VENOUS THROMBOEMBOLISM

Raed Abughazaleh, PharmD, BCPS PHAR 551: Pharmacotherapy I Birzeit University

Definitions

- DVT and PE
- · Provoked vs. unprovoked
- Risk factors: surgery, trauma, hx VTE, hypercoagulable D/O, age, immobility, major surgery, malignancy, pregnancy, estrogen, HIT, critical illness
- Caused by imbalance between thrombogenic and antithrombotic forces

Homeostasis and Thrombosis



Source: DiPiro J $\it{et~al.}$ Pharmacotherapy Principles and Practice, $\it{2}^{nd}$ Ed. 2010

Epidemiology

- 1/3 of pts with symptomatic VTE have PE
- > 50% have silent VTE
- · 1 month mortality
 - DVT = 6%, PE = 12%
- In pts post-trauma or post-orthopedic surgery to lower limbs, VTE risk > 50% w/o prophylaxis

Signs, Symptoms, Diagnosis

- DVT
 - Most likely to form in lower extremity
 - Leg pain, swelling, cyanosis, pain, warmth
 - D-Dimer, doppler ultrasonography, and clinical exam used to diagnose DVT
- PE
 - SOB, tachypnea, tachycardia, hemoptysis
 - D-Dimer, CT scan, Wells Criteria, and clinical exam used to diagnose PE

VTE Complications

- Post-Thrombotic Syndrome (PTS)
 - 2/2 vein damage from thrombosis, resulting in chronic venous insufficiency
 - S/S: pain, swelling, skin discoloration, ulceration $\,$
- Recurrent VTE
- DVT progression to PE
- Death

VTE Prevention

- · CHEST guidelines!
- Identify pts at risk for VTE, classify risk
- Non-pharmacologic
 - Graduated Compression Stockings (GCS)
 - Intermittent Pneumatic Compression (IPC)
 - Early mobilization post surgery
 - Inferior vena cava filter (IVCF)



VTE Prevention

- · Pharmacologic Prophylaxis
 - SQ: LMWH, Fondaparinux, UFH
 - PO: warfarin, dabigatran, apixaban, rivaroxiban
 - THA/TKA
 - 1st line: LMWH
 - 2nd line: fondaparinux, UFH, dabigatran, apixaban, rivaroxiban
 - 3rd line: ASA, warfarin
 - All other pts: LMWH or UFH as 1st line
 - Factors to consider: pt preference, compliance, dose frequency, price, availability..

VTE Treatment

- · CHEST guidelines!
- · Goals of therapy
 - Prevent progression of thrombus, embolization, death
- Prevent long-term complications of VTE/recurrence
- · Principles of therapy
 - Minimal symptoms and low risk for extension (i.e., superficial vein thrombosis): monitor, no need for AC
 - PE with hemodynamic instability: consider lytics
 - PE without hemodynamic instability, or DVT: initiate parenteral AC
 - Transition to oral therapy for long-term treatment
 - Stop AC when clot resolves and per guideline recommendations

VTE Treatment DVT/Hemodynamically Stable PE

- 1. Initiate parenteral AC ASAP, continue ≥ 5d
 - LMWH and fondaparinux preferred over UFH
 - LMWH: enoxaparin 1.5 mg/kg SQ QD or 1 mg/kg BID
 - Fondaparinux: 7.5 mg SQ QD, with dose adjustment required for pt weight
 - UFH: SQ or IV infusion to target therapeutic aPTT
- 2. Initiate PO warfarin as early as possible
 - Usual starting dose 5 mg/d, unless confounding factors (see warfarin section)

VTE Treatment DVT/Hemodynamically Stable PE

- 3. Stop parenteral AC once INR >2 x24h, and after at least 5d of parenteral AC
- 4. Continue warfarin for 3 mo for most pts, with careful monitoring
- 5. Some pts require > 3 mo of warfarin (i.e. based on risk for recurrence or bleeding)
- 6. In patients with cancer and VTE, LMWH is preferred over warfarin for the entire duration of treatment

VTE Treatment Hemodynamically Unstable PE

- In pts with PE associated with HoTN (i.e. SBP < 90 mmHg) with low bleeding risk, thrombolytic therapy is recommended
- Once thrombolytic therapy is complete, treat as hemodynamically-stable PE

VTE Treatment Pharmacotherapy: UFH

- IV continuous infusion for VTE dosed per wt, although also available in IV bolus and SQ
- Dose is adjusted per aPTT
- Generally, aPTT is measured a baseline, 6h after initiation, and 6h after any dose change
- No renal adjustment necessary
- Major AEs: bleeding (aPTT-related), HIT (non-aPTT-related)
- C/I: active bleeding, history of HIT
- · Antidote: protamine sulfate
- · Monitoring: PLT, aPTT, INR, Hg

Heparin-Induced Thrombocytopenia

- Immune-mediated reaction resulting in thrombocytopenia and possible thrombosis at the same time
- · Not related to AC intensity
- 4 Ts: <u>Thrombocytopenia</u>, <u>Thrombosis</u>, <u>Timing</u>, other causes for <u>Thrombocytopenia</u>
- PLT drop > 50% of baseline or < 100,000
- Onset: 5-14 d
- Compare with HAT

Heparin-Induced Thrombocytopenia Management

- D/C all sources of heparin, do not re-challenge
- Initiate alternative AC immediately unless C/I (lepirudin, argatroban..) and regardless of thrombosis presence
- Initiate warfarin once PLT > 150,000 /mL
 - Otherwise high risk for venous limb gangrene and warfarin induced skin necrosis
 - Reverse warfarin if already started
- · Avoid PLT transfusion unless actively bleeding
- For px in pts with HIT, fondaparinux can be used

VTE Treatment Anticoagulants

- · General principles:
 - AC do not lyse a clot, they only stop its growth and propagation
 - Labs that you'll need to monitor at baseline and periodically after: Hg, PLT, INR, aPTT, Cr with certain drugs

VTE Treatment Pharmacotherapy: LMWH

- Enoxaparin, dalterparin, tinzaparin
- Smaller heparin fragments than UFH
- Given SQ
- At least as safe and effective as UFH for VTE, more effective in THA/TKA surgeries
- Enoxaparin
 - Prophylaxis: 40 mg QD or 30 mg BID
 - Treatment: 1.5 mg/kg QD or 1 mg/kg BID
 - Requires renal adjustment CrCl < 30 mL/min

VTE Treatment Pharmacotherapy: LMWH

- No therapeutic monitoring required except in special populations (large or small size, CKD..)
- Cr should be checked at baseline
- · Can be safely used at home-convenient
- · AEs: HIT (10x less likely than UFH), bleeding
- C/I: HIT, active bleeding
- Antidote: protamine sulfate (partial efficacy)

VTE Treatment

Pharmacotherapy: Factor Xa Inhibitors

- Fondaparinux, rivaroxaban, apixaban
- Only fondaparinux and rivaroxaban approved by FDA for treatment of VTE
- Synthetic, do not cross-react with heparins, ideal in pts with HIT history or heparin allergy
- · No therapeutic monitoring required
- · Not reversed by protamine
- Fondaparinux dosing: (SQ)
 - Prophylaxis: 2.5 mg QD
 - Treatment: < 50 kg: 5 mg QD; 50-100 kg: 7.5 mg QD; > 100 kg: 10 mg QD

VTE Treatment

Pharmacotherapy: DTIs

- Dabigatran (PO), lepirudin (IVI), bivalirudin (IVI), argatroban (IVI), desirudin (SQ)
- Dabigatran is the only PO DTI, requires no therapeutic monitoring, and has quick onset
 - Requires at least 5d of parenteral anticoagulation before initiation
- Lepirudin is cleared renally, argatroban hepatically. Both used in pts with HIT
- Bivalirudin is used in PCI
- · Desirudin is used SQ for VTE Px after THA
- DTIs have no antidote and falsely elevate INR

VTE Treatment

Pharmacotherapy: Warfarin

- Inhibitor of Vitamin K-dependent coagulation factors (II, VII, IX, X, protein C, protein S)
- No effect on existing factors- anticoagulation will start when factors eliminated (5-10 d)
- · Metabolized by CYP450 2C9
- Dosed to target INR which takes several doses to reach
- Acute drop in protein C before depletion of clotting factors results in paradoxical hypercoagulable state in the first few days of therapy

VTE Treatment

Pharmacotherapy: Warfarin

- Dosing and bleeding risk depends on environmental and genetic factors
 - Age, nutritional status, liver disease, pt wt, hyperthyroidism, genetic polymorphisms, prior used doses, FAMES (<u>Fluconazole</u>, <u>Fluoroquinolones</u>, <u>A</u>miodarone, <u>M</u>etronidazole, <u>E</u>rythromycin, <u>S</u>ulfas), rifampin, phenobarbital, phenytoin, vitamin K, NSAIDs/antiplatelets/anticoagulants, gut flora modifiers, general health status, infections...

VTE Treatment

Pharmacotherapy: Warfarin

- Dosing strategy:
 - 1. Obtain baseline labs (CBC, INR, aPTT)
 - 2. Initiate 5 mg PO QD for most pts
 - Avoid loading dose
 - Initiate at lower or higher doses according to your pt evaluation
 - 3. Obtain INR more frequently during initial dosing, less frequently once therapeutic on stable dose
 - Example for initial phase: QD if in hospital setting; twice a week if in outpatient setting
 - Example for late phase: every other day if in hospital setting; every 2-4 wks if in outpatient setting

VTE Treatment Pharmacotherapy: Warfarin

- Dosing strategy/Cont:
 - Make dose adjustments in small increments (0.5-1 mg per adjustment) until INR becomes therapeutic
 - Normal pace to achieve therapeutic INR is 5-7d
 - If INR starts rising too fast, lower dose
 - 5. Pick a chronic dose, follow-up on INR less frequently, adjust dose if necessary
 - Educate pt on importance of diet consistency, need to avoid drug interactions, avoiding injuries and bleeding risks, ways to recognize bleeding..

VTE Treatment Pharmacotherapy: Warfarin

- DTIs falsely elevate INR
- ΔFc
 - Warfarin-induced skin necrosis and venous limb gangrene
 - Bleeding
 - Teratogenicity (Category X)
- Antidote:
 - Vitamin K (1-10 mg IV/PO x1)
 - SQ/IV: peaks in 12h, given in urgent cases with bleeding
 - PO: peaks in 24h, given in less urgent cases
 - No reversal necessary if INR < 10 with no bleeding

VTE Treatment Pharmacotherapy: Thrombolytics

- Can be used in pts with PE and hemodynamic instability
- High bleeding risk- careful pt selection needed
- Streptokinase, urokinase, t-PA are FDAapproved for PE treatment
- All are equally efficacious, t-PA has shortest infusion time