 2 or less. RN may bolus *** every *** hours
	as needed. If pain persists, may increase PCA demand dose by *** mcg
	ONCE. If more than 2 bolus doses in 12 hours or if pain persists after
	increase in demand dose, call ordering prescriber.
	Adjust doses for age, renal function or other factors.
[] Nursing PCA Orders	Deutine Den wit grote est
[] Vital signs - T/P/R/BP	Routine, Per unit protocol
	- Initially and every 30 minutes for 1 hour after PCA started, bolus
	administration or dose change; then
	<ul> <li>Every hour x 2 starting second hour after PCA started, bolus administered or dose change; then</li> </ul>
	- Every 4 hours until PCA therapy is discontinued.
	- Immediately following PCA administration tubing change
[] PCA Documentation	Routine, Every 12 hours
	At the beginning (or end of each shift), prior to clearing PCA pump data,
	the following must be documented: doses delivered, number of attempts
	total amount of medication infused (in mg or mcg), and volume
	remaining in syringe (residual volume).
[] Patient education Pain pump	Routine, Once, Starting S For 1 Occurrences
	Patient/Family:
	Education for: Pain pump
	Provide patient education on appropriate use of PCA including no PCA
	by proxy. Only the patient may press the dosing button.
[] Pasero Opioid-induced Sedation Scale	Routine, Every 6 hours, Starting S
	Assess POSS while patient has an active PCA order. Contact provider it
	score 3 or 4.

[] Notify Physician	Routine, Until discontinued, Starting S, - PCA pump infusion discontinued for any reason - Inadequate analgesia
	<ul> <li>Prior to administration of any other narcotics, antiemetics, or sedatives other than those ordered by the prescriber responsible for IV</li> </ul>
	PCA therapy
	<ul> <li>PCA pump discontinued by any service other than the prescribe responsible for IV PCA therapy</li> </ul>
[] Stop the PCA pump and call ordering physician and/or CERT team for any of the	Routine, Until discontinued, Starting S, - Respiratory rate 10 per minute or less
following:	<ul> <li>Severe and/or recent confusion or disorientation</li> <li>POSS sedation level 4: Somnolent and difficult to arouse</li> <li>Sustained hypotension (SBP less than 90)</li> </ul>
	<ul> <li>Excessive nausea or vomiting</li> <li>Urinary retention</li> </ul>
[] IV Fluids for provision of PCA Therapy (Single Response)	
() sodium chloride 0.9 % infusion	30 mL/hr, intravenous, continuous
() dextrose 5% infusion	30 mL/hr, intravenous, continuous
hydromorPHONE PCA (DILAUDID) 30 mg/30 m Nursing PCA Orders	
[] hydromorPHONE PCA (DILAUDID) 30 mg/30 Response)	mL (Single
<ul> <li>() hydromorPHONE (DILAUDID) 30 mg/30 mL in sodium chloride 0.9% PCA for Opioid Naive</li> </ul>	Nurse Loading Dose: Not Ordered PCA Dose: 0.2 mg Lockou Interval: 10 Minutes MAX (Four hour dose limit): 3 mg intravenous, continuous
	For breakthrough pain: Administer only if respiratory rate 12 per minute
	or more and POSS level of 2 or less. RN may bolus *** every *** hours
	as needed. If pain persists, may increase PCA demand dose by *** mg
	ONCE. If more than 2 bolus doses in 12 hours or if pain persists after increase in demand dose, call ordering prescriber.
	Adjust doses for age, renal function or other factors.
[] Nursing PCA Orders	
[] Vital signs - T/P/R/BP	Routine, Per unit protocol - Initially and every 30 minutes for 1 hour after PCA started, bolus administration or dose change; then
	- Every hour x 2 starting second hour after PCA started, bolus
	administered or dose change; then
	<ul> <li>Every 4 hours until PCA therapy is discontinued.</li> <li>Immediately following PCA administration tubing change</li> </ul>
[] PCA Documentation	- Immediately following PCA administration tubing change Routine, Every 12 hours
[] PCA Documentation	<ul> <li>Immediately following PCA administration tubing change</li> <li>Routine, Every 12 hours</li> <li>At the beginning (or end of each shift), prior to clearing PCA pump data,</li> </ul>
[] PCA Documentation	<ul> <li>Immediately following PCA administration tubing change</li> <li>Routine, Every 12 hours</li> <li>At the beginning (or end of each shift), prior to clearing PCA pump data, the following must be documented: doses delivered, number of attempts total amount of medication infused (in mg or mcg), and volume</li> </ul>
<ul><li>[] PCA Documentation</li><li>[] Patient education Pain pump</li></ul>	<ul> <li>Immediately following PCA administration tubing change</li> <li>Routine, Every 12 hours</li> <li>At the beginning (or end of each shift), prior to clearing PCA pump data, the following must be documented: doses delivered, number of attempts total amount of medication infused (in mg or mcg), and volume remaining in syringe (residual volume).</li> <li>Routine, Once, Starting S For 1 Occurrences</li> </ul>
	<ul> <li>Immediately following PCA administration tubing change</li> <li>Routine, Every 12 hours</li> <li>At the beginning (or end of each shift), prior to clearing PCA pump data, the following must be documented: doses delivered, number of attempts total amount of medication infused (in mg or mcg), and volume remaining in syringe (residual volume).</li> </ul>
	<ul> <li>Immediately following PCA administration tubing change</li> <li>Routine, Every 12 hours</li> <li>At the beginning (or end of each shift), prior to clearing PCA pump data, the following must be documented: doses delivered, number of attempts total amount of medication infused (in mg or mcg), and volume remaining in syringe (residual volume).</li> <li>Routine, Once, Starting S For 1 Occurrences Patient/Family:</li> </ul>
	<ul> <li>Immediately following PCA administration tubing change</li> <li>Routine, Every 12 hours</li> <li>At the beginning (or end of each shift), prior to clearing PCA pump data, the following must be documented: doses delivered, number of attempts total amount of medication infused (in mg or mcg), and volume remaining in syringe (residual volume).</li> <li>Routine, Once, Starting S For 1 Occurrences</li> <li>Patient/Family:</li> <li>Education for: Pain pump</li> <li>Provide patient education on appropriate use of PCA including no PCA</li> </ul>
[] Patient education Pain pump	<ul> <li>Immediately following PCA administration tubing change</li> <li>Routine, Every 12 hours</li> <li>At the beginning (or end of each shift), prior to clearing PCA pump data, the following must be documented: doses delivered, number of attempts total amount of medication infused (in mg or mcg), and volume remaining in syringe (residual volume).</li> <li>Routine, Once, Starting S For 1 Occurrences</li> <li>Patient/Family:</li> <li>Education for: Pain pump</li> <li>Provide patient education on appropriate use of PCA including no PCA by proxy. Only the patient may press the dosing button.</li> <li>Routine, Every 6 hours, Starting S</li> <li>Assess POSS while patient has an active PCA order. Contact provider if score 3 or 4.</li> <li>Routine, Until discontinued, Starting S, - PCA pump infusion discontinued for any reason</li> </ul>
<ul> <li>[] Patient education Pain pump</li> <li>[] Pasero Opioid-induced Sedation Scale</li> </ul>	<ul> <li>Immediately following PCA administration tubing change</li> <li>Routine, Every 12 hours</li> <li>At the beginning (or end of each shift), prior to clearing PCA pump data, the following must be documented: doses delivered, number of attempts total amount of medication infused (in mg or mcg), and volume remaining in syringe (residual volume).</li> <li>Routine, Once, Starting S For 1 Occurrences</li> <li>Patient/Family:</li> <li>Education for: Pain pump</li> <li>Provide patient education on appropriate use of PCA including no PCA by proxy. Only the patient may press the dosing button.</li> <li>Routine, Every 6 hours, Starting S</li> <li>Assess POSS while patient has an active PCA order. Contact provider if score 3 or 4.</li> <li>Routine, Until discontinued, Starting S, - PCA pump infusion discontinued for any reason</li> <li>Inadequate analgesia</li> </ul>
<ul> <li>[] Patient education Pain pump</li> <li>[] Pasero Opioid-induced Sedation Scale</li> </ul>	<ul> <li>Immediately following PCA administration tubing change</li> <li>Routine, Every 12 hours</li> <li>At the beginning (or end of each shift), prior to clearing PCA pump data, the following must be documented: doses delivered, number of attempts total amount of medication infused (in mg or mcg), and volume remaining in syringe (residual volume).</li> <li>Routine, Once, Starting S For 1 Occurrences</li> <li>Patient/Family:</li> <li>Education for: Pain pump</li> <li>Provide patient education on appropriate use of PCA including no PCA by proxy. Only the patient may press the dosing button.</li> <li>Routine, Every 6 hours, Starting S</li> <li>Assess POSS while patient has an active PCA order. Contact provider if score 3 or 4.</li> <li>Routine, Until discontinued, Starting S, - PCA pump infusion discontinued for any reason</li> <li>Inadequate analgesia</li> <li>Prior to administration of any other narcotics, antiemetics, or</li> </ul>
<ul> <li>[] Patient education Pain pump</li> <li>[] Pasero Opioid-induced Sedation Scale</li> </ul>	<ul> <li>Immediately following PCA administration tubing change</li> <li>Routine, Every 12 hours</li> <li>At the beginning (or end of each shift), prior to clearing PCA pump data, the following must be documented: doses delivered, number of attempts total amount of medication infused (in mg or mcg), and volume remaining in syringe (residual volume).</li> <li>Routine, Once, Starting S For 1 Occurrences</li> <li>Patient/Family:</li> <li>Education for: Pain pump</li> <li>Provide patient education on appropriate use of PCA including no PCA by proxy. Only the patient may press the dosing button.</li> <li>Routine, Every 6 hours, Starting S</li> <li>Assess POSS while patient has an active PCA order. Contact provider if score 3 or 4.</li> <li>Routine, Until discontinued, Starting S, - PCA pump infusion discontinued for any reason</li> <li>Inadequate analgesia</li> <li>Prior to administration of any other narcotics, antiemetics, or sedatives other than those ordered by the prescriber responsible for IV</li> </ul>
<ul> <li>[] Patient education Pain pump</li> <li>[] Pasero Opioid-induced Sedation Scale</li> </ul>	<ul> <li>Immediately following PCA administration tubing change</li> <li>Routine, Every 12 hours</li> <li>At the beginning (or end of each shift), prior to clearing PCA pump data, the following must be documented: doses delivered, number of attempts total amount of medication infused (in mg or mcg), and volume remaining in syringe (residual volume).</li> <li>Routine, Once, Starting S For 1 Occurrences</li> <li>Patient/Family:</li> <li>Education for: Pain pump</li> <li>Provide patient education on appropriate use of PCA including no PCA by proxy. Only the patient may press the dosing button.</li> <li>Routine, Every 6 hours, Starting S</li> <li>Assess POSS while patient has an active PCA order. Contact provider if score 3 or 4.</li> <li>Routine, Until discontinued, Starting S, - PCA pump infusion discontinued for any reason</li> <li>Inadequate analgesia</li> <li>Prior to administration of any other narcotics, antiemetics, or</li> </ul>

<ul> <li>Stop the PCA pump and call ordering physician and/or CERT team for any of the</li> </ul>	Routine, Until discontinued, Starting S, - Respiratory rate 10 per minute or less
following:	<ul> <li>Severe and/or recent confusion or disorientation</li> </ul>
	<ul> <li>POSS sedation level 4: Somnolent and difficult to arouse</li> </ul>
	- Sustained hypotension (SBP less than 90)
	- Excessive nausea or vomiting
	- Urinary retention
[] IV Fluids for provision of PCA Therapy (Single Response)	
() sodium chloride 0.9 % infusion	30 mL/hr, intravenous, continuous
() dextrose 5% infusion	30 mL/hr, intravenous, continuous
) morPHINE PCA 30 mg/30 mL + Nursing PCA Ord	ers
[] morPHINE PCA 30 mg/30 mL (Single Response	e)
() morPHINE PCA 30 mg/30 mL in sodium	Nurse Loading Dose: Not Ordered PCA Dose: 1 mg Lockout
chloride 0.9% for Opioid Naive	Interval: 10 Minutes MAX (Four hour dose limit): 20 mg
·	intravenous, continuous
	For breakthrough pain: Administer only if respiratory rate 12 per minute
	or more and POSS level of 2 or less. RN may bolus *** every *** hours
	as needed. If pain persists, may increase PCA demand dose by *** mg
	ONCE. If more than 2 bolus doses in 12 hours or if pain persists after
	increase in demand dose, call ordering prescriber.
	Adjust doses for age, renal function or other factors.
[] Nursing PCA Orders	
[] Vital signs - T/P/R/BP	Routine, Per unit protocol
	- Initially and every 30 minutes for 1 hour after PCA started, bolus
	administration or dose change; then
	- Every hour x 2 starting second hour after PCA started, bolus
	administered or dose change; then
	- Every 4 hours until PCA therapy is discontinued.
	- Immediately following PCA administration tubing change
[] PCA Documentation	Routine, Every 12 hours
	At the beginning (or end of each shift), prior to clearing PCA pump data,
	the following must be documented: doses delivered, number of attempts
	total amount of medication infused (in mg or mcg), and volume
	remaining in syringe (residual volume).
[] Patient education Pain pump	Routine, Once, Starting S For 1 Occurrences
	Patient/Family:
	Education for: Pain pump
	Provide patient education on appropriate use of PCA including no PCA
	by proxy. Only the patient may press the dosing button.
[] Pasero Opioid-induced Sedation Scale	Routine, Every 6 hours, Starting S
	Assess POSS while patient has an active PCA order. Contact provider if
	score 3 or 4.
[] Notify Physician	Routine, Until discontinued, Starting S, - PCA pump infusion
	discontinued for any reason
	- Inadequate analgesia
	- Prior to administration of any other narcotics, antiemetics, or
	sedatives other than those ordered by the prescriber responsible for IV
	PCA therapy
	<ul> <li>PCA pump discontinued by any service other than the prescriber</li> </ul>
	responsible for IV PCA therapy
[] Stop the PCA pump and call ordering	Routine, Until discontinued, Starting S, - Respiratory rate 10 per minute
physician and/or CERT team for any of the	or less
following:	- Severe and/or recent confusion or disorientation
	- POSS sedation level 4: Somnolent and difficult to arouse
	- Sustained hypotension (SBP less than 90)
	- Excessive nausea or vomiting
	- Urinary retention
[] IV Fluids for provision of PCA Therapy (Single Response)	
() sodium chloride 0.9 % infusion	30 mL/hr, intravenous, continuous
() dextrose 5% infusion	30 mL/hr, intravenous, continuous

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#### **Respiratory Depression and Somnolence**

[X] naloxone (NARCAN) injection	<ul> <li>0.2 mg, intravenous, once PRN, respiratory depression, as needed for respiratory rate 8 per minute or less OR patient somnolent and difficult to arouse (POSS GREATER than 3)., Post-op</li> <li>Repeat Naloxone 0.2 mg once in 2 minutes if necessary (MAXIMUM 0.4 mg).</li> <li>If naloxone is needed, please call the ordering physician and/or CERT</li> <li>team. Monitor vital signs (pulse oximetry, P/R/BP) every 15 minutes for 3 times.</li> </ul>

#### Antiemetics - HMSL, HMWB Only

.] ondansetron (ZOFRAN) IV or Oral (Selection Req	uired) "Or" Linked Panel
[X] ondansetron ODT (ZOFRAN-ODT)	4 mg, oral, every 8 hours PRN, nausea, vomiting, Post-op
disintegrating tablet	Give if patient is able to tolerate oral medication.
[X] ondansetron (ZOFRAN) 4 mg/2 mL injection	4 mg, intravenous, every 8 hours PRN, nausea, vomiting, Post-op
	Give if patient is UNable to tolerate oral medication OR if a faster onset of
	action is required.
[] promethazine (PHENERGAN) IV or Oral or Rectal	"Or" Linked Panel
[X] promethazine (PHENERGAN) injection	12.5 mg, intravenous, every 6 hours PRN, nausea, vomiting, PACU &
	Post-op
	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.
[X] promethazine (PHENERGAN) tablet	12.5 mg, oral, every 6 hours PRN, nausea, vomiting, Post-op
	Give if ondansetron (ZOFRAN) is ineffective and patient is able to tolerate oral medication.
[X] promethazine (PHENERGAN) suppository	12.5 mg, rectal, every 6 hours PRN, nausea, vomiting, Post-op Give if ondansetron (ZOFRAN) is ineffective and patient is UNable to tolerate oral medication.
	disintegrating tablet [X] ondansetron (ZOFRAN) 4 mg/2 mL injection (] promethazine (PHENERGAN) IV or Oral or Rectal [X] promethazine (PHENERGAN) injection [X] promethazine (PHENERGAN) tablet

#### Antiemetics - HMH, HMSJ, HMW, HMSTC Only

[X] ondansetron (ZOFRAN) IV or Oral (Selection Req	uired) "Or" Linked Panel	
[X] ondansetron ODT (ZOFRAN-ODT)	4 mg, oral, every 8 hours PRN, nausea, vomiting, Post-op	
disintegrating tablet	Give if patient is able to tolerate oral medication.	
[X] ondansetron (ZOFRAN) 4 mg/2 mL injection	4 mg, intravenous, every 8 hours PRN, nausea, vomiting, Post-op Give if patient is UNable to tolerate oral medication OR if a faster onset of action is required.	
[X] promethazine (PHENERGAN) IV or Oral or Recta	Or" Linked Panel	
[X] promethazine (PHENERGAN) 12.5 mg IV	12.5 mg, intravenous, every 6 hours PRN, nausea, vomiting, Post-op 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.	
[X] promethazine (PHENERGAN) tablet	12.5 mg, oral, every 6 hours PRN, nausea, vomiting, Post-op Give if ondansetron (ZOFRAN) is ineffective and patient is able to tolerate oral medication.	
[X] promethazine (PHENERGAN) suppository	12.5 mg, rectal, every 6 hours PRN, nausea, vomiting, Post-op Give if ondansetron (ZOFRAN) is ineffective and patient is UNable to tolerate oral medication.	

#### Antiemetics - HMSTJ Only

[X] ondansetron (ZOFRAN) IV or Oral (Selection Rec	uired) "Or" Linked Panel
[X] ondansetron ODT (ZOFRAN-ODT)	4 mg, oral, every 8 hours PRN, nausea, vomiting, Post-op
disintegrating tablet	Give if patient is able to tolerate oral medication.
[X] ondansetron (ZOFRAN) 4 mg/2 mL injection	4 mg, intravenous, every 8 hours PRN, nausea, vomiting, Post-op Give if patient is UNable to tolerate oral medication OR if a faster onset of action is required.
[X] promethazine (PHENERGAN) IVPB or Oral or Re	ectal "Or" Linked Panel

<ul> <li>[X] promethazine (PHENERGAN) 25 mg in sodium chloride 0.9 % 50 mL IVPB</li> </ul>	vomiting Give if o tolerate	, intravenous, for 30 Minutes, every 6 hours PRN, nausea, , Post-op ndansetron (ZOFRAN) is ineffective and patient is UNable to oral or rectal medication OR if a faster onset of action is required.
[X] promethazine (PHENERGAN) tablet	Give if o oral med	
[X] promethazine (PHENERGAN) suppository	Give if o	, rectal, every 6 hours PRN, nausea, vomiting, Post-op ndansetron (ZOFRAN) is ineffective and patient is UNable to oral medication.
Laxatives		
<ul> <li>[] sennosides-docusate sodium (SENOKOT-S) 8.6 per tablet</li> </ul>	-50 mg	2 tablet, oral, nightly PRN, constipation, Post-op
<ul> <li>[] magnesium hydroxide suspension - NOT RECOMMENDED FOR CHRONIC KIDNEY DISI STAGE 3 OR WORSE</li> </ul>	EASE	30 mL, oral, every 6 hours PRN, constipation, Post-op Do not give if patient is on hemodialysis or is in chronic renal failure.
[] bisacodyl (DULCOLAX) EC tablet		10 mg, oral, daily PRN, constipation, Post-op
[]         bisacodyl (DULCOLAX) suppository		10 mg, rectal, daily PRN, constipation, Post-op
[] polyethylene glycol (MIRALAX) packet		17 g, oral, daily PRN, constipation, Post-op
[] docusate sodium (COLACE) capsule		100 mg, oral, 2 times daily PRN, constipation, Post-op
Symptom Management		
[] pantoprazole (PROTONIX) EC tablet		40 mg, oral, daily at 0600, Post-op Indication(s) for Proton Pump Inhibitor (PPI) Therapy:
[] acetaminophen (TYLENOL) tablet		650 mg, oral, every 6 hours PRN, headaches, fever, Temperature GREATER than 101 F., Post-op
[] alum-mag hydroxide-simeth (MAALOX) 200-200- mL suspension		30 mL, oral, every 6 hours PRN, indigestion, heartburn, Post-op
[] magnesium hydroxide OR magnesium citrate (Si Response)		
() magnesium hydroxide suspension - NOT RECOMMENDED FOR CHRONIC KIDNEY DISEASE STAGE 3 OR WORSE		oral, every 6 hours PRN, constipation, Post-op ommended for Chronic Kidney Disease Stage 3 Or Worse
<ul> <li>magnesium citrate solution - NOT RECOMMENDED FOR CHRONIC KIDNEY DISEASE STAGE 3 OR WORSE</li> </ul>		oral, once PRN, constipation, Post-op ommended for Chronic Kidney Disease Stage 3 Or Worse
Itching: For Patients LESS than 70 years old (Sin	gle Respo	nse)
() diphenhydrAMINE (BENADRYL) tablet		25 mg, oral, every 6 hours PRN, itching, Post-op
() hydrOXYzine (ATARAX) tablet		10 mg, oral, every 6 hours PRN, itching, Post-op
() cetirizine (ZyrTEC) tablet		5 mg, oral, daily PRN, itching, Post-op
<ol> <li>fexofenadine (ALLEGRA) tablet - For eGFR LES 80 mL/min, reduce frequency to once daily as ne</li> </ol>		60 mg, oral, 2 times daily PRN, itching, Post-op
Itching: For Patients between 70-76 years old (Si	ngle Resp	onse)
() cetirizine (ZyrTEC) tablet		5 mg, oral, daily PRN, itching, Post-op
Itching: For Patients GREATER than 77 years old	l (Single R	esponse)
() cetirizine (ZyrTEC) tablet		5 mg, oral, daily PRN, itching, Post-op
Insomnia: For Patients LESS than 70 years old (S	Single Res	ponse)
() zolpidem (AMBIEN) tablet		5 mg, oral, nightly PRN, sleep, Post-op
() ramelteon (ROZEREM) tablet		8 mg, oral, nightly PRN, sleep, Post-op
Insomnia: For Patients GREATER than or EQUAL	to 70 yea	rs old (Single Response)
() ramelteon (ROZEREM) tablet		8 mg, oral, nightly PRN, sleep, Post-op
VTE		

DVT Risk and Prophylaxis Tool (Single Response VTE/DVT Risk Definitions	e) (Selection Required) URL:
Anticoagulation Guide for COVID patients	"\\appt1\epicappprod\Restricted\OrderSets\VTEDVTRISK DEFINITIONS.pdf" URL: "https://formweb.com/files/houstonmethodist/documents/C OVID-19 Anticoagulation Guideline - 8.20.2021v15.pdf"
() Patient currently has an active order for therapeu anticoagulant or VTE prophylaxis with Risk Strati (Single Response) (Selection Required)	
() Moderate Risk - Patient currently has an active	e order for
therapeutic anticoagulant or VTE prophylaxis ( Required)	Selection
[] Moderate risk of VTE	Routine, Once, PACU & Post-op
<ul> <li>Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis</li> </ul>	Routine, Once No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication. Therapy for the following: PACU & Post-op
[] Place sequential compression device (Single	
<ul> <li>Contraindications exist for mechanical prophylaxis</li> </ul>	Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s): PACU & Post-op
<ul> <li>Place/Maintain sequential compression device continuous</li> </ul>	Routine, Continuous, PACU & Post-op
() Moderate Risk - Patient currently has an active therapeutic anticoagulant or VTE prophylaxis ( Required)	Selection
[] Moderate risk of VTE	Routine, Once, PACU & Post-op
<ul> <li>Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis</li> </ul>	Routine, Once No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication. Therapy for the following: PACU & Post-op
[] Place sequential compression device (Single	
<ul> <li>() Contraindications exist for mechanical prophylaxis</li> </ul>	Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s): PACU & Post-op
() Place/Maintain sequential compression device continuous	Routine, Continuous, PACU & Post-op
<ul> <li>High Risk - Patient currently has an active order therapeutic anticoagulant or VTE prophylaxis ( Required)</li> </ul>	
[] High risk of VTE	Routine, Once, PACU & Post-op
[] Patient currently has an active order for	Routine, Once
therapeutic anticoagulant or VTE prophylaxis	No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication. Therapy for the following: PACU & Post-op
[] Place sequential compression device (Single	
() Contraindications exist for mechanical prophylaxis	Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s): PACU & Post-op
() Place/Maintain sequential compression device continuous	Routine, Continuous, PACU & Post-op
() High Risk - Patient currently has an active order therapeutic anticoagulant or VTE prophylaxis ( Required)	
[] High risk of VTE	Routine, Once, PACU & Post-op

[] Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis	Routine, Once No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication. Therapy for the following: PACU & Post-op
[] Place sequential compression device (Single F	
() Contraindications exist for mechanical prophylaxis	Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s): PACU & Post-op
() Place/Maintain sequential compression device continuous	Routine, Continuous, PACU & Post-op
() LOW Risk of DVT (Selection Required)	
Low Risk Definition Age less than 60 years and NO other VTE risk fac	tors
[] Low Risk (Single Response) (Selection Require	d)
() Low risk of VTE	Routine, Once Low risk: Due to low risk, no VTE prophylaxis is needed. Will encourgae early ambulation PACU & Post-op
MODERATE Risk of DVT - Surgical (Selection Re	
Moderate Risk Definition	echanical prophylaxis is optional unless pharmacologic is
stroke, rheumatologic disease, sickle cell disease, Age 60 and above Central line History of DVT or family history of VTE Anticipated length of stay GREATER than 48 hour Less than fully and independently ambulatory Estrogen therapy Moderate or major surgery (not for cancer) Major surgery within 3 months of admission	leg swelling, ulcers, venous stasis and nephrotic syndrome
[] Moderate Risk (Selection Required)	
[] Moderate risk of VTE [] Moderate Risk Pharmacological Prophylaxis - S	Routine, Once, PACU & Post-op
Patient (Single Response) (Selection Required)	•
() Contraindications exist for pharmacologic prop BUT order Sequential compression device	hylaxis "And" Linked Panel
<ul> <li>Contraindications exist for pharmacologic prophylaxis</li> </ul>	Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op
[] Place/Maintain sequential compression device continuous	Routine, Continuous, PACU & Post-op
<ul> <li>Contraindications exist for pharmacologic prop AND mechanical prophylaxis</li> </ul>	hylaxis "And" Linked Panel
<ul> <li>Contraindications exist for pharmacologic prophylaxis</li> </ul>	Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op
<ul> <li>Contraindications exist for mechanical prophylaxis</li> </ul>	Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s): PACU & Post-op
() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)	PACU & Post-op

(Selection Required) Printed on 9/6/2022 at 2:23 PM from POC environment

()	enoxaparin (LOVENOX) syringe	40 mg, subcutaneous, daily at 0600, Starting S+1, PACU & Post-op Indication(s): VTE Prophylaxis			
()	) patients with CrCL LESS than 30 mL/min	30 mg, subcutaneous, daily at 0600, Starting S+1, PACU & Post-op For Patients with CrCL LESS than 30 mL/min Indication(s): VTE Prophylaxis			
()	patients weight between 100-139 kg AND CrCl GREATER than 30 mL/min	30 mg, subcutaneous, 2 times daily at 0600, 1800, Starting S+1, PAC & Post-op For Patients weight between 100-139 kg and CrCl GREATER than 30			
		mL/min Indication(s): VTE Prophylaxis			
()	patients weight 140 kg or GREATER AND CrCl GREATER than 30 mL/min	40 mg, subcutaneous, 2 times daily at 0600, 1800, Starting S+1, PAC & Post-op For Patients weight 140 kg or GREATER and CrCl GREATER than 30			
		mL/min Indication(s): VTE Prophylaxis			
()	fondaparinux (ARIXTRA) injection	2.5 mg, subcutaneous, daily, Starting S+1, PACU & Post-op If the patient does not have a history of or suspected case of			
		Heparin-Induced Thrombocytopenia (HIT) do NOT order this medicatic Contraindicated in patients LESS than 50kg, prior to surgery/invasive procedure, or CrCI LESS than 30 mL/min.			
		This patient has a history of or suspected case of Heparin-Induced Thrombocytopenia (HIT):			
	heparin (porcine) injection	5,000 Units, subcutaneous, every 8 hours, S+1 at 6:00 AM, PACU & Post-op			
()	heparin (porcine) injection (Recommended for patients with high risk of bleeding, e.g. weight < 50kg and age > 75yrs)	5,000 Units, subcutaneous, every 12 hours, S+1 at 6:00 AM, PACU & Post-op Recommended for patients with high risk of bleeding, e.g. weight LESS			
$\overline{()}$	HEParin (porcine) injection - For Patients	than 50kg and age GREATER than 75yrs. 7,500 Units, subcutaneous, every 8 hours, S+1 at 6:00 AM, PACU &			
()	with weight GREATER than 100 kg	Post-op For patients with weight GREATER than 100 kg.			
()	warfarin (COUMADIN) tablet	oral, daily at 1700, Starting S+1, PACU & Post-op Indication:			
()	Pharmacy consult to manage warfarin (COUMADIN)	STAT, Until discontinued, Starting S Indication:			
Ē	Mechanical Prophylaxis (Single Response) (Sel Required)				
()	Contraindications exist for mechanical prophylaxis	Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s PACU & Post-op			
()	Place/Maintain sequential compression device continuous	Routine, Continuous, PACU & Post-op			
Re	DERATE Risk of DVT - Non-Surgical (Selectio quired)	n			
Pha	derate Risk Definition armacologic prophylaxis must be addressed. M ntraindicated.	echanical prophylaxis is optional unless pharmacologic is			
СН		nation, dehydration, varicose veins, cancer, sepsis, obesity, previous leg swelling, ulcers, venous stasis and nephrotic syndrome			
Age Cei	e 60 and above ntral line				
Ant	tory of DVT or family history of VTE ticipated length of stay GREATER than 48 hour as than fully and independently ambulatory	s			
Est	rogen therapy derate or major surgery (not for cancer)				
	jor surgery within 3 months of admission				
1 1	Moderate Risk (Selection Required)				

hylaxis - "And" Linked Panel
Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op
Routine, Continuous, PACU & Post-op
hylaxis "And" Linked Panel
Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op
Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s): PACU & Post-op
oonse)
40 mg, subcutaneous, daily at 1700, Starting S+1, PACU & Post-op Indication(s): VTE Prophylaxis
30 mg, subcutaneous, daily at 1700, Starting S+1, PACU & Post-op For Patients with CrCL LESS than 30 mL/min Indication(s): VTE Prophylaxis
30 mg, subcutaneous, every 12 hours at 0900, 2100, Starting S+1, PACU & Post-op For Patients weight between 100-139 kg and CrCI GREATER than 30 mL/min Indication(s): VTE Prophylaxis
40 mg, subcutaneous, every 12 hours at 0900, 2100, Starting S+1, PACU & Post-op For Patients weight 140 kg or GREATER and CrCI GREATER than 30 mL/min Indication(s): VTE Prophylaxis
2.5 mg, subcutaneous, daily, PACU & Post-op 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 CrCI LESS than 30 mL/min This patient has a history of or suspected case of Heparin-Induced Thrombocytopenia (HIT):
5,000 Units, subcutaneous, every 8 hours, PACU & Post-op
5,000 Units, subcutaneous, every 12 hours, PACU & Post-op Recommended for patients with high risk of bleeding, e.g. weight LESS than 50kg and age GREATER than 75yrs.
7,500 Units, subcutaneous, every 8 hours, PACU & Post-op For patients with weight GREATER than 100 kg.
oral, daily at 1700, PACU & Post-op Indication:
STAT, Until discontinued, Starting S Indication:
lection
Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s) PACU & Post-op

() HIGH Risk of DVT - Surgical (Selection Required	)
High Risk Definition	a must be addressed
Both pharmacologic AND mechanical prophylaxis One or more of the following medical conditions:	s must be addressed.
Thrombophilia (Factor V Leiden, prothrombin vari or protein S deficiency; hyperhomocysteinemia; n Severe fracture of hip, pelvis or leg	iant mutations, anticardiolipin antibody syndrome; antithrombin, protein C nyeloproliferative disorders)
Acute spinal cord injury with paresis	
Multiple major traumas Abdominal or pelvic surgery for CANCER	
Acute ischemic stroke	
History of PE	
[] High Risk (Selection Required)	
[] High risk of VTE	Pouting Onco
	Routine, Once
[] High Risk Pharmacological Prophylaxis - Surgi (Single Response) (Selection Required)	
() Contraindications exist for pharmacologic	Routine, Once
prophylaxis	No pharmacologic VTE prophylaxis due to the following contraindication(s):
() enoxaparin for VTE Prophylaxis (Single Resp	
	30 mg, subcutaneous, daily at 1700
() enoxaparin (LOVENOX) 30 mg daily at 1700	Indication(s): VTE Prophylaxis
() enoxaparin (LOVENOX) 30 mg every 12	30 mg, subcutaneous, every 12 hours
hours	Indication(s): VTE Prophylaxis
() enoxaparin (LOVENOX) 40 mg daily at	40 mg, subcutaneous, daily at 1700
1700	Indication(s): VTE Prophylaxis
() enoxaparin (LOVENOX) 40 mg every 12 hours	40 mg, subcutaneous, every 12 hours Indication(s): VTE Prophylaxis
() fondaparinux (ARIXTRA) injection	2.5 mg, subcutaneous, daily, Starting 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 CrCI LESS than 30 mL/min.
	This patient has a history of or suspected case of Heparin-Induced
() honorin (norcina) injection	Thrombocytopenia (HIT):
() heparin (porcine) injection	5,000 Units, subcutaneous, every 8 hours, S+1 at 6:00 AM
() heparin (porcine) injection (Recommended for patients with high risk of bleeding, e.g.	5,000 Units, subcutaneous, every 12 hours, S+1 at 6:00 AM Recommended for patients with high risk of bleeding, e.g. weight LESS
weight < 50kg and age > 75yrs)	than 50kg and age GREATER than 75yrs.
() HEParin (porcine) injection - For Patients	7,500 Units, subcutaneous, every 8 hours, Starting S+1
with weight GREATER than 100 kg	For patients with weight GREATER than 100 kg.
() warfarin (COUMADIN) tablet	oral, daily at 1700, Starting S+1
	Indication:
() Pharmacy consult to manage warfarin (COUMADIN)	STAT, Until discontinued, Starting S Indication:
[] Mechanical Prophylaxis (Single Response) (Se Required)	
() Contraindications exist for mechanical	Routine, Once
prophylaxis	No mechanical VTE prophylaxis due to the following contraindication(s):
() Place/Maintain sequential compression	Routine, Continuous
device continuous	·
() HIGH Risk of DVT - Non-Surgical (Selection Req	uired)

High Risk Definition Both pharmacologic AND mechanical prophylaxis	must be addressed.
One or more of the following medical conditions:	
	nt mutations, anticardiolipin antibody syndrome; antithrombin, protein C
or protein S deficiency; hyperhomocysteinemia; my	veloproliferative disorders)
Severe fracture of hip, pelvis or leg	
Acute spinal cord injury with paresis	
Multiple major traumas	
Abdominal or pelvic surgery for CANCER	
Acute ischemic stroke History of PE	
[] High Risk (Selection Required)	
[] High risk of VTE	Routine, Once, PACU & Post-op
[] High Risk Pharmacological Prophylaxis - Non-S Patient (Single Response) (Selection Required)	urgical
() Contraindications exist for pharmacologic	Routine, Once
prophylaxis	No pharmacologic VTE prophylaxis due to the following
	contraindication(s):
	PACU & Post-op
<ul> <li>enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> </ul>	· · · · · · · · · · · · · · · · · · ·
() enoxaparin (LOVENOX) syringe	40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op Indication(s): VTE Prophylaxis
() patients with CrCL LESS than 30 mL/min	30 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op
	For Patients with CrCL LESS than 30 mL/min
	Indication(s): VTE Prophylaxis
() patients weight between 100-139 kg AND	30 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op
CrCl GREATER than 30 mL/min	For Patients weight between 100-139 kg and CrCl GREATER than 30
	mL/min
() notionto weight 140 kg or ODEATED AND	Indication(s): VTE Prophylaxis
<ul> <li>() patients weight 140 kg or GREATER AND CrCl GREATER than 30 mL/min</li> </ul>	40 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op For Patients weight 140 kg or GREATER and CrCl GREATER than 30
CICI GREATER INAN 30 ML/MM	mL/min
	Indication(s): VTE Prophylaxis
() fondaparinux (ARIXTRA) injection	2.5 mg, subcutaneous, daily, PACU & Post-op
	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 CrCI LESS than 30 mL/min.
	This patient has a history of or suspected case of Heparin-Induced
	Thrombocytopenia (HIT):
() heparin (porcine) injection	5,000 Units, subcutaneous, every 8 hours, PACU & Post-op
() heparin (porcine) injection (Recommended	5,000 Units, subcutaneous, every 12 hours, PACU & Post-op
for patients with high risk of bleeding, e.g.	Recommended for patients with high risk of bleeding, e.g. weight LESS
weight < 50kg and age > 75yrs)	than 50kg and age GREATER than 75yrs.
<ul> <li>HEParin (porcine) injection - For Patients with weight GREATER than 100 kg</li> </ul>	7,500 Units, subcutaneous, every 8 hours, PACU & Post-op For patients with weight GREATER than 100 kg.
() warfarin (COUMADIN) tablet	oral, daily at 1700, PACU & Post-op Indication:
() Pharmacy consult to manage warfarin (COUMADIN)	STAT, Until discontinued, Starting S Indication:
[] Mechanical Prophylaxis (Single Response) (Sel Required)	ection
() Contraindications exist for mechanical prophylaxis	Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s):
	PACU & Post-op
() Place/Maintain sequential compression device continuous	Routine, Continuous, PACU & Post-op
<ul> <li>HIGH Risk of DVT - Surgical (Hip/Knee) (Selection Required)</li> </ul>	1

High Risk Definition Both pharmacologic AND mechanical prophylaxis One or more of the following medical conditions: Thrombophilia (Factor V Leiden, prothrombin vari or protein S deficiency; hyperhomocysteinemia; n Severe fracture of hip, pelvis or leg Acute spinal cord injury with paresis Multiple major traumas Abdominal or pelvic surgery for CANCER Acute ischemic stroke History of PE	ant mutations, anticardiolipin antibody syndrome; antithrombin, protein C
[1] Llick Diele (Colorition Derwined)	
[] High Risk (Selection Required) [] High risk of VTE	Routine, Once, PACU & Post-op
<ul> <li>[] High Risk Pharmacological Prophylaxis - Hip o (Arthroplasty) Surgical Patient (Single Respons (Selection Required)</li> </ul>	r Knee
() Contraindications exist for pharmacologic prophylaxis	Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op
() aspirin chewable tablet	162 mg, oral, daily, Starting S+1, PACU & Post-op
() aspirin (ECOTRIN) enteric coated tablet	162 mg, oral, daily, Starting S+1, PACU & Post-op
() Apixaban and Pharmacy Consult (Selection R	
[] apixaban (ELIQUIS) tablet	2.5 mg, oral, 2 times daily, Starting S+1, PACU & Post-op Indications: VTE prophylaxis
[] Pharmacy consult to monitor apixaban (ELIQUIS) therapy	STAT, Until discontinued, Starting S Indications: VTE prophylaxis
<ul> <li>() enoxaparin (LOVENOX) injection (Single Res (Selection Required)</li> <li>() enoxaparin (LOVENOX) syringe</li> </ul>	40 mg, subcutaneous, daily at 0600, Starting S+1, PACU & Post-op
() enoxaparin (LOVENOX) syringe	Indication(s): VTE Prophylaxis 30 mg, subcutaneous, 2 times daily at 0600, 1800, Starting S+1, PACU & Post-op
() enoxaparin (LOVENOX) syringe - For Patients with CrCL LESS than 30 mL/min	Indication(s): VTE Prophylaxis 30 mg, subcutaneous, daily at 0600, Starting S+1, PACU & Post-op For Patients with CrCL LESS than 30 mL/min. Indication(s): VTE Prophylaxis
() enoxaparin (LOVENOX) syringe - For Patients weight between 100-139 kg and CrCl GREATER than 30 mL/min	30 mg, subcutaneous, 2 times daily at 0600, 1800, Starting S+1, PACU & Post-op For Patients weight between 100-139 kg and CrCl GREATER than 30 mL/min. Indication(s): VTE Prophylaxis
<ul> <li>enoxaparin (LOVENOX) syringe - For Patients weight between 140 kg or GREATER and CrCI GREATER than 30 mL/min</li> </ul>	40 mg, subcutaneous, 2 times daily at 0600, 1800, Starting S+1, PACU & Post-op For Patients weight 140 kg or GREATER and CrCI GREATER than 30 mL/min Indication(s): VTE Prophylaxis
() fondaparinux (ARIXTRA) injection	<ul> <li>2.5 mg, subcutaneous, daily, Starting S+1, PACU &amp; Post-op</li> <li>If the patient does not have a history or suspected case of</li> <li>Heparin-Induced Thrombocytopenia (HIT) do NOT order this medication.</li> <li>Contraindicated in patients LESS than 50kg, prior to surgery/invasive</li> <li>procedure, or CrCl LESS than 30 mL/min</li> <li>This patient has a history of or suspected case of Heparin-Induced</li> <li>Thrombocytopenia (HIT):</li> </ul>
() heparin (porcine) injection	5,000 Units, subcutaneous, every 8 hours, S+1 at 6:00 AM, PACU & Post-op
<ul> <li>() heparin (porcine) injection (Recommended for patients with high risk of bleeding, e.g. weight &lt; 50kg and age &gt; 75yrs)</li> </ul>	5,000 Units, subcutaneous, every 12 hours, S+1 at 6:00 AM, PACU & Post-op Recommended for patients with high risk of bleeding, e.g. weight LESS than 50kg and age GREATER than 75yrs.

() HEParin (porcine) injection - For Patients	
with weight GREATER than 100 kg	7,500 Units, subcutaneous, every 8 hours, S+1 at 6:00 AM, PACU & Post-op For patients with weight GREATER than 100 kg.
() Rivaroxaban and Pharmacy Consult (Selection	· · · · · · · · · · · · · · · · · · ·
Required)	
<ul> <li>rivaroxaban (XARELTO) tablet for hip or knee arthroplasty planned during this admission</li> </ul>	10 mg, oral, daily at 0600 (TIME CRITICAL), PACU & Post-op Indications: VTE prophylaxis
[] Pharmacy consult to monitor rivaroxaban (XARELTO) therapy	STAT, Until discontinued, Starting S Indications: VTE prophylaxis
() warfarin (COUMADIN) tablet	oral, daily at 1700, Starting S+1, PACU & Post-op Indication:
() Pharmacy consult to manage warfarin (COUMADIN)	STAT, Until discontinued, Starting S Indication:
[] Mechanical Prophylaxis (Single Response) (Se Required)	
() Contraindications exist for mechanical prophylaxis	Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s) PACU & Post-op
<ul> <li>Place/Maintain sequential compression device continuous</li> </ul>	Routine, Continuous, PACU & Post-op
VT Risk and Prophylaxis Tool (Single Response	)
VTE/DVT Risk Definitions	URL: "\\appt1\epicappprod\Restricted\OrderSets\VTEDVTRISK DEFINITIONS.pdf"
Anticoagulation Guide for COVID patients	URL: "https://formweb.com/files/houstonmethodist/documents/C OVID-19 Anticoagulation Guideline - 8.20.2021v15.pdf"
<ul> <li>(Single Response) (Selection Required)</li> <li>Moderate Risk - Patient currently has an active therapeutic anticoagulant or VTE prophylaxis (</li> </ul>	
Poduirod)	
Required)	Routine Once PACU & Post-op
<ul> <li>Required)</li> <li>Moderate risk of VTE</li> <li>Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis</li> </ul>	Routine, Once, PACU & Post-op Routine, Once No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication. Therapy for the following: PACU & Post-op
<ul> <li>Moderate risk of VTE</li> <li>Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis</li> <li>Place sequential compression device (Single</li> </ul>	Routine, Once No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication. Therapy for the following: PACU & Post-op
<ul> <li>Moderate risk of VTE</li> <li>Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis</li> </ul>	Routine, Once No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication. Therapy for the following: PACU & Post-op Response) Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s):
<ul> <li>[] Moderate risk of VTE</li> <li>[] Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis</li> <li>[] Place sequential compression device (Single () Contraindications exist for mechanical</li> </ul>	Routine, Once No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication. Therapy for the following: PACU & Post-op Response) Routine, Once No mechanical VTE prophylaxis due to the following
<ul> <li>Moderate risk of VTE</li> <li>Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis</li> <li>Place sequential compression device (Single () Contraindications exist for mechanical prophylaxis</li> <li>Place/Maintain sequential compression</li> </ul>	Routine, Once No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication. Therapy for the following: PACU & Post-op Response) Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s): PACU & Post-op Routine, Continuous, PACU & Post-op
<ul> <li>Moderate risk of VTE</li> <li>Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis</li> <li>Place sequential compression device (Single         <ul> <li>() Contraindications exist for mechanical prophylaxis</li> <li>() Place/Maintain sequential compression device continuous</li> <li>() Moderate Risk - Patient currently has an active therapeutic anticoagulant or VTE prophylaxis ( Required)</li> <li>[] Moderate risk of VTE</li> </ul> </li> </ul>	Routine, Once         No pharmacologic VTE prophylaxis because: patient is already on         therapeutic anticoagulation for other indication.         Therapy for the following:         PACU & Post-op         Response)         Routine, Once         No mechanical VTE prophylaxis due to the following         contraindication(s):         PACU & Post-op         Routine, Continuous, PACU & Post-op         e order for         Selection         Routine, Once, PACU & Post-op
<ul> <li>Moderate risk of VTE</li> <li>Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis</li> <li>Place sequential compression device (Single         <ul> <li>() Contraindications exist for mechanical prophylaxis</li> <li>() Place/Maintain sequential compression device continuous</li> <li>() Moderate Risk - Patient currently has an active therapeutic anticoagulant or VTE prophylaxis ( Required)</li> </ul> </li> </ul>	Routine, Once       No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication.         Therapy for the following:       PACU & Post-op         Response)       Routine, Once         No mechanical VTE prophylaxis due to the following contraindication(s):       PACU & Post-op         Routine, Continuous, PACU & Post-op       Routine, Continuous, PACU & Post-op         Routine, Once, PACU & Post-op       Routine, Once         No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication.         Therapy for the following:
<ul> <li>Moderate risk of VTE</li> <li>Patient currently has an active order for therapeutic anticoagulant or VTE prophylaxis</li> <li>Place sequential compression device (Single         <ul> <li>() Contraindications exist for mechanical prophylaxis</li> <li>() Place/Maintain sequential compression device continuous</li> <li>() Moderate Risk - Patient currently has an active therapeutic anticoagulant or VTE prophylaxis ( Required)</li> <li>[] Moderate risk of VTE</li> <li>[] Patient currently has an active order for therapeutic anticoagulant or VTE</li> </ul> </li> </ul>	Routine, Once       No pharmacologic VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication.         Therapy for the following:       PACU & Post-op         Response)       Routine, Once         No mechanical VTE prophylaxis due to the following contraindication(s):       PACU & Post-op         Routine, Continuous, PACU & Post-op       Routine, Continuous, PACU & Post-op         Routine, Once, No mechanical VTE prophylaxis because: patient is already on therapeutic anticoagulation for other indication.         Therapy for the following:         PACU & Post-op

() Place/Maintain sequential compression device continuous	Routine, Continuous, PACU & Post-op
() High Risk - Patient currently has an active order therapeutic anticoagulant or VTE prophylaxis (	
Required) [] High risk of VTE	Routine, Once, PACU & Post-op
<ul> <li>Patient currently has an active order for</li> </ul>	Routine, Once
therapeutic anticoagulant or VTE	No pharmacologic VTE prophylaxis because: patient is already on
prophylaxis	therapeutic anticoagulation for other indication.
propristante	Therapy for the following:
	PACU & Post-op
[] Place sequential compression device (Single	
<ul> <li>Contraindications exist for mechanical prophylaxis</li> </ul>	Routine, Once
	No mechanical VTE prophylaxis due to the following
	contraindication(s):
	PACU & Post-op
<ul> <li>Place/Maintain sequential compression device continuous</li> </ul>	Routine, Continuous, PACU & Post-op
<ul> <li>High Risk - Patient currently has an active order therapeutic anticoagulant or VTE prophylaxis ( Required)</li> </ul>	
[] High risk of VTE	Routine, Once, PACU & Post-op
<ul> <li>Patient currently has an active order for</li> </ul>	Routine, Once
therapeutic anticoagulant or VTE	No pharmacologic VTE prophylaxis because: patient is already on
prophylaxis	therapeutic anticoagulation for other indication.
	Therapy for the following:
	PACU & Post-op
[] Place sequential compression device (Single	Response)
() Contraindications exist for mechanical	Routine, Once
prophylaxis	No mechanical VTE prophylaxis due to the following
	contraindication(s):
	PACU & Post-op
<ul> <li>Place/Maintain sequential compression device continuous</li> </ul>	Routine, Continuous, PACU & Post-op
) LOW Risk of DVT (Selection Required)	
Low Risk Definition	
Age less than 60 years and NO other VTE risk fa	
[] Low Risk (Single Response) (Selection Requir	
() Low risk of VTE	Routine, Once
	Low risk: Due to low risk, no VTE prophylaxis is needed. Will encourga
	early ambulation
MODERATE Risk of DV/T Surgical (Selection R	PACU & Post-op
MODERATE Risk of DVT - Surgical (Selection Re	
Moderate Risk Definition Pharmacologic prophylaxis must be addressed. N contraindicated.	lechanical prophylaxis is optional unless pharmacologic is
One or more of the following medical conditions:	
	mation, dehydration, varicose veins, cancer, sepsis, obesity, previous
	e, leg swelling, ulcers, venous stasis and nephrotic syndrome
Age 60 and above	
Central line	
History of DVT or family history of VTE	
Anticipated length of stay GREATER than 48 hou	ırs
Less than fully and independently ambulatory	
Estrogen therapy	
Moderate or major surgery (not for cancer)	
Major surgery within 3 months of admission	
[] Moderate Risk (Selection Required)	

[] Moderate risk of VTE	Routine, Once, PACU & Post-op
] Moderate Risk Pharmacological Prophylaxis - Su Patient (Single Response) (Selection Required)	urgical
<ul> <li>() Contraindications exist for pharmacologic propl BUT order Sequential compression device</li> </ul>	hylaxis "And" Linked Panel
[] Contraindications exist for pharmacologic	Routine, Once
prophylaxis	No pharmacologic VTE prophylaxis due to the following
	contraindication(s):
	PACU & Post-op
[] Place/Maintain sequential compression device continuous	Routine, Continuous, PACU & Post-op
() Contraindications exist for pharmacologic propl AND mechanical prophylaxis	hylaxis "And" Linked Panel
[] Contraindications exist for pharmacologic	Routine, Once
prophylaxis	No pharmacologic VTE prophylaxis due to the following
	contraindication(s):
	PACU & Post-op
[] Contraindications exist for mechanical	Routine, Once
prophylaxis	No mechanical VTE prophylaxis due to the following
	contraindication(s): PACU & Post-op
() enoxaparin (LOVENOX) injection (Single Resp. (Selection Required)	
() enoxaparin (LOVENOX) syringe	40 mg, subcutaneous, daily at 0600, Starting S+1, PACU & Post-op
() choxapann (EOVENOX) synnge	Indication(s): VTE Prophylaxis
() patients with CrCL LESS than 30 mL/min	30 mg, subcutaneous, daily at 0600, Starting S+1, PACU & Post-op
	For Patients with CrCL LESS than 30 mL/min
	Indication(s): VTE Prophylaxis
() patients weight between 100-139 kg AND	30 mg, subcutaneous, 2 times daily at 0600, 1800, Starting S+1, PACL
CrCl GREATER than 30 mL/min	& Post-op
	For Patients weight between 100-139 kg and CrCI GREATER than 30
	Indication(s): VTE Prophylaxis
() patients weight 140 kg or GREATER AND CrCl GREATER than 30 mL/min	40 mg, subcutaneous, 2 times daily at 0600, 1800, Starting S+1, PACL
CICI GREATER than 50 mL/mm	& Post-op For Patients weight 140 kg or GREATER and CrCl GREATER than 30
	mL/min
	Indication(s): VTE Prophylaxis
() fondaparinux (ARIXTRA) injection	2.5 mg, subcutaneous, daily, Starting S+1, PACU & Post-op
	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 CrCI LESS than 30 mL/min.
	This patient has a history of or suspected case of Heparin-Induced
	Thrombocytopenia (HIT):
() heparin (porcine) injection	5,000 Units, subcutaneous, every 8 hours, S+1 at 6:00 AM, PACU & Post-op
() heparin (porcine) injection (Recommended	5,000 Units, subcutaneous, every 12 hours, S+1 at 6:00 AM, PACU &
for patients with high risk of bleeding, e.g.	Post-op
weight < 50kg and age > 75yrs)	Recommended for patients with high risk of bleeding, e.g. weight LESS
	than 50kg and age GREATER than 75yrs.
() HEParin (porcine) injection - For Patients	7,500 Units, subcutaneous, every 8 hours, S+1 at 6:00 AM, PACU &
with weight GREATER than 100 kg	Post-op
() warfarin (COUMADIN) tablet	For patients with weight GREATER than 100 kg. oral, daily at 1700, Starting S+1, PACU & Post-op
	Indication:
() Pharmacy consult to manage warfarin	STAT, Until discontinued, Starting S
(COUMADIN)	Indication:

Required)

· · ·	Contraindications exist for mechanical prophylaxis	Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s) PACU & Post-op
• •	Place/Maintain sequential compression device continuous	Routine, Continuous, PACU & Post-op
	DERATE Risk of DVT - Non-Surgical (Selection juired)	1
Pha con One CHI stro Age Cen Hist	traindicated. or more of the following medical conditions: F, MI, lung disease, pneumonia, active inflamma ke, rheumatologic disease, sickle cell disease, l 60 and above tral line ory of DVT or family history of VTE	echanical prophylaxis is optional unless pharmacologic is ation, dehydration, varicose veins, cancer, sepsis, obesity, previous leg swelling, ulcers, venous stasis and nephrotic syndrome
Les Estr Mod	cipated length of stay GREATER than 48 hours s than fully and independently ambulatory ogen therapy derate or major surgery (not for cancer) or surgery within 3 months of admission	5
[] N	Ioderate Risk (Selection Required)	
	Moderate risk of VTE	Routine, Once, PACU & Post-op
N	Ioderate Risk Pharmacological Prophylaxis - Ion-Surgical Patient (Single Response) (Selecti Required)	on
	Contraindications exist for pharmacologic proph Order Sequential compression device	hylaxis - "And" Linked Panel
[]	Contraindications exist for pharmacologic prophylaxis	Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op
[]	Place/Maintain sequential compression device continuous	Routine, Continuous, PACU & Post-op
• • •	Contraindications exist for pharmacologic proph AND mechanical prophylaxis	-
[]	Contraindications exist for pharmacologic prophylaxis	Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op
[]	Contraindications exist for mechanical prophylaxis	Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s): PACU & Post-op
	enoxaparin (LOVENOX) injection (Single Responsion (Single Responsion (Selection Required)	onse)
()	enoxaparin (LOVENOX) syringe	40 mg, subcutaneous, daily at 1700, Starting S+1, PACU & Post-op Indication(s): VTE Prophylaxis
()	patients with CrCL LESS than 30 mL/min	30 mg, subcutaneous, daily at 1700, Starting S+1, PACU & Post-op For Patients with CrCL LESS than 30 mL/min Indication(s): VTE Prophylaxis
()	patients weight between 100-139 kg AND CrCl GREATER than 30 mL/min	30 mg, subcutaneous, every 12 hours at 0900, 2100, Starting S+1, PACU & Post-op For Patients weight between 100-139 kg and CrCI GREATER than 30 mL/min Indication(s): VTE Prophylaxis
()	patients weight 140 kg or GREATER AND CrCl GREATER than 30 mL/min	40 mg, subcutaneous, every 12 hours at 0900, 2100, Starting S+1, PACU & Post-op For Patients weight 140 kg or GREATER and CrCI GREATER than 30 mL/min Indication(s): VTE Prophylaxis
ited o	n 9/6/2022 at 2:23 PM from POC environment	

	2.5 mg, subcutaneous, daily, PACU & Post-op 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 CrCI LESS than 30 mL/min This patient has a history of or suspected case of Heparin-Induced Thrombocytopenia (HIT):
() heparin (porcine) injection	5,000 Units, subcutaneous, every 8 hours, PACU & Post-op
() heparin (porcine) injection (Recommended	5,000 Units, subcutaneous, every 12 hours, PACU & Post-op
for patients with high risk of bleeding, e.g. weight < 50kg and age > 75yrs)	Recommended for patients with high risk of bleeding, e.g. weight LESS than 50kg and age GREATER than 75yrs.
<ul> <li>HEParin (porcine) injection - For Patients with weight GREATER than 100 kg</li> </ul>	7,500 Units, subcutaneous, every 8 hours, PACU & Post-op For patients with weight GREATER than 100 kg.
() warfarin (COUMADIN) tablet	oral, daily at 1700, PACU & Post-op Indication:
<ul> <li>Pharmacy consult to manage warfarin (COUMADIN)</li> </ul>	STAT, Until discontinued, Starting S Indication:
[] Mechanical Prophylaxis (Single Response) (Sel Required)	ection
() Contraindications exist for mechanical prophylaxis	Routine, Once No mechanical VTE prophylaxis due to the following contraindication(s): PACU & Post-op
<ul> <li>Place/Maintain sequential compression device continuous</li> </ul>	Routine, Continuous, PACU & Post-op
HIGH Risk of DVT - Surgical (Selection Required)	
Severe fracture of hip, pelvis or leg Acute spinal cord injury with paresis	
Acute spinal cord injury with paresis Multiple major traumas Abdominal or pelvic surgery for CANCER Acute ischemic stroke	
Acute spinal cord injury with paresis Multiple major traumas Abdominal or pelvic surgery for CANCER Acute ischemic stroke History of PE [] High Risk (Selection Required) [] High risk of VTE	Routine, Once
Acute spinal cord injury with paresis Multiple major traumas Abdominal or pelvic surgery for CANCER Acute ischemic stroke History of PE [] High Risk (Selection Required) [] High risk of VTE [] High Risk Pharmacological Prophylaxis - Surgio (Single Response) (Selection Required)	al Patient
Acute spinal cord injury with paresis Multiple major traumas Abdominal or pelvic surgery for CANCER Acute ischemic stroke History of PE [] High Risk (Selection Required) [] High risk of VTE [] High Risk Pharmacological Prophylaxis - Surgio (Single Response) (Selection Required) () Contraindications exist for pharmacologic	Routine, Once
Acute spinal cord injury with paresis Multiple major traumas Abdominal or pelvic surgery for CANCER Acute ischemic stroke History of PE [] High Risk (Selection Required) [] High risk of VTE [] High Risk Pharmacological Prophylaxis - Surgio (Single Response) (Selection Required)	al Patient Routine, Once No pharmacologic VTE prophylaxis due to the following
Acute spinal cord injury with paresis Multiple major traumas Abdominal or pelvic surgery for CANCER Acute ischemic stroke History of PE [] High Risk (Selection Required) [] High risk of VTE [] High Risk Pharmacological Prophylaxis - Surgic (Single Response) (Selection Required) () Contraindications exist for pharmacologic prophylaxis	al Patient Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s):
Acute spinal cord injury with paresis Multiple major traumas Abdominal or pelvic surgery for CANCER Acute ischemic stroke History of PE [] High Risk (Selection Required) [] High risk of VTE [] High Risk Pharmacological Prophylaxis - Surgio (Single Response) (Selection Required) () Contraindications exist for pharmacologic prophylaxis () enoxaparin for VTE Prophylaxis (Single Response)	al Patient Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s): onse)
Acute spinal cord injury with paresis Multiple major traumas Abdominal or pelvic surgery for CANCER Acute ischemic stroke History of PE [] High Risk (Selection Required) [] High risk of VTE [] High Risk Pharmacological Prophylaxis - Surgic (Single Response) (Selection Required) () Contraindications exist for pharmacologic prophylaxis () enoxaparin for VTE Prophylaxis (Single Respondent () enoxaparin (LOVENOX) 30 mg daily at 1700	Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s): onse) 30 mg, subcutaneous, daily at 1700 Indication(s): VTE Prophylaxis
Acute spinal cord injury with paresis Multiple major traumas Abdominal or pelvic surgery for CANCER Acute ischemic stroke History of PE [] High Risk (Selection Required) [] High Risk of VTE [] High Risk Pharmacological Prophylaxis - Surgic (Single Response) (Selection Required) () Contraindications exist for pharmacologic prophylaxis () enoxaparin for VTE Prophylaxis (Single Respondent () enoxaparin (LOVENOX) 30 mg daily at 1700 () enoxaparin (LOVENOX) 30 mg every 12 hours	Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s): mse) 30 mg, subcutaneous, daily at 1700 Indication(s): VTE Prophylaxis 30 mg, subcutaneous, every 12 hours Indication(s): VTE Prophylaxis
Acute spinal cord injury with paresis Multiple major traumas Abdominal or pelvic surgery for CANCER Acute ischemic stroke History of PE [] High Risk (Selection Required) [] High Risk Of VTE [] High Risk Pharmacological Prophylaxis - Surgic (Single Response) (Selection Required) () Contraindications exist for pharmacologic prophylaxis () enoxaparin for VTE Prophylaxis (Single Respondent () enoxaparin (LOVENOX) 30 mg daily at 1700 () enoxaparin (LOVENOX) 30 mg every 12 hours () enoxaparin (LOVENOX) 40 mg daily at 1700	Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s): onse) 30 mg, subcutaneous, daily at 1700 Indication(s): VTE Prophylaxis 30 mg, subcutaneous, every 12 hours Indication(s): VTE Prophylaxis 40 mg, subcutaneous, daily at 1700 Indication(s): VTE Prophylaxis
Acute spinal cord injury with paresis Multiple major traumas Abdominal or pelvic surgery for CANCER Acute ischemic stroke History of PE [] High Risk (Selection Required) [] High Risk of VTE [] High Risk Pharmacological Prophylaxis - Surgic (Single Response) (Selection Required) () Contraindications exist for pharmacologic prophylaxis () enoxaparin for VTE Prophylaxis (Single Respondent () enoxaparin (LOVENOX) 30 mg daily at 1700 () enoxaparin (LOVENOX) 30 mg every 12 hours () enoxaparin (LOVENOX) 40 mg daily at	Routine, Once No pharmacologic VTE prophylaxis due to the following contraindication(s): onse) 30 mg, subcutaneous, daily at 1700 Indication(s): VTE Prophylaxis 30 mg, subcutaneous, every 12 hours Indication(s): VTE Prophylaxis 40 mg, subcutaneous, daily at 1700

() heparin (porcine) injection (Recommended	
for a sector of the last state of the sector	5,000 Units, subcutaneous, every 12 hours, S+1 at 6:00 AM
for patients with high risk of bleeding, e.g. weight < 50kg and age > 75yrs)	Recommended for patients with high risk of bleeding, e.g. weight LESS than 50kg and age GREATER than 75yrs.
() HEParin (porcine) injection - For Patients	7,500 Units, subcutaneous, every 8 hours, Starting S+1
with weight GREATER than 100 kg	For patients with weight GREATER than 100 kg.
() warfarin (COUMADIN) tablet	oral, daily at 1700, Starting S+1
	Indication:
() Pharmacy consult to manage warfarin	STAT, Until discontinued, Starting S
(COUMADIN)	Indication:
[] Mechanical Prophylaxis (Single Response) (Sel	ection
Required) () Contraindications exist for mechanical	Routine, Once
prophylaxis	No mechanical VTE prophylaxis due to the following contraindication(s):
() Place/Maintain sequential compression	Routine, Continuous
device continuous	
) HIGH Risk of DVT - Non-Surgical (Selection Requ	ired)
High Risk Definition	
Both pharmacologic AND mechanical prophylaxis One or more of the following medical conditions:	must be addressed.
	ant mutations, anticardiolipin antibody syndrome; antithrombin, protein C
or protein S deficiency; hyperhomocysteinemia; m	
Severe fracture of hip, pelvis or leg	
Acute spinal cord injury with paresis	
Multiple major traumas	
Abdominal or pelvic surgery for CANCER Acute ischemic stroke	
History of PE	
[] High Risk (Selection Required)	
[] High risk of VTE	Routine, Once, PACU & Post-op
[] High Risk Pharmacological Prophylaxis - Non-S Patient (Single Response) (Selection Required)	
() Contraindications exist for pharmacologic	Routine, Once
prophylaxis	No pharmacologic VTE prophylaxis due to the following
	No pharmacologic VTE prophylaxis due to the following contraindication(s):
prophylaxis	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op
() enoxaparin (LOVENOX) injection (Single Resp	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op
<ul> <li>prophylaxis</li> <li>() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> </ul>	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op ponse)
() enoxaparin (LOVENOX) injection (Single Resp	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op ponse) 40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op
<ul> <li>prophylaxis</li> <li>() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> <li>() enoxaparin (LOVENOX) syringe</li> </ul>	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op ponse) 40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op Indication(s): VTE Prophylaxis
<ul> <li>prophylaxis</li> <li>() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> </ul>	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op ponse) 40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op
<ul> <li>prophylaxis</li> <li>() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> <li>() enoxaparin (LOVENOX) syringe</li> <li>() patients with CrCL LESS than 30 mL/min</li> </ul>	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op ponse) 40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op Indication(s): VTE Prophylaxis 30 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op For Patients with CrCL LESS than 30 mL/min Indication(s): VTE Prophylaxis
<ul> <li>prophylaxis</li> <li>() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> <li>() enoxaparin (LOVENOX) syringe</li> <li>() patients with CrCL LESS than 30 mL/min</li> <li>() patients weight between 100-139 kg AND</li> </ul>	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op bonse) 40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op Indication(s): VTE Prophylaxis 30 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op For Patients with CrCL LESS than 30 mL/min Indication(s): VTE Prophylaxis 30 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op
<ul> <li>prophylaxis</li> <li>() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> <li>() enoxaparin (LOVENOX) syringe</li> <li>() patients with CrCL LESS than 30 mL/min</li> </ul>	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op ponse) 40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op Indication(s): VTE Prophylaxis 30 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op For Patients with CrCL LESS than 30 mL/min Indication(s): VTE Prophylaxis 30 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op For Patients weight between 100-139 kg and CrCl GREATER than 30
<ul> <li>prophylaxis</li> <li>() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> <li>() enoxaparin (LOVENOX) syringe</li> <li>() patients with CrCL LESS than 30 mL/min</li> <li>() patients weight between 100-139 kg AND</li> </ul>	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op ponse) 40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op Indication(s): VTE Prophylaxis 30 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op For Patients with CrCL LESS than 30 mL/min Indication(s): VTE Prophylaxis 30 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op For Patients weight between 100-139 kg and CrCl GREATER than 30 mL/min
<ul> <li>prophylaxis</li> <li>() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> <li>() enoxaparin (LOVENOX) syringe</li> <li>() patients with CrCL LESS than 30 mL/min</li> <li>() patients weight between 100-139 kg AND CrCl GREATER than 30 mL/min</li> </ul>	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op Donse) 40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op Indication(s): VTE Prophylaxis 30 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op For Patients with CrCL LESS than 30 mL/min Indication(s): VTE Prophylaxis 30 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op For Patients weight between 100-139 kg and CrCl GREATER than 30 mL/min Indication(s): VTE Prophylaxis
<ul> <li>prophylaxis</li> <li>() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> <li>() enoxaparin (LOVENOX) syringe</li> <li>() patients with CrCL LESS than 30 mL/min</li> <li>() patients weight between 100-139 kg AND CrCl GREATER than 30 mL/min</li> <li>() patients weight 140 kg or GREATER AND</li> </ul>	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op Donse) 40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op Indication(s): VTE Prophylaxis 30 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op For Patients with CrCL LESS than 30 mL/min Indication(s): VTE Prophylaxis 30 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op For Patients weight between 100-139 kg and CrCl GREATER than 30 mL/min Indication(s): VTE Prophylaxis 40 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op
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<ul> <li>prophylaxis</li> <li>() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> <li>() enoxaparin (LOVENOX) syringe</li> <li>() patients with CrCL LESS than 30 mL/min</li> <li>() patients weight between 100-139 kg AND CrCl GREATER than 30 mL/min</li> <li>() patients weight 140 kg or GREATER AND</li> </ul>	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op bonse) 40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op Indication(s): VTE Prophylaxis 30 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op For Patients with CrCL LESS than 30 mL/min Indication(s): VTE Prophylaxis 30 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op For Patients weight between 100-139 kg and CrCl GREATER than 30 mL/min Indication(s): VTE Prophylaxis 40 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op For Patients weight 140 kg or GREATER and CrCl GREATER than 30 mL/min Indication(s): VTE Prophylaxis
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<ul> <li>prophylaxis</li> <li>() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> <li>() enoxaparin (LOVENOX) syringe</li> <li>() patients with CrCL LESS than 30 mL/min</li> <li>() patients weight between 100-139 kg AND CrCl GREATER than 30 mL/min</li> <li>() patients weight 140 kg or GREATER AND CrCl GREATER than 30 mL/min</li> </ul>	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op ponse) 40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op Indication(s): VTE Prophylaxis 30 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op For Patients with CrCL LESS than 30 mL/min Indication(s): VTE Prophylaxis 30 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op For Patients weight between 100-139 kg and CrCI GREATER than 30 mL/min Indication(s): VTE Prophylaxis 40 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op For Patients weight 140 kg or GREATER and CrCI GREATER than 30 mL/min Indication(s): VTE Prophylaxis 2.5 mg, subcutaneous, daily, PACU & Post-op If the patient does not have a history of or suspected case of
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<ul> <li>prophylaxis</li> <li>() enoxaparin (LOVENOX) injection (Single Resp (Selection Required)</li> <li>() enoxaparin (LOVENOX) syringe</li> <li>() patients with CrCL LESS than 30 mL/min</li> <li>() patients weight between 100-139 kg AND CrCl GREATER than 30 mL/min</li> <li>() patients weight 140 kg or GREATER AND CrCl GREATER than 30 mL/min</li> </ul>	No pharmacologic VTE prophylaxis due to the following contraindication(s): PACU & Post-op ponse) 40 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op Indication(s): VTE Prophylaxis 30 mg, subcutaneous, daily at 1700, Starting S, PACU & Post-op For Patients with CrCL LESS than 30 mL/min Indication(s): VTE Prophylaxis 30 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op For Patients weight between 100-139 kg and CrCl GREATER than 30 mL/min Indication(s): VTE Prophylaxis 40 mg, subcutaneous, 2 times daily, Starting S, PACU & Post-op For Patients weight 140 kg or GREATER and CrCl GREATER than 30 mL/min Indication(s): VTE Prophylaxis 2.5 mg, subcutaneous, daily, PACU & Post-op 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.
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() heparin (porcine) injection (Recommended	5,000 Units, subcutaneous, every 12 hours, PACU & Post-op
for patients with high risk of bleeding, e.g. weight < 50kg and age > 75yrs)	Recommended for patients with high risk of bleeding, e.g. weight LESS than 50kg and age GREATER than 75yrs.
() HEParin (porcine) injection - For Patients	7,500 Units, subcutaneous, every 8 hours, PACU & Post-op
with weight GREATER than 100 kg	For patients with weight GREATER than 100 kg.
() warfarin (COUMADIN) tablet	oral, daily at 1700, PACU & Post-op Indication:
<ul> <li>Pharmacy consult to manage warfarin</li> <li>(COUMADIN)</li> </ul>	STAT, Until discontinued, Starting S Indication:
[] Mechanical Prophylaxis (Single Response) (Se Required)	election
() Contraindications exist for mechanical	Routine, Once
prophylaxis	No mechanical VTE prophylaxis due to the following contraindication(s): PACU & Post-op
<ul> <li>Place/Maintain sequential compression device continuous</li> </ul>	Routine, Continuous, PACU & Post-op
) HIGH Risk of DVT - Surgical (Hip/Knee) (Selectio Required)	n
High Risk Definition	
Both pharmacologic AND mechanical prophylaxis	s must be addressed.
One or more of the following medical conditions:	
	ant mutations, anticardiolipin antibody syndrome; antithrombin, protein C
or protein S deficiency; hyperhomocysteinemia; m	nyeloproliferative disorders)
Severe fracture of hip, pelvis or leg	
Acute spinal cord injury with paresis	
Multiple major traumas Abdominal or pelvic surgery for CANCER	
Acute ischemic stroke	
History of PE	
[] High Risk (Selection Required)	
[] High risk of VTE	Routine, Once, PACU & Post-op
[] High Risk Pharmacological Prophylaxis - Hip o (Arthroplasty) Surgical Patient (Single Response	r Knee
(Selection Required)	
() Contraindications exist for pharmacologic	Routine, Once
prophylaxis	No pharmacologic VTE prophylaxis due to the following
	contraindication(s):
	PACU & Post-op
() aspirin chewable tablet	162 mg, oral, daily, Starting S+1, PACU & Post-op
<ul> <li>() aspirin (ECOTRIN) enteric coated tablet</li> <li>() Apixaban and Pharmacy Consult (Selection R</li> </ul>	162 mg, oral, daily, Starting S+1, PACU & Post-op
[] apixaban (ELIQUIS) tablet	2.5 mg, oral, 2 times daily, Starting S+1, PACU & Post-op
	Indications: VTE prophylaxis
[] Pharmacy consult to monitor apixaban	STAT, Until discontinued, Starting S
(ELIQUIS) therapy	Indications: VTE prophylaxis
<ul> <li>enoxaparin (LOVENOX) injection (Single Res (Selection Required)</li> </ul>	ponse)
() enoxaparin (LOVENOX) syringe	40 mg, subcutaneous, daily at 0600, Starting S+1, PACU & Post-op Indication(s): VTE Prophylaxis
() enoxaparin (LOVENOX) syringe	30 mg, subcutaneous, 2 times daily at 0600, 1800, Starting S+1, PACL
	& Post-op Indication(s): VTE Prophylaxis
() enoxaparin (LOVENOX) syringe - For	30 mg, subcutaneous, daily at 0600, Starting S+1, PACU & Post-op
Patients with CrCL LESS than 30 mL/min	For Patients with CrCL LESS than 30 mL/min. Indication(s): VTE Prophylaxis
() enoxaparin (LOVENOX) syringe - For	30 mg, subcutaneous, 2 times daily at 0600, 1800, Starting S+1, PACL
Patients weight between 100-139 kg and	& Post-op
CrCl GREATER than 30 mL/min	For Patients weight between 100-139 kg and CrCl GREATER than 30
	mL/min.

() enoxaparin (LOVENOX) syringe - For	40 mg, subcutaneous, 2 times daily at 0600, 1800, Starting S+1, PACU
Patients weight between 140 kg or GREATER and CrCI GREATER than 30	& Post-op For Patients weight 140 kg or GREATER and CrCI GREATER than 30
mL/min	mL/min Indication(s): VTE Prophylaxis
() fondaparinux (ARIXTRA) injection	2.5 mg, subcutaneous, daily, Starting S+1, PACU & Post-op
	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
	This patient has a history of or suspected case of Heparin-Induced
	Thrombocytopenia (HIT):
() heparin (porcine) injection	5,000 Units, subcutaneous, every 8 hours, S+1 at 6:00 AM, PACU & Post-op
() heparin (porcine) injection (Recommended	5,000 Units, subcutaneous, every 12 hours, S+1 at 6:00 AM, PACU &
for patients with high risk of bleeding, e.g.	Post-op
weight < 50kg and age > 75yrs)	Recommended for patients with high risk of bleeding, e.g. weight LESS than 50kg and age GREATER than 75yrs.
() HEParin (porcine) injection - For Patients	7,500 Units, subcutaneous, every 8 hours, S+1 at 6:00 AM, PACU &
with weight GREATER than 100 kg	Post-op
() Rivaroxaban and Pharmacy Consult (Selection	For patients with weight GREATER than 100 kg.
Required)	
[] rivaroxaban (XARELTO) tablet for hip or	10 mg, oral, daily at 0600 (TIME CRITICAL), PACU & Post-op
knee arthroplasty planned during this admission	Indications: VTE prophylaxis
[] Pharmacy consult to monitor rivaroxaban	STAT, Until discontinued, Starting S
(XARELTO) therapy	Indications: VTE prophylaxis
() warfarin (COUMADIN) tablet	oral, daily at 1700, Starting S+1, PACU & Post-op Indication:
<ul> <li>Pharmacy consult to manage warfarin</li> <li>(COUMADIN)</li> </ul>	STAT, Until discontinued, Starting S Indication:
[] Mechanical Prophylaxis (Single Response) (Sele	ection
Required) () Contraindications exist for mechanical	Routine, Once
prophylaxis	No mechanical VTE prophylaxis due to the following contraindication(s):
	PACU & Post-op
<ul> <li>Place/Maintain sequential compression device continuous</li> </ul>	Routine, Continuous, PACU & Post-op
abs	
abs POD #1	
CBC with platelet and differential	AM draw repeats For 2 Occurrences, Post-op
Basic metabolic panel     Prothrombin time with INR	AM draw repeats For 2 Occurrences, Post-op AM draw repeats For 2 Occurrences, Post-op
-	Aim draw repeats 1 or 2 occurrences, 1 ost-op
Cardiology	
Diagnostic Imaging	
naging	
] XR Tibia Fibula 2 Vw Left	Routine, 1 time imaging, Starting S at 1:00 AM For 1 Occurrences, Post-op
] XR Tibia Fibula 2 Vw Right	Routine, 1 time imaging, Starting S at 1:00 AM For 1
1 VP Anklo 2 V/w Pilotorol	Occurrences, Post-op
] XR Ankle 3 Vw Bilateral	Routine, 1 time imaging, Starting S at 1:00 AM For 1, Post-op
] XR Ankle 3+ Vw Left	Routine, 1 time imaging, Starting S at 1:00 AM For 1,
XR Ankle 3+ Vw Right	Post-op Routine, 1 time imaging, Starting S at 1:00 AM For 1 ,
	Post-op

Post-op

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[] XR Foot 3 Vw Bilateral	Routine, 1 time imaging, Starting S at 1:00 AM For 1 , Post-op
[] XR Foot 3+ Vw Left	Routine, 1 time imaging, Starting S at 1:00 AM For 1, Post-op
[] XR Foot 3+ Vw Right	Routine, 1 time imaging, Starting S at 1:00 AM For 1, Post-op

# Other Diagnostic Studies

# Respiratory

### Rehab

### Consults

Ancillary Consult

[] Consult to Case Management	Consult Reason: Discharge Planning
	Post-op, And post discharge equipment needs. Plan
	discharge on POD #2-3.
[] Consult to Social Work	Reason for Consult: Discharge Placement
	Post-op, And post discharge equipment needs. Plan to
	discharge on POD #2-3.
[X] Consult PT eval and treat	Special Instructions: evaluate equipment needs at discharge
	Weight Bearing Status:
	Post-op
[] Consult OT eval and treat	Special Instructions: Instruct on use of hip kit.
	Weight Bearing Status:
	Post-op
[] Consult to Fracture Liaison Service	Clinical Indications:
	Post-op