

# Venous Thromboembolism (VTE)

# **Prevention protocol for adult patients**

Version 1.1

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# Aim and scope:

To standardize Venous Thromboembolism (VTE) risk assessment that delivers decision support to the point of care and standardize the clinical practice for VTE prevention to reduce morbidity and mortality related to thrombosis. The VTE prevention protocol developed to cover all related clinical specialties.

# Targeted end users:

This protocol intended to be used by the physicians and other Health Care Providers working at MOH hospitals.

# **Targeted population:**

All adult patients admitted to MOH hospitals.

# Level of Evidence:

Review of best practice and expert opinion.

# **Disclaimer:**

This living guidance is subject to updates with new emerging data or within 2 years. The task force members have no conflict of interest. This protocol is not attached to any funding.

# Scoring

VTE prevention protocols selected VTE and bleeding risk assessment based on:

- Modified Caprini tool for all cases except obstetric.
- Royal College of Obstetrics & Gynecology (RCOG) VTE and bleeding risk assessment tool for Obstetric cases only (Antenatal & Postnatal)



# **Modified Caprini**

RISK FACTORS				
<ul> <li>1 score for each</li> <li>Age 41-60 years</li> <li>BMI &gt; 25 Kg/m2</li> <li>Minor surgery</li> <li>Swollen legs (current)</li> <li>Varicose veins</li> <li>Major Surgery (in the past month)</li> <li>lung disease (e.g., emphysema or COPD)</li> <li>Currently on bed rest or restricted mobility</li> <li>History of Inflammatory bowel disease</li> <li>Acute myocardial infarction</li> <li>Congestive heart failure (&lt;1 month)</li> <li>Sepsis/ Pneumonia (&lt;1month)/</li> <li>History of unexplained or recurrent spontaneous abortion (&gt;3)</li> <li>Pregnant or post-partum (&lt;1 month)</li> <li>Oral contraceptives or hormone replacement</li> </ul>	<ul> <li>2 score for each</li> <li>Age: 61-74 years</li> <li>Arthroscopic Surgery</li> <li>Laparoscopy Surgery (&gt;45 min)</li> <li>Major open Surgery (&gt;45 min)</li> <li>Cancer (current or previous)</li> <li>Immobilizing Plaster cast</li> <li>Bed bound for more than 72hrs</li> <li>Central venous access</li> </ul>	3 score for each         Age≥ 75 years         History of DVT/PE         Family history of VTE         Factor V Leiden         Prothrombin 20210A         Lupus anticoagulant         Anticardiolipin antibodies         Elevated serum         homocysteine         Heparin-induced         thrombocytopenia         Other congenital or         acquired thrombophilia	<ul> <li><u>5 score for each</u></li> <li>Hip, pelvis or leg fracture (within the past month)</li> <li>Stroke (within past month)</li> <li>Multiple trauma (within past month)</li> <li>Elective major lower extremity arthroplasty</li> <li>Acute Spinal cord injury – paralysis (within the past month)</li> </ul>	

# Based on the calculation of scores from the selected risk factors the patient should fall in one of the following risk levels:

RISK LEVEL			
If total scores equal to 0 or	If total scores equal to 2:	If total scores equal to 3 or 4:	If total scores equal to or more than 5: <b><u>Highest</u> risk</b>
1: <u>Low</u> risk	<u>Moderate</u> risk	<u><b>High</b></u> risk	



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#### VTE prophylaxis based on Modified Caprini risk levels

#### For all MEDICAL and GENERAL SURGICAL conditions:

Category	Supportive Care	Pharmacotherapy	Precautions
• Low Risk	Encourage ambulation if     not restricted	No thromboprophylaxis required	
• <u>Moderate</u> <u>Risk</u>	<ul> <li>Encourage ambulation if not restricted</li> <li>Offer mechanical prophylaxis if pharmacological prophylaxis contraindicated</li> </ul>	<ul> <li>Enoxaparin 40 mg SC <u>once</u> daily OR</li> <li>Unfractionated Heparin 5000 Units SC BID or TID OR</li> <li>Fondaparinux dose 2.5 mg SC q24h</li> </ul>	If CrCl < 30ml/min, Enoxaparin 30 mg subcutaneously <u>once</u> daily and avoid Fondaparinux
• <u>High Risk</u>	<ul> <li>Encourage ambulation if not restricted <u>with or</u> <u>without</u> mechanical prophylaxis</li> </ul>	<ul> <li>Enoxaparin 40mg SC <u>once</u> daily OR</li> <li>Unfractionated Heparin 5000 Units SC TID OR</li> <li>Fondaparinux dose 2.5 mg SC q24h</li> </ul>	If CrCl < 30ml/min, Enoxaparin 30 mg subcutaneously <u>once</u> daily and avoid Fondaparinux
• <u>Highest Risk</u>	<ul> <li>Encourage ambulation if not restricted <u>with</u> mechanical prophylaxis</li> </ul>	<ul> <li>Enoxaparin 40mg SC <u>once</u> daily OR</li> <li>Unfractionated Heparin 5000 Units SC TID OR</li> <li>Fondaparinux dose 2.5 mg SC q24h</li> </ul>	If CrCl < 30ml/min, Enoxaparin 30 mg subcutaneously <u>once</u> daily and avoid Fondaparinux

#### Prophylactic Dose Anticoagulation based on BMI and CrCI:

CrCl (ml/min)	BMI (Kg/m²)	Enoxaparin	Fondaparinux	Unfractionated heparin
>30	<40	40 mg SC q24h	2.5 mg SC q24h	5000 units SC q8-12h
	>40	40 mg SC q12h	5 mg SC q24h	7500 units SC q8h
<30	<40		7500 units SC q8h	
	>40		UFH 7500 units SC q8h	

#### **Special consideration:**

**Oncology cases:** 

- Start prophylaxis early administration (postoperative, within 12 hours) or late administration (postoperative, after 12 hours) of antithrombotic prophylaxis in major surgical patients including cancer depending on bleeding risk
- Duration of anticoagulant for abdominal cancer surgery or previous VTE is **30 days**

#### Critical cases:

- For patient admitted to critical care units, routine assessment for VTE & bleeding risk is recommended and routine thrombo-prophylaxis is administered for at risk patients.
- For critical care patients who are at high-risk of bleeding, we recommend the optimal use of mechanical thromboprophylaxis with IPC at least until the bleeding risk decreases. When the high bleeding risk decreases.
- When the high bleeding risk decreases, we recommend that pharmacologic thromboprophylaxis be substituted for or added to the mechanical thromboprophylaxis.



# II- ORTHOPEDIC Surgery:

Category	Supportive Care	Pharmacotherapy	Precautions
A. Elective hip repla	acement		
For patient undergoing		Recommended thromboprophylaxis	
elective total hip		either:	
replacement (THR)		a. LMWH:	
		- At a usual high-risk dose 40 mg SC	
		q24h initiated 12 h before surgery	
		OR - At a usual high-risk dose 30 mg SC	
		q24h initiated 12 to 24 h <u>after</u> surgery	
		OR	
		b. Fondaparinux dose 2.5 mg SC	
		q24h initiated 6-8 hr after surgery	
		OR	
		<b>c. Apixaban 2.5</b> mg twice daily initiated 12-24 hr after surgery	
		OR	
		d. Adjusted-dose VKA (Warfarin)	
		started preoperatively the evening of	
		the surgical day (INR target 2.5, INR	
For a stight on down sing. TUD	Ontimal upp of a	range: 2.0 – 3.0 for 35 days)	Detiente placed en
For patient undergoing THR who have a high risk of	Optimal use of a mechanical method	When the high bleeding risk decreases, pharmacologic thrombo-	Patients placed on mechanical prophylaxis after
bleeding	with IPC	prophylaxis be substituted for or	surgery because of a high risk
bleeding		added to the mechanical thrombo-	of bleeding should have their
		prophylaxis	risk of bleeding consistently
			reassessed, with
			pharmacologic prophylaxis
			started as soon as the
			bleeding risk is decreased
B. Elective Knee Re	eplacement		
	-		
For patient undergoing total		Recommended thromboprophylaxis either:	
knee replacement (TKR)		a. LMWH:	
		- At a usual high-risk dose 30 mg SC	
		q24h initiated 12 to 24 h after surgery	
		<u>OR</u>	
		b. Fondaparinux dose 2.5 mg SC q24h initiated 6-8 hr after surgery	
		OR	
		<b>c. Apixaban 2.5</b> mg twice daily	
		initiated 12-24 hr after surgery	
		OR	
		d. Adjusted-dose VKA (Warfarin)	
		started preoperatively of the evening	
		of the surgical day <i>(INR target 2.5, INR range: 2.0 – 3.0 for 35 days)</i>	
For patient undergoing TKR	Optimal use of a	When the high bleeding risk	
who have a high risk of	mechanical method	decreases, pharmacologic thrombo-	
bleeding	with IPC	prophylaxis be substituted for or	
<u>_</u>		added to the mechanical thrombo-	
		prophylaxis to extend	
		pharmacological prophylaxis beyond 10 days after discharge	
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Category	Supportive Care	Pharmacotherapy	Precautions
C. Hip Fracture Sur	gery (HFS)		
For patient undergoing HFS		Routine thromboprophylaxis minimum 10 days up to 35 days is recommended: <b>a. Fondaparinux</b> 2.5 mg SC q24h initiated 6-8h after surgery <b>OR</b> <b>b. LMWH</b> 30mg SC q12h initiated 12- 24hr after surgery <b>OR</b> <b>c. Adjusted dose VKA (Warfarin)</b> preoperatively (INR target. 2.5. INR range. 2.0 to 3.0)	
D. Elective Spine Se	urgery		
Low risk	Encourage ambulation	No thromboprophylaxis required	
Moderate Risk such as:     Advanced age     Malignancy     Neurological deficit     Previous VT     An anterior surgical     approach	Optimal use of peri- operative IPC	The recommended thromboprphylaxis options: a. Enoxaparin 40 mg SC once daily OR b. Unfractionated Heparin 5000 Units SC or TID	VTE prophylaxis after elective spinal surgery can typically be initiated 12–24 hours postoperatively. Prophylaxis may need to be delayed if the surgical site remains open
• <u>Highest Risk</u>	Optimal use of a mechanical method (i.e. GCS and/or IPC)	The recommended thromboprophylaxis is one of the pharmacological thromboprophylaxis options combined with mechanical method: <b>a. Enoxaparin</b> 40 mg SC once daily <u>OR</u> <b>b. Unfractionated Heparin</b> 5000 Units SC or TID	
E. Knee arthroscop	У	•	
Low risk	Encourage ambulation	No thromboprophylaxis required	
High risk (multiple risk factors or following a complicated procedure)	Early mobilization	The recommended thromboprophylaxis is one of the pharmacological thromboprophylaxis options combined with mechanical method: LMWH minimum of 10 days. <b>a. Enoxaparin</b> 40 mg SC once daily <u>OR</u> <b>b. Unfractionated Heparin</b> 5000 Units SC or TID	
F. Isolated Lower Ex	ctremity Injuries Dis	tal to the Knee	
For patient with Isolated Lower Extremity Injuries Distal to the Knee		Routine use of thromboprophylaxis is <b>NOT</b> suggested	



#### III. UROLOGIC Surgery:

Category	Supportive Care	Pharmacotherapy	Precautions
For patient undergoing transurethral or other low risk procedures	Early mobilization	The recommendation is <u>against</u> the use of thromboprophylaxis	
For patient undergoing major open urologic procedures		The recommendation is to use <u>routine</u> thromboprophylaxis with: Pharmacological prophylaxis alone: a. Enoxaparin 40 mg SC once daily <u>OR</u> b. Unfractionated Heparin 5000 Units SC TID <u>OR</u> Pharmacological plus mechanical prophylaxis	Patients with very high risk for bleeding, we recommend the optimal use of mechanical thrombo- prophylaxis with GCS and/or IPC at least until the bleeding risk decreases. When the high bleeding risk decreases, we recommend pharmacologic thrombo-prophylaxis substituted for or added to the mechanical thrombo- prophylaxis.

# IV. LAPRAROSCOPIC Surgery:

Category	Supportive Care	Pharmacotherapy	Precautions
For patient undergoing entirely laparoscopic procedures who don't have additional risk factors	Early mobilization	The recommendation is <u>against</u> the use of thromboprophylaxis	
For patient undergoing entirely laparoscopic procedures who don't have additional risk factors	Optimal use of a mechanical method (i.e., GCS and/or IPC)	The recommendation is the use of <u>routine</u> thromboprophylaxis with either: Pharmacological prophylaxis alone: a. Enoxaparin 40 mg SC once daily <u>OR</u> b. Unfractionated Heparin 5000 Units SC TID OR	
		Pharmacological plus mechanical prophylaxis	

# V. BARIATRIC Surgery:

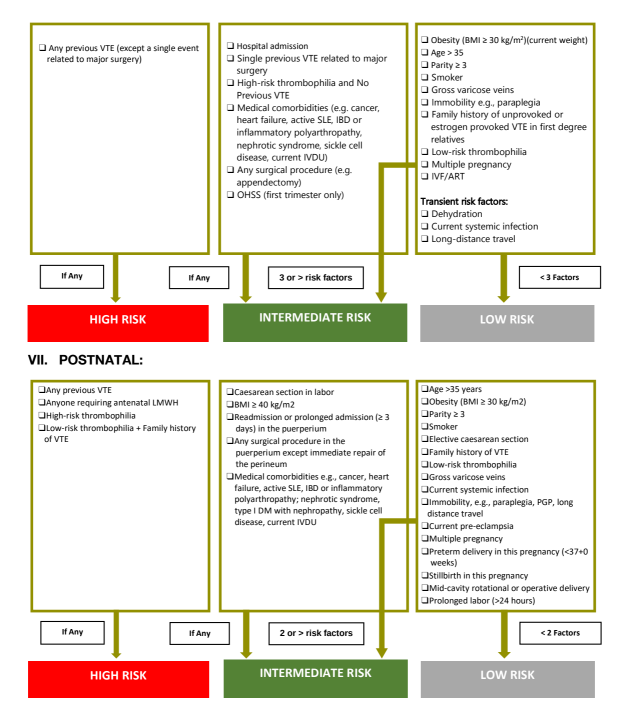
Category	Supportive Care	Pharmacotherapy	Precautions
For patient undergoing inpatient bariatric surgery	Optimal use of a mechanical method (i.e., GCS and/or IPC)	The recommendation is the use of <u>routine</u> thromboprophylaxis with either: Pharmacological prophylaxis alone: a. Enoxaparin 40 mg SC once daily <u>OR</u> b. Unfractionated Heparin 5000 Units SC TID	



Category	Supportive Care	Pharmacotherapy	Precautions
		OR Pharmacological plus mechanical prophylaxis	

# RCOG VTE risk factors (refer to RCOG risk factor calculator):

#### VI. ANTENATAL:





#### **VTE Prophylaxis based on RCOG risk levels**

Risk factors in pregnancy and the puerperium				
Pre-existing risk factors	Score			
Previous VTE (except a single event related to major surgery)	4			
Previous VTE provoked by major surgery	3			
Known high-risk thrombophilia	3			
Medical comorbidities e.g., cancer, heart failure; active systemic lupus erythematous, inflammatory polyarthropathy or inflammatory bowel disease in phrotic syndrome; type I diabetes mellitus with nephropathy; sickle cell disease. Current intravenous drug user	3			
Family history of unprovoked or estrogen related VTE in first-degree relative	1			
Known low-risk thrombophilia (no VTE)	1a			
Age (> 35 years)	1			
Obesity (body mass index [BMI] 30 0 kg/m2 or higher) either pre pregnancy or in early pregnancy	1 or 2b			
Parity ≥ 3	1			
Smoker	1			
Gross varicose veins	1			

Obstetric risk factors	Score
Previous VTE (except a single event related to major surgery)	4
Previous VTE provoked by major surgery	3
Known high-risk thrombophilia	3
Medical comorbidities e.g., cancer, heart failure; active systemic lupus erythematous, inflammatory polyarthropathy or inflammatory bowel disease; Interprete provide the syndrome; type I diabetes mellitus with nephropathy; sickle cell disease. Current intravenous drug user	3
Pre-eclampsia in current pregnancy	1
ART/IVF (antenatal only)	1
Multiple pregnancy	1
Caesarean section in labor	2
Elective caesarean section	1
Mid-cavity or rotational operative delivery	1
Prolonged labor (> 24 hours)	1
PPH (> 1 liter or transfusion)	1
Preterm birth < 37+0 weeks in current pregnancy	1
Stillbirth in current pregnancy	1

Transient risk factors	Score
Any surgical procedure in pregnancy or puerperium except immediate repair of the 3 perinea, e.g., appendicectomy, postpartum sterilization	3
Hyperemesis	4
OHSS (first trimester only)	1
Current systemic infection	1
Immobility, dehydration	1

• If total score  $\geq$  4 antenatally, consider thromboprophylaxis from the first trimester.

• If total score 3 antenatally, consider thromboprophylaxis from 28 weeks.

If total score ≥ 2 postnatally, consider thromboprophylaxis for at least 10 days.



- If admitted to hospital antenatally consider thromboprophylaxis.
- If prolonged admission (≥ 3 days) or readmission to hospital within the puerperium, consider thromboprophylaxis.

# VTE prophylaxis for OBSTETRICS (Ante and Post-natal):

- Pharmacological thromboprophylaxis should be avoided, discontinued or postponed in women at risk of bleeding after careful consideration of the balance of risks of bleeding and thrombosis.

- LMWH is safe and easy to use postpartum and has the advantage of not requiring monitoring.

- For those women receiving LMWH antenatally (and therefore for 6 weeks postpartum) or for those requiring 10 days' postpartum thromboprophylaxis, it is the agent of choice.

- Experience of LMWH in the puerperium reports no problems during breastfeeding

	Category	Supportive Care	Pharmacotherapy	Precautions
•	Low Risk	- Early mobilization & avoid dehydration	- No thromboprophylaxis required	
•	<u>Moderate</u> <u>Risk</u>	<ul> <li>Encourage ambulation</li> <li>Intermittent pneumatic compression or Graduated compression stockings</li> </ul>	The recommendation is the use of routine         thromboprophylaxis with either:         a. Enoxaparin SC once daily according to current         weight as the following:         Weight       Enoxaparin         < 50 kg       □20 mg daily         50–90 kg       □40 mg daily         91–130 kg       □60 mg daily         131–170 kg       □80 mg daily         > 170 kg       □0.6 mg/kg/ day         OR       b. Unfractionated Heparin 5000 Units SC BID or TID         Antenatal prophylaxis from 28 weeks in pregnancy.	
•	<u>High Risk</u>	<ul> <li>Encourage ambulation</li> <li>Intermittent pneumatic compression or Graduated compression stockings</li> </ul>	The recommendation is the use of routine thromboprophylaxis with either:a. Enoxaparin SC once daily according to current weight as the following:WeightEnoxaparin< 50 kg□20 mg daily50–90 kg□40 mg daily91–130 kg□60 mg daily131–170 kg□80 mg daily> 170 kg□0.6 mg/kg/ dayOROb. Unfractionated Heparin 5000 Units SC BID Antenatal prophylaxis from first trimester.	



	Med	ication Related Ir	formation	
Medication	Contraindication	Major Drug Interactions	Required dose adjustment	Pregnancy
Unfractionate d Heparin (UFH)	<ul> <li>Severe thrombocytopenia</li> <li>Uncontrolled active bleeding; except when due to DIC</li> </ul>	Apixaban Dabigatran Endoxaban Mifepristone Rivaroxaban Streptokinase Urokinase	Renal impairment: No specific recommendations are available Hepatic impairment: No specific recommendations are available Geriatric: No adjustment necessary; however, a higher incidence of bleeding has been reported in patients over 60 years of age, especially women, therefore lower doses of heparin may be indicated in these patients.	Fetal risk cannot be ruled out
Enoxaparin	<ul> <li>Active major bleeding</li> <li>History of immune-mediated heparin-induced thrombocytopenia within the past 100 days or in presence of circulating antibodies</li> <li>Hypersensitivity to benzyl alcohol (present in multi-dose formulation)</li> <li>Hypersensitivity to enoxaparin sodium, heparin, or pork products</li> </ul>	Apixaban Dabigatran Endoxaban Mifepristone Rivaroxaban Urokinase	Renal impairment (CrCl 30 to 80 mL/min): No adjustment necessary. Renal impairment (CrCl less than 30 mL/min): Unfractionated heparin recommended instead of low- molecular-weight heparin (LMWH); if LMWH is used, reduce usual recommended dose by 50%. Renal impairment (CrCl less than 30 mL/min) in prevention of DVT following abdominal surgery: 30 mg subQ once daily. Renal impairment (CrCl less than 30 mL/min) in prevention of DVT following hip or knee replacement surgery: 30 mg subQ once daily. Renal impairment (CrCl less than 30 mL/min) in prevention of DVT following hip or knee replacement surgery: 30 mg subQ once daily. Renal impairment (CrCl less than 30 mL/min) in prevention of DVT in medical patients during acute illness: 30 mg subQ once daily.	Fetal risk cannot be ruled out
Warfarin	<ul> <li>Blood dyscrasias</li> <li>Cerebral aneurysms</li> <li>CNS hemorrhage</li> <li>Dissecting aorta</li> <li>Eclampsia, preeclampsia, threatened abortion</li> <li>Gastrointestinal, genitourinary, or respiratory tract ulcerations or overt bleeding</li> <li>Hemorrhagic tendencies</li> <li>Hypersensitivity to warfarin or any component of the product</li> <li>Major regional or lumbar block anesthesia</li> <li>Malignant hypertension</li> <li>Pregnancy, except in pregnant women with mechanical heart valves, who are at high risk of thromboembolism</li> <li>Recent or potential surgery of central nervous system or eye</li> <li>Recent or potential traumatic surgery resulting in large open surface</li> <li>Spinal puncture and other procedures with potential for uncontrollable bleeding Unsupervised and potentially noncompliant patients</li> </ul>	Tamoxifen Streptokinase Urokinase Allopurinol Amiodarone Barbiturates Cholestyramine resin	Renal impairment: No adjustment necessary; monitor INR more frequently in patients with compromised renal function to maintain INR within the therapeutic range Geriatric: Consider using lower initial and maintenance dosage Pregnancy, mechanical valve: Warfarin to goal INR plus aspirin 75 mg to 100 mg/day during second and third trimesters; during first trimester, warfarin may be continued in patients who can achieve therapeutic INR with doses of 5 mg/day or less. Frequent monitoring required. Discontinue warfarin and initiate continuous infusion unfractionated heparin prior to planned vaginal delivery (guideline dosing)	Contraindicate d



Medication	Contraindication	Major Drug	Required dose adjustment	Pregnancy
Fondaparinux	<ul> <li>Contraincication</li> <li>Contraincication</li> <li>Contraincication</li> <li>Contraincication</li> <li>With a CrCl &lt; 30 mL/min/1.73 m2 Body weight less than 50 kg in VTE prophylaxis</li> <li>Active major bleeding</li> <li>Thrombocytopenia associated with positive in vitro test for antiplatelet antibody in the presence of fondaparinux sodium</li> <li>History of serious hypersensitivity reaction (eg, angioedema, anaphylactoid or anaphylactic reactions)</li> </ul>	Apixaban Dabigatran Endoxaban Mifepristone Rivaroxaban	Required dose adjustment Renal impairment (CrCl 30 to 50 mL/min): Use with caution; may cause prolonged anticoagulation. Hepatic impairment (mild to moderate): No dosage adjustment required; however, observe closely for signs/symptoms of bleeding. Geriatric: Pay particular attention to dosing directions and concomitant medications (especially anti-platelet medication). Hemodiafiltration in patients with heparin-induced thrombocytopenia: Initiate at 0.03 mg/kg post dialysis body weight, administered via the efferent line of the dialyzer; titrate in increments of 0.01 mg/kg post dialysis anti-Xa activity.	Fetal risk cannot be ruled out
Apixaban	<ul> <li>Contraindicated in patients with a CrCl &lt; 25 mL/min/1.73 m2 SCr &gt; 2.5 mg/dL</li> <li>Active pathological bleeding</li> <li>Severe hypersensitivity (eg, anaphylactic reactions) to apixaban</li> </ul>	Rifampin, phenytoin, carbamazepine, St. John's wort) protease inhibitors, itraconazole, ketoconazole	50% dose reduction if receiving 5 or 10 mg twice daily with strong CYP3A4 and P-gp inhibitor (e.g., protease inhibitors, itraconazole, ketoconazole, conivaptan)	Fetal risk cannot be ruled out



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مستشفى :	AGE:	سنهYEARS	العمر: يومDAYS شهر MONTHS						
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القسم/الوحدة: Dept./Unit:	GENDER:	Male Femal	الجنس: 🔄 نكر 🔄 انثى e						
Adult In-Patient Venous Thromboembolism (VTE) Assessment and Prophylaxis									
Note: (To be assessed for all adult ( $\geq$ 18 year	s) patients during	admission and rep	eated if patients' condition changed)						
Diagnosis:									
Admission     Post-surgical procedu	re 🗆 C	hange in condition	Other						
STEP 1 : N	lark risk factors the	n calculate the total s	score						
Risk Factor Score =1	Risk Facto	or Score = 2	Risk Factor Score = 3						
① Age 41 to 60 years	② Age 61- 74 years		③ Age ≤ 75 years						
0 Medical patient at bed rest (e.g. Sickle cell	② Arthroscopic surg	gery	③ Personal history of DVT/PE						
disease, dehydration, diabetes, etc )	② Malignancy (pres	sent or previous)	③ Family history of thrombosis						
① Minor surgery planned	② Major surgery (> 4	15 minutes) under G.A.	③ Positive Factor V Leiden						
${ m I}{ m D}$ History of prior major surgery (< 1 month)	<sup>2</sup> Laparoscopic sur	gery(> 45 minutes)	③ Elevated serum homocysteine						
① Varicose veins	<sup>②</sup> Patient confined t		③ Positive lupus anticoagulant						
① History of inflammatory bowel disease		ter cast for lower limbs	③ Elevated anticardiolipin antibodies						
① Swollen legs (current)	(< 1 month)		③ Positive prothrombin 20210A						
① Obesity (BMI > 25)	<ul> <li>Central venous ac</li> </ul>	cess	③ Heparin-induced thrombocytopenia (HIT)						
① Acute myocardial infarction			③ Other congenital or acquired thrombophi						
① Congestive heart failure (< 1 month)			: Protein C, Protein S, Antithrombin III						
① Sepsis(< 1 month)			Risk Factor Score = 5						
${\rm \textcircled{O}}$ Serious lung disease incl. pneumonia (< 1 month)			<sup>⑤</sup> Elective Knee or Hip Arthroplasty						
① Abnormal pulmonary function (COPD)			⑤ Hip and / or Pelvis fracture (< 1 month)						
① Oral contraceptives or hormone replacement			<pre>⑤ Stroke(&lt; 1 month)</pre>						
therapy			<ul> <li>Subject (1 month)</li> <li>Multiple trauma(&lt; 1 month)</li> </ul>						
① Pregnancy or postpartum (refer to antenatal and			<ul> <li>S Acute spinal cord (paralysis), (&lt; 1 month)</li> </ul>						
postnatal VTE prophylaxis forms)									
① History of unexplained stillborn infant,									
recurrent spontaneous abortion (≥ 3), premature birth with toxemia or growth restricted infant									
	otal Risk Factor Scor								
			ith any of the following						
STEP 2 : Assess risk versus the	a benefit of proping	laxis in the patients w							
Contraindications	aria / Fandanarianu /	— Listen, of easterintee	Warnings/Precaution						
<ul> <li>Patient on therapeutic doses of: Heparin / Enoxap.</li> <li>Warfarin / Rivaroxaban / Dabigatran / Apixaban</li> </ul>	ann / Fondapannux /		tinal bleed of Hemorriagic scroke						
□ Hypersensitivity to low molecular weight heparin,		□Renal failure with (	Creatinine clearance less than 30 ml/min (fo						
unfractionated heparin, (including heparin-induced		Enoxaparin-modify the							
thrombocytopenia)		Enoxuputin moury the							
Active bleeding / Fall Patients		Coagulopathy (high a graduate of the second seco	aPTT. PT/INR ≥ 1.5)						
□ Uncontrolled HTN (SBP >185 and /or DBP > 110 mm	Hg)	Clinically significant thrombocytopenia (Platelet count less than 50)							
Epidural anesthesia (within last 12 hours or planned			urgery or intracranial surgery						
If the patient has any of the abo									
Sequential Compression Device (SCD)[ first priority]		-							
If there are any contraindications to (SCD) & (ECS): Ga	ngrene; Recent Skin G	iraft; Suspected existing	s lower limb Deep Venous Thrombosis: Use						
electric stimulation device.									
GDOH- MRA-COR-IP(VTE)-073 AVTE	10	OF 2 ISSUED DATE: 3	0/12/2021 update date 23/11/2023 SN						

رقم الملف الطبي

	STEP 3 :	MANDATORY to Select One or More of the Risk level and Treatment Options	
Risk Score	Risk Level	Pharmacologic	Mechanical Device
1-0	□Low	Early ambulation	
1-0         2         4-3         or more 5         *The recomm         n Oncology-su         ischarge (4-5 w)         No orders for         If the patient         Labs: Check base         Iurse intervent         The nurse no         Providing VTE         The nurse pro         The nurse ap         Compression         urse'/Midwife         Date, Time and         Patient education	□Moderate	LMWH*:(CrCl > 30mL/min)	
		LMWH: If BMI $\ge$ 40: $\Box$ Enoxaparin 60 mg subcutaneously once daily	
		OR   Enoxaparin 40 mg subcutaneously BID	
		Heparin 5000 units subcutaneously every 12 hrs.	
		Fondaparinux dose 2.5 mg SC q24h (HIT or Allergy) avoid if CrCl < 30ml/min	
<i>1</i> _2		LMWH*:(CrCl > 30mL/min)   Enoxaparin 40 mg subcutaneously once daily	
4-3	□High	LMWH:(CrCl < 30mL/min)   Enoxaparin 30 mg subcutaneously once daily	
		LMWH: If BMI $\ge$ 40: $\Box$ Enoxaparin 60 mg subcutaneously once daily	
		OR 🗆 Enoxaparin 40 mg subcutaneously BID	
		□Heparin 5000 units subcutaneously every 8 hrs.	
		Fondaparinux dose 2.5 mg SC q24h (HIT or Allergy) avoid if CrCl < 30ml/min	
or more 5		LMWH*:(CrCl > 30mL/min)   Enoxaparin 40 mg subcutaneously once daily	Plus: SCD
	□Highest	LMWH:(CrCl < 30mL/min)   Enoxaparin 30 mg subcutaneously once daily	
		LMWH: If BMI $\ge$ 40: $\Box$ Enoxaparin 60 mg subcutaneously once daily	
		OR   Enoxaparin 40 mg subcutaneously BID	
		Heparin 5000 units subcutaneously every 8 hrs.	
		Fondaparinux dose 2.5 mg SC q24h (HIT or Allergy) avoid If CrCl < 30ml/min	
		be used as alternative according to hospital formulary	
	rgery, Orthopedic ( /eeks): Enoxaparin	TKR,THR,HFS), abdominal surgery and Bariatric surgery: consider extend or DOAC	led- prophylaxis after
	prophylaxis, Reas		
	's condition changes or i	s is a general guideline and the physician's clinical judgment may override it. If there is a procedure with bleeding risk, the risk stratification must be revised using a new t every 72 hours thereafter. Notify physician if platelet count less than 100,000 or	
<b>lurse intervent</b> The nurse not	ions tified the physician	to fill out the form	
Providing VTE	mechanical proph	ylaxis devices.	
=	-	ly education (the patient received his/her injection by him/her-self.	
-	-	n about administration.	tooching foot los aver-is-
Compression, Iurse'/Midwife	/elastic stockings ry Name and Stamp	easures (non-pharmacologic measures):  Assist in early mobilization.	teaching foot-leg exercise
Patient educa ffectetc.	ited by pharmacist	(medication information: indication, duration, frequency, important for a	dherence, suspected side
Patient educa	ited by health educ		
ain Responsible Pl	nysician's Name and Sta	mp: Date, Time and Signature:	

: الاسم MRN؛

NAME of Patient:

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ارة الصحة Ministry of Hea	MRN: NAME of Pat NATIONALII	<b>Y</b> :YEAF	 RSمنس MON'	رقم الملف الطبي رقم الملف الطبي الاسم : الاسم : الجنسية: العمر : يومDAYS شهر THS		
Hospital:	مستشفى :	DATE OF BI	RTH: 20_	//	تاريخ الميلاد/ هـ	
Region:	المنطقة /المحافظة	GENDER:	Male	Female	الجنس: 🔄 ذكر 🔄 انثى	
Dept./Unit: Antenatal V		mboemboli			and Prophylaxis	
(to be assessed at firs	st visit to obste	trics clinic and i	repeated i	n 2nd and 3r	d trimester or if admitted)	
□Any previous Venous thromboembolism (VTE) except a single event related to major surgery	ion. VTE related to majo pophilia + no VTE idities e.g., cancer, emic lupus erythem drome or inflammat ephrotic syndrome, thy, sickle cell disea user ocedure e.g., stimulation syndrom ester only)	heart atosus, ory type I ase	□ Family history estrogen provol □ Low-risk thro □ Multiple preg □ IVF/ART (Ant Fertilization (IV Transient risk	se veins eclampsia .g., paraplegia, pelvic girdle pain y of unprovoked or ked VTE in first-degree relative mbophilia nancy tiretroviral Therapy / In Vitro F) pregnancy factors: Dehydration/ Current systemic Infection.		
If Any	If Any	-				
HIGH RISK Requires antenatal prophylax with LMWH. Refer to Consultant in pregnancy expert/team	kis Consid prophyla:	IEDIATE RISK der antenatal xis with LMWH.	r antenatal prophyla:		Fewer than Three risk factors. Mobilization and s. avoidance of dehydration	
		antenatal and post alternative according to ho			Mark contraindications to LMWH	
Weight	Enoxaparin	Tinzaparin (75 u/kg/day)		CrCl	C Known bleeding disorder	
< 50 kg	□20 mg daily	□3500 units daily		30 ml/min	□ Active bleeding	
50–90 kg	□40 mg daily	□4500 units daily	Enoxap to Wigh	parin dose based	Thrombocytopenia (platelet count < 50,000)	
91–130 kg	□60 mg daily	□7000 units daily		<u>30 ml/min</u> n 5000 IU	Acute stroke in previous 4	
131–170 kg	□80 mg daily*	□9000 units daily	subcuta	aneously of	weeks (hemorrhagic or	
> 170 kg	□0.6 mg/kg/day	□75 u/kg/day	could b	ionated heparin be used and	ischemic)	
High prophylactic dose for women weighing 50–90 kg	4500 units 12 hour	hours u	ed every 12 until LMWH can umed after y.	Uncontrolled hypertension		
Patient at significant risks for Use Sequential Compressi Properly fitted graduated co	on Device (SCD) [f	irst priority]	gulation: 🗅		ations of SCD: Gangrene; Recent uspected existing Deep Venous	
<ul> <li>Special Cases:</li> <li>Previous VTE + Anti phospholipid syndrome (APS): High dose. (The same dose of LMWH but TWICE daily)</li> <li>Previous VTE + Anti Thrombin deficiency: High dose. (The same dose of LMWH but TWICE daily)</li> <li>Recurrent VTE (2 or more): High dose. (The same dose of LMWH but TWICE daily)</li> </ul>						

NAME of Patient: :	וצייים MRN:								الملف الطبي
<ul> <li>Unfractionated Heparin: Indications</li> <li>Around the time of delivery in women at very high risk of thrombosis ( there may be reluctance to use LMWH in case regional anesthetic tec are required)</li> <li>In women at increased risk of hemorrhage</li> <li>The required interval between a prophylactic dose of unfractionated h and regional analgesia or anesthesia is less (4 hours) than with LMWI hours)</li> </ul>	ery in women at very high risk of thrombosis (when e to use LMWH in case regional anesthetic techniques risk of hemorrhage etween a prophylactic dose of unfractionated heparin etween a prophylactic dose of unfractionated heparin								
Admission Date& time Physicians Name: Date &time :				Signa	ature:	:			
Nurse interventions:									
□ The nurse notified the physician to fill out the form									
Providing VTE mechanical prophylaxis devices.									
□ The nurse provided patient/family education (the patient)	atient receiv	ved his/h	er injec	tion b	y him/	her-s	self.		
The patient receive only education about administra	ition.								
□ The nurse applies prevention measures (nonpharm	nacologic m	neasures	):						
Assist in early mobilization.									
teaching foot-leg exercises.									
Compression/elastic stockings									
Nurse'/Midwifery Name and Stamp: Signature:					D	ate,	, Time	e and	t
Patient educated by pharmacist (medication inform adherence, suspected side effectetc.	ation: indic	cation, du	ration, f	freque	ency, ir	npor	rtant fo	or	
Patient educated by health educator									

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: MRN الأسم MRN رقم الملف الطبى NAME of Patient: **Risk factors for VTE** Pre-existing risk factors Tick Score Tick Score Previous VTE (except a single event related to major surgery) 4 Previous VTE provoked by major surgery 3 Known high-risk thrombophilia 3 Medical comorbidities e.g. cancer, heart failure; active systemic lupus erythematous, inflammatory polyarthropathy or inflammatory bowel 3 disease; nephrotic syndrome; type I diabetes mellitus with nephropathy; sickle cell disease; current intravenous drug user Family history of unprovoked or estrogen-related VTE in first-degree 1 relative Known low-risk thrombophilia (no VTE) 1a Age (> 35 years) 1 Obesity 1 or 2b Parity  $\geq 3$ 1 Smoker 1 Gross varicose veins 1 Obstetric risk factors Pre-eclampsia in current pregnancy 1 ART/IVF (antenatal only) 1 Multiple pregnancy 1 Caesarean section in labor 2 2 **Elective caesarean section** 1 Mid-cavity or rotational operative delivery 1 Prolonged labour (> 24 hours) 1 PPH (> 1 litre or transfusion) 1 Preterm birth < 37+0 weeks in current pregnancy 1 Stillbirth in current pregnancy 1 Transient risk factors Any surgical procedure in pregnancy or puerperium except immediate 3 repair of the 3 perineum, e.g. appendicectomy, postpartum sterilisation **Hyperemesis** 3 OHSS (first trimester only) 4 4 Current systemic infection 1 Immobility, dehydration 1 Total

• If total score  $\geq$  4 antenatally, consider thromboprophylaxis from the first trimester.

• If total score 3 antenatally, consider thromboprophylaxis from 28 weeks.

• If total score  $\geq$  2 postnatally, consider thromboprophylaxis for at least 10 days.

• If admitted to hospital antenatally consider thromboprophylaxis.

If prolonged admission (≥ 3 days) or readmission to hospital within the puerperium consider thromboprophylaxis

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محية Ministry of Hospitals Regions Dept./Units	مستشفى : المنطقة /المحافظ لقسم/الوحدة:	MRN:						
Postnata	al Ven	ous Thre	omboembolis	m Ass	essment ar	d Prophylaxis		
(to be	e assess	ed after deliv	very and repeated d	uring obst	etrics clinic visit or	if re-admitted)		
Thromboembolism (VTE)□ BMI ≥ 40 kg/□ Anyone requiring antenatal LMWH□ Readmission (≥ 3 puerperium□ High-risk thrombophilia□ Any surgical puerperium exit□ Low-risk thrombophilia + Family history□ Medical com heart failure, are erythematosus syndrome or in			ion or prolonged 2 3 days) in the cal procedure in the except immediate represent pmorbidities e.g., cance , active systemic lupus sus, irritable bowel r inflammatory	m2       □ Obesity (BMI ≥ 30 kg/m2)         □ or prolonged days) in the       □ Parity ≥ 3         □ procedure in the cept immediate repair n       □ Elective caesarean section         □ Family history of VTE       □ Low-risk thrombophilia         □ Gross varicose veins       □ Current systemic infection         □ Immobility, e.g., paraplegia, pelvic girdle pain, long distance travel         □ Current pre-eclampsia         □ Multiple pregnancy         y; nephrotic syndrome,				
If Any		If Any	Drug user			or operative delivery		
HIGH RISK At least 6 week postnatal prophyl LMWH		I At Note: If	NTERMEDIATE RIS least 10 days 'post prophylactic LMW persisting or > 3 ris consider extendin boprophylaxis with	natal 'H sk factor Ig	Ē	LOWER RISK r than two risk factors) arly mobilization and idance of dehydration		
Thro		ylaxis doses f	or antenatal and postn d as alternative accordi	atal LMWH	ital formulary	Mark contraindications to		
Weight	Enoxapa		Tinzaparin (75 u/kg/day)		CrCl	C Known bleeding disorder		
< 50 kg	□20 mg	daily	□3500 units daily	<u>CrCl &gt; 30</u>	<b>ml/min</b> Enoxaparin	<ul> <li>Active bleeding</li> <li>Thrombocytopenia</li> </ul>		
50–90 kg	□40 mg	daily	□4500 units daily	dose base CrCl < 30		(platelet count < 50,000)		
91–130 kg	<b>□</b> 60 mg	daily	□7000 units daily	Heparin 50 subcutane	000 IU	Acute stroke in previous 4 weeks (hemorrhagic or		
131–170 kg	<b>□</b> 80 mg	daily	□9000 units daily	unfractiona	ated heparin could be	ischemic)		
High prophylactic dose			□75 u/kg/day □4500 units 12 hourly	hours until	epeated every 12 LMWH can be fter delivery.	Uncontrolled hypertension		
□Use Sequential Com	Patient at significant risks for bleeding or contraindication to anticoagulation:Contraindications of SCD: Gangrene; RecentUse Sequential Compression Device (SCD) [first priority]Skin Graft; Suspected existing Deep Venous ThrombosisProperly fitted graduated compression stockings (15-30mmHg)Thrombosis							
Previous VTE + Anti Th	Previous VTE + Anti phospholipid syndrome (APS): High dose. (The same dose of LMWH but TWICE daily)							
GDOH-MRA-INP-0072B			1of2 ISSUED DA	ATE: 20/0	8/2017 Update	date 23/11/2023 SN		

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<ul> <li>Unfractionated Heparin: Indications</li> <li>In women at increased risk of hemorrhage</li> <li>The required interval between a prophylactic dose of unfractionated heparin and regional analgesia or anesth less (4 hours) than with LMWH (12 hours)</li> </ul>		<ul> <li>This is a general guideline and the physician's clinical judgment may override it.</li> <li>If the patient's condition changes or if there is a procedure with bleeding risk, the risk stratification must be revised using a new form by the Primary Team</li> <li>Labs: Check baseline CBC and at least every 72 hours thereafter. Notify physician if platelet count less than 100,000 or drop by 50% from baseline, or renal impairment (CrCl &lt; 30mL/min)</li> </ul>									form er.	
Admission Date& time												
Physicians Name: Signature:			_									
Date &time :												
Nurse interventions:												
□ The nurse notified the physician to fill out	the form											
Providing VTE mechanical prophylaxis de	evices.											
The nurse provided patient/family education	ion (the pat	tient re	ceiv	red ł	nis/he	r inje	ectio	n by l	nim/h	ner-s	self.	
The patient receive only education about a	administrati	ion.										
The nurse applies prevention measures (	nonpharma	acologi	c m	eası	ures):							
Assist in early mobilization.												
teaching foot-leg exercises.												
Compression/elastic stockings												
Nurse'/Midwifery Name and Stamp:									Da	ate,	Time	e an
Patient educated by pharmacist (medicat important for adherence, suspected side effective		ition: ir	ndica	atior	, dur	ation	, fre	quenc	су,			
Patient educated by health educator												