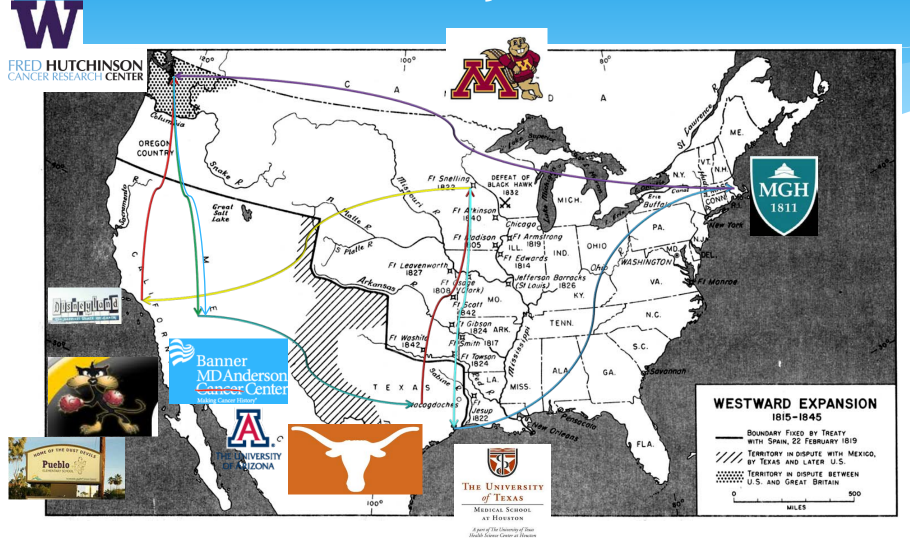


Blood and Clots

Matthew Ulrickson, MD
Banner MD Anderson Cancer Center
Matthew.Ulrickson@bannerhealth.com
September 12, 2017

Where are you from?



Words to live by

* "If you don't take care of yourself and the people you love, you won't be able to take care of strangers"

- Dr. Hasan Bazari

* "Take care of yourself, take care of your teammates, the rest takes care of itself"

- Coach Eddie Reese



Objectives

- Discuss case-based approach to patients with coagulopathy – both acquired and inherited
- * If interested for background:

Coagulation Pathways (1):

https://drive.google.com/file/d/0B8U1h90l29uGbXZ4Umx3Mngtdm8/view?usp=sharing

Platelets (2):

https://drive.google.com/file/d/0B8U1h90l29uGaHh6NkxFM2JIQnM/view?usp=sharing

Bleeding Thrombotic (3):

https://drive.google.com/file/d/0B8U1h90l29uGdFhBTzh5ZU5VSmM/view?usp=sharing



The Bleeding History

- * 1. Have you or a relative ever been told you had a bleeding problem? Bleeding after surgery? After dental work? With trauma? During childbirth or had heavy menses? Have you ever had bruises with lumps?
- * 2. Have you ever required a blood transfusion or had abnormal blood counts? Do you have liver disease?
- * 3. Are you currently taking or have you recently taken anticoagulation or antiplatelet medications (warfarin, heparin, aspirin, NSAIDs, clopidogrel)?

Concerning Bleeding symptoms

- * Have you ever had any of the following symptoms?
 - * Bleeding from trivial wounds <u>lasting >15 minutes</u> or **recurring spontaneously** during the 7 days after the injury?
 - * Heavy, prolonged, or recurrent bleeding after surgical procedures?
 - * Bruising with minimal or no apparent trauma, especially if you could **feel a lump under the bruise**?
 - * Spontaneous nosebleed lasting >10 minutes or that required medical attention?
 - * Heavy, prolonged, or recurrent bleeding after dental extractions that required **medical attention**?
 - * Blood in your stool that required medical attention and was unexplained by an anatomic lesion (stomach ulcer, colon polyp)?
 - * Anemia that required a **blood transfusion** or other type of treatment?
 - * Heavy menses characterized by **clots >1 inch** in diameter, changing a pad or tampon **more than hourly**, or resulting in **anemia** or low iron?

Categorize Bleeding Symptoms

- Characterize bleeding
 - * Superficial (mucocutaneous) vs. deep (muscle/joint)
 - Primary Hemostasis (plt, vWF)

- Coagulation factors
- * Spontaneous vs. Secondary (trauma, surgery, tooth extraction, menses, pregnancy/post partum)
- * Immediate vs. delayed
- Acute (acquired) vs. lifelong (hereditary)
- Family history (X-linked/autosomal)
- * Medications (e.g. aspirin, warfarin, EtOH)
- * Comorbid disease (liver disease, uremia, malignancy)



Case 1-Presentation

- 22-year old man presents to the ED
- Spontaneous knee and hip pain; similar to prior episodes. Also RLQ pain
- No prior surgeries
- Maternal grandfather died of bleeding complications
- Exam: Chronic knee & elbow joint deformities, RLQ pain worse with leg straight

First Test

* What is the first test to order based on this history to evaluate risk of bleeding diathesis?

Case 1 - Laboratory Results

Normal Value			, 1				
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Platelet count 250,000/μl 150 – 400,000/μl

Fibrinogen 300 mg/dl 150 – 400 mg/dl

Prothrombin time 11 sec 11 – 13.6 sec

(INR=0.8)

Partial thromboplastin time 130 sec 24 – 36 sec



Case 1 - Laboratory Results

Normal	1/2	1100
NOTHA	val	iues

Platelet count 250,000/µl

150 – 400,000/µl

Fibrinogen 300 mg/dl

150 – 400 mg/dl

Prothrombin time 11 sec

11 – 13.6 sec

(INR=0.8)

Partial thromboplastin time 130 sec

24 – 36 sec

1:1 mixing study leads to correction of PTT to 26 sec



Case 1 Laboratory Results

Specific Factor Activity Assay:

Normal Range

Factor VIII:C = 90%

50 – 150%

Factor IX:C = < 1%

50 - 150%

What is the diagnosis?



Case 1 Diagnosis of Hemophilia



Inheritance: X-linked recessive (no male/male transmission)

Severity: Varies between families/mutations; ~ half severe

(corrects with 1:1 mixing)

Confirm with genetic testing

Specific:

Clotting activity

Α δ

В

↓ FIX:C



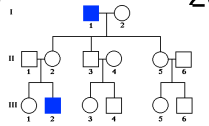
↓ (normal VWF:Ag)

Frequency

75-80%

↓ FVIII:C

20-25%



Cryo contains FVIII but must use FFP for FIX

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Treat by replacing missing factor with recombinant product

Emerging therapy for hemophilia

TABLE III. Gene Therapy Trials for Hemophilia A and B

Product	Manufacturer	Vector	Promotor-transgene	Clinical trial status	Trials identifier	Human response
Factor VIII BMN270	BioMarin	AAV5	Active Factor VIII with B domain deletion of Refacto TM	Enrollment suspended	NCT02576795	8 patients dosed; 6 at high dose level, 6 × 10 ¹³ vg/kg, with FVIII activity level of 4-60% with maximum of 16 weeks of follow-up, prophylactic corticosteroids initiated with patient 4 and beyond
Factor IX AAV8-hFIX19	Spark Therapeutics	ssAAV8	HCRhATT-hFIXco	Enrollment completed	NCT01620801	No study results posted
SPK-9001 (SPX-FIX)	Spark Therapeutics/Pfizer	novel AAV vector	high specific activity FIX variant	Enrolling	NCT02484092	4 patients dosed at initial dose level (5 \times 10 11 vg/kg) with FIX activity ranging from 26 to 41% with 7 to 26 weeks of follow up
scAAV2/8-LP1-hFIXco	St. Jude Children's Research Hospital	scAAV2/8	LP1-hFIXco	Enrolling	NCT00979238	10 patients dosed in 3 dosing cohorts, mean steady state FIX activity 2.9-7.2% with follow-up of at least 16 months.
AskBio009 (BAX 335)	Shire	scAAV8	TTR-FIXR338L (Padua)	Enrollment closed	NCT01687608	7 patients dosed in 3 dosing cohorts, 2 patients with transient FIX activity > 50%, only 1 persisted (medium dose cohort, 1 × 10 ¹² vg/kg)
DTX101	Dimension Therapeutics	AAVrh10	hFiX	Enrolling	NCT02618915	No study results posted
АМТ-060	UniQure	AAV5	LP1-hFIXco	Enrolling	NCT02396342	5 patient enrolled in initial dose level (5×10^{12} gc/kg), two with at least week 12 follow-up had FIX expression levels of 4.5–5.5%
SB-FIX	Sangamo Biosciences	AAV2/6	ZNF mediated gene editing Three components of SB-FIX (ZFN1, ZFN2, and FIX cDNA donor)	Anticipated	NCT02695160	No study results posted

Hartmann and Croteau. AmJ Heme 2016. 91:12:1252

Case 1 Family Testing

- 20-year old sister's factor IX:C = 60%
- DNA: Factor IX gene heterozygous for brother's hemophilic nonsense mutation

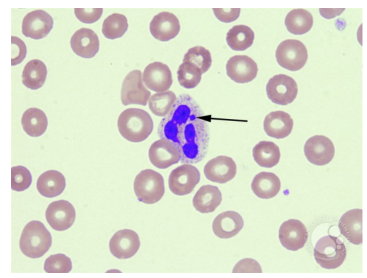




Image Courtesy of Jon Fukumoto

Case 1 Family Testing

- 20-year old sister's factor IX:C = 60%
- DNA: Factor IX gene heterozygous for brother's hemophilic nonsense mutation

Females can have symptoms of mild hemophilia based on X-inactivation pattern

> Usually must have factor <40% to have bleeding symptoms



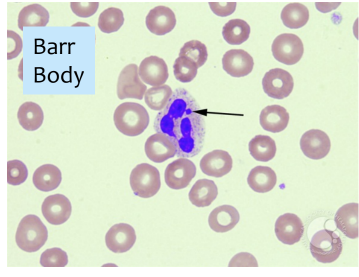


Image Courtesy of Jon Fukumoto

Case 2 - Presentation

- 30yo male physician, presents with melena, UGI bleed
- PMHx: transfused at 15yo for spontaneous GI bleed; oozed 5 days post prior tooth extraction
- Father with history of abnormal bleeding
- Upper endoscopy is negative for focal lesion



Case 2 Laboratory Results

<u>patient</u>

normal values

Platelet count = $250,000/\mu$ l

150 – 400,000/µl

Prothrombin time = 12 sec

11 - 13.6 sec

(INR=1.0)

Partial thromboplastin time = 58 sec

 $24 - 36 \sec$

Thrombin time = 20 sec

Fibrinogen = 294 mg/dl

18 – 28 sec

150 – 400 mg/dl



Case 2 Laboratory Results

<u>patient</u>

normal values

Platelet count = $250,000/\mu$ l

150 – 400,000/µl

Prothrombin time = 12 sec

11 - 13.6 sec

(INR=1.0)

Partial thromboplastin time = 58 sec

 $24 - 36 \sec$

Thrombin time = 20 sec

 $18 - 28 \sec$

Fibrinogen = 294 mg/dl

150 – 400 mg/dl



Mixing time corrects PTT to 27 sec

Case 2: vWF Roles in Hemostasis

1. Enhance platelet function:

platelet adhesion to vascular endothelium

- binds to platelet membrane glycoprotein lb
- depends upon high mol wt VWF multimers

2. Facilitate coagulation:

binds & stabilizes circulating FVIII

- depends upon amino-terminal VWF residues



vWD is the most common bleeding disorder

Present in ~15% of women who undergo hysterectomy for menorrhagia (without structural cause)

Case 2 - Diagnosis of vWD

Clinical: varies from mild, type 1, to severe, type 3

Laboratory:

<u>Screen</u>

Specific assays

1. Platelet function

Tbleeding time plt function

↓ vWF:Antigen (except type 2)
 ↓ vWF activity (except 2N)
 (ristocetin cofactor assay)

2. FVIII activity

↑PTT

mild ↓ Factor VIII:C level

Specific subtype: VWF multimer analysis/genotype (types 2A/B)



vWD Subtypes

Туре	Inheritance	Deficiency
Type 1	Autosomal dominant	Quantitative
Type 2	Autosomal dominant	Qualitative
Type 3	Autosomal recessive	Severe/absent

Case 2 - Specific Assay Results

<u>patient</u>

normal values

vWF antigen level = 30%

50 – 150%

Ristocetin cofactor assay = 25% (vWF activity)

50 - 180%

FVIII:C activity = 20%

50 - 180%

Multimer analysis: normal pattern



Treatment of VWD

- DDAVP (des-amino-D-arginine vasopressin)
 - stimulates VWF/FVIII vascular endothelial release
 - useful to treat or prevent bleeding in mild VWD
 - not helpful in VWD type 2B
- vWF containing FVIII concentrates (e.g. Humate-P)
- vWF concentrates (recombinant completed ph III trial)
- Cryoprecipitate, can use if concentrate not available



Case 3 - Presentation

- 60yo man presents with thigh hematoma
- No prior bleeding history
- * No family history of bleeding
- Prior diagnosis of rheumatoid arthritis



Case 3 Laboratory Results

<u>patient</u>

normal values

Platelet count

 $= 250,000/\mu l$

150 – 400,000/µl

Prothrombin time

= 12 sec

11 - 13.6 sec

(INR=1.0)

Partial thromboplastin time = 100 sec

 $24 - 36 \sec$

Thrombin time

= 20 sec

 $18 - 28 \sec$

Fibrinogen

= 294 mg/dl

150 – 400 mg/dl



Case 3 Laboratory Results

patient

normal values

Platelet count = $250,000/\mu$ l

150 - 400,000/µl

Prothrombin time = 12 sec

11 - 13.6 sec

(INR=1.0)

Partial thromboplastin time = 100 sec

24 – 36 sec

Thrombin time = 20 sec

 $18 - 28 \sec$

Fibrinogen = 294 mg/dl

150 - 400 mg/dl

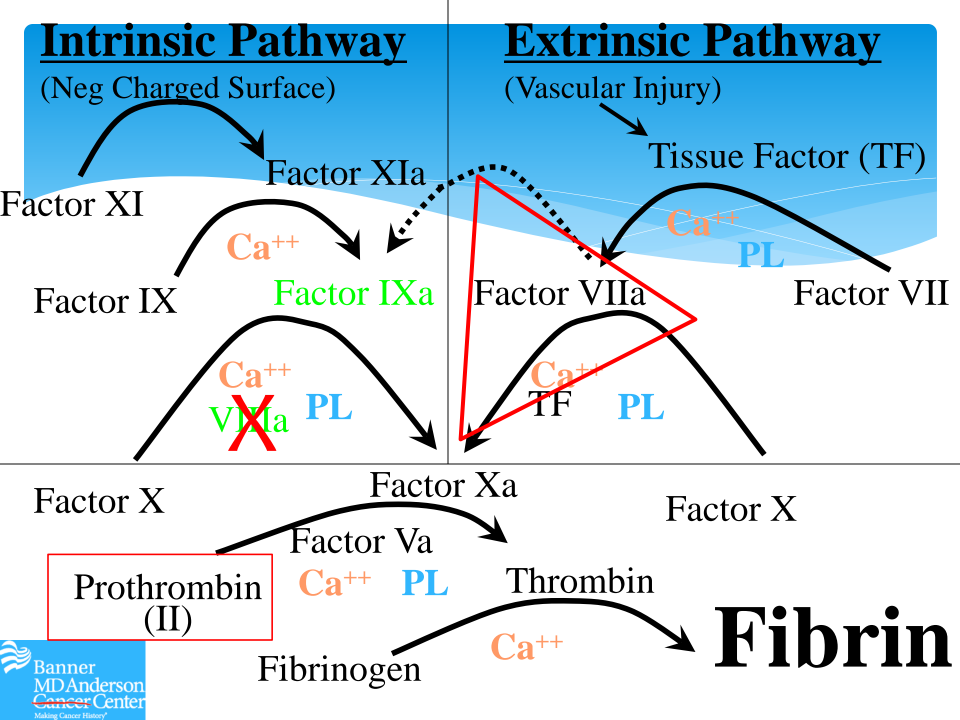
1:1 Mixing initially corrects the PTT to normal, but at one hour the incubated PTT returns to 100 sec



Factor VIII Inhibitors

- * Measured in 'Bethesda units'
- Consume Factor VIII 'acquired hemophilia'
- * Associated with autoimmune and malignant diagnoses, can also <u>rarely</u> occur post-partum
- Significant morbidity and mortality associated
- * Treat bleeding with bypass agents (rFVIIa or prothrombin complex concentrate (PCC))
 - FFP will not correct coagulopathy from inhibitor
- * Treat inhibitor with immune suppression (steroids, rituximab)





Case 4

- * A 66yo alcoholic man presents with hematemesis
- He has a prior history of IVDU and hepatitis C
- * On exam he is icteric with palmar erythema, spider angiomata, gynecomastia, and caput. He has very limited peripheral veins noted on exam
- * HR 115 BP 96/42
- * CBC 2.4>7.1<42 ANC 1200
- * Albumin 2.1 INR 2.8 PTT 65 sec

Case 4

- * A 66yo alcoholic man presents with hematemesis
- He has a prior history of IVDU and hepatitis C
- * On exam he is icteric with palmar erythema, spider angiomata, gynecomastia, and caput. He has very limited peripheral veins noted on exam
- * HR 115 BP 96/42
- * CBC 2.4>7.1<42 ANC 1200
- * Albumin 2.1 INR 2.8 PTT 65 sec

What additional hematologic test would you order?

Case 4 - Cirrhosis

- * Fibrinogen = 65 (thrombin time 37 sec (18-28sec))
 - Decreased production
 - * Abnormal function (increased thrombin time)
 - Level <75 can spuriously increase the INR and PTT</p>
 - Treatment: Replacement with cryoprecipitate for level
 <100

Liver Disease and Hemostatic Defects

Screening Test Result

Platelets

- Thrombocytopenia

Coagulation

- Prolonged PT & PTT
- Prolonged thrombin time
- Low fibrinogen

Etiology

```
    ↓thrombopoietin (made by liver)
    Folate deficiency (possible)
    Toxic EtOH effects
    ↑ splenic pooling (splenomegaly)
```

- ↓ vitamin K-dependent carboxylation
- ↓ factor synthesis (II,VII,IX & X)

Dysfibrinogenemia

- ↓ FDP clearance
- ↓ synthesis



Remember – dysfibrinogenemia can lead to normal level, but inadequate activity

Case 4 - cont

- * The nurse informs you that they are unable to get peripheral access.
- * What do you recommend?

Can you place a line?

- * Prospective study (N = 658) of patients with liver disease and coagulopathy
- * All underwent CVC insertion
- * 1 major bleeding complication (hemothorax) due to inadvertent subclavian artery puncture.
- * Average INR of patients was 2.4; all thrombocytopenic
- * Rates of superficial hematoma and ooze were increased compared to other populations, though these correlated more with number of passes required and ease of guidewire insertion than with INR or platelet count.
 - * Intensive Care Med (1999) 25: 481-485

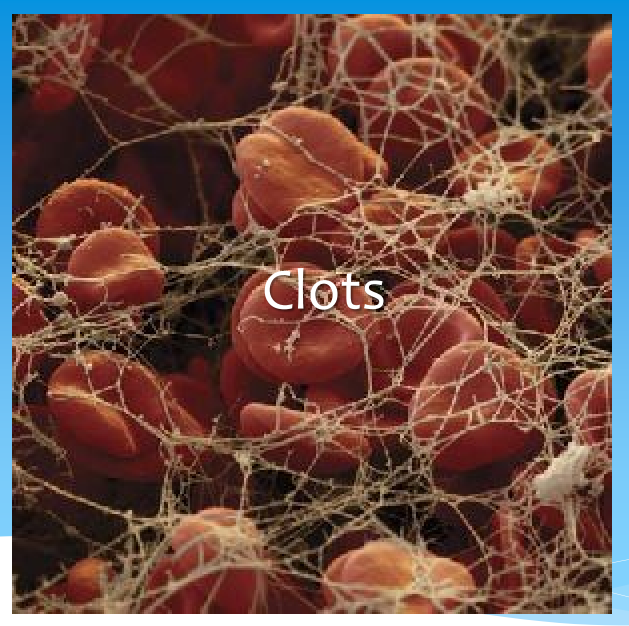


How about IR?

- * Tunneled lines placed in interventional radiology
 - * at least 25k platelets
 - * INR less than 2.0
 - * N=626 with either platelets <50k, INR >1.5, or both
 - No bleeding complications noted
- * J Vasc Interv Radiol 2010;21:212–217

Transfusion Recs

- Platelets (usually last 3-5 days)
 - For major bleeding or on anticoagulation, >50k
 - For minor bleeding (epistaxis, gum bleeding) >30k
 - * With no bleeding >10k (Stanworth, NEJM 2013. 368:1771)
- * FFP
 - * If active bleeding or need for procedure and INR >2
 - * Effects wane after 4 hours, so must time procedure well
 - * This often precludes a 'check then send' approach unless sent stat and procedure team immediately available
 - If no bleeding, no FFP regardless of INR
 - * (*possibly for anticoagulation reversal)
- * Cryo
 - * 1 unit per 10kg body weight for fibrinogen <100 in setting of bleeding</p>



Discuss case-based approach to patients with thrombophilia

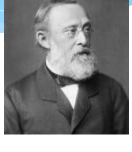
'Provoked' DVT

* Immobility/Stasis



* Trauma (surgery)





Dr. Rudolf Virchow 1821-1902

- * Hypercoagulable
 - * Malignancy
 - * Age-appropriate cancer screening
 - * Hormones
 - * Pregnancy, OCP/HRT



Abrofa

Dr. Armaund Trousseau 1801-1867



SOME study (Screening for Occult Malignancy in VTE) NEJM 2015

New unprovoked VTE

Limited Screening

Limited + CT Abd/Pelvis

- * PSA*
- * Mammogram*
- * CXR
- * Blood work
- * Pap smear*

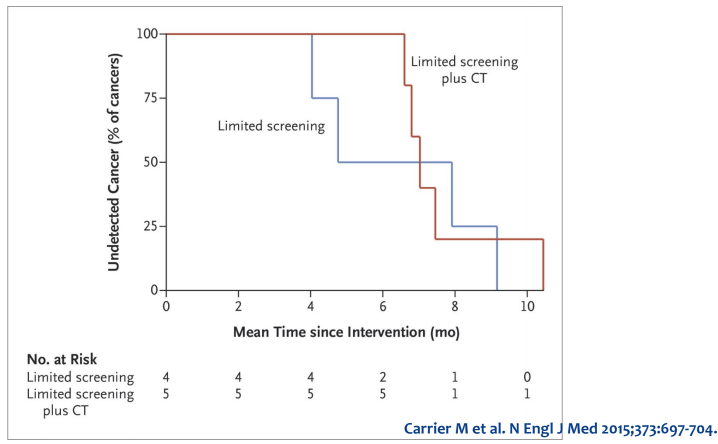
Carrier M et al. N Engl J Med 2015;373:697-704.

Primary endpoint: cancer missed by the screening but detected within 1 year of screening

SOME study (Screening for Occult Malignancy in VTE) NEJM 2015

New unprovoked VTE

Kaplan-Meier Curves for Time to Detection of Missed Occult Cancer.



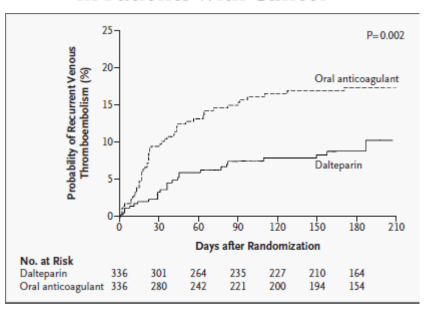


CLOT

- * Dalteparin vs warfarin
 - Randomized, N=676
 - * 6 months of treatment
 - * Dalteparin (5 days) → Warfarin
 - * Dalteparin
 - All with active malignancy within 6 months of enrollment
 - * Recurrent VTE
 - * Warfarin 17%, Dalteparin 9%
 - Major Bleeding
 - * Warfarin 4%, Dalteparin 6%
 - * Any Bleeding
 - * Warfarin 19%, Dalteparin 4%

Lee, et al. NEJM 2003. 349:146

Low-Molecular-Weight Heparin versus a Coumarin for the Prevention of Recurrent Venous Thromboembolism in Patients with Cancer



When to Consider Underlying Hypercoagulable State

- Recurrent <u>unexplained</u> episodes of VTE
- VTE at a young age (<40 years)
- Family history of unprovoked VTE
- Venous thrombosis at an unusual site

 (e.g. axillary vein, mesenteric vein, portal vein)

 American Society of Hematology (ASH) advises against sending hypercoagulable testing in patients with provoked VTE.



Case 1 - Presentation

- * 35yo s/p arthroscopy to her R knee 2 weeks ago
- Presents with RLE swelling and pain
- * RLE DVT is diagnosed and she is started on anticoagulation
- * She is referred to you because recent testing revealed low levels of protein C and protein S, and that she has a gene change in MTHFR
- * What are your recommendations?

When to send hypercoagulable testing (if at all)

Table 7-1 Conditions associated with acquired coagulation factor deficiencies.

Factor	Conditions associated with decreased factor level	
Protein C	Acute thrombosis Warfarin therapy Liver disease Protein-losing enteropathy	
Protein S	Acute thrombosis Warfarin therapy Liver disease Inflammatory states	
Antithrombin	Estrogens (contraceptives, pregnancy, postpartum state, hormone replacement therapy) Protein-losing enteropathy Acute thrombosis Heparin therapy Liver disease Nephrotic syndrome Protein-losing enteropathy	

Table 7-2 Influence of acute thrombosis, heparin, and vitamin K antagonists on thrombophilia test results.

Test	Acute thrombosis	Unfractionated heparin	Low molecular weight heparin	Vitamin K antagonists
Factor V Leiden genetic test	Reliable	Reliable	Reliable	Reliable
APC resistance assay	Reliable*	???*	??? [†]	Reliable*
Prothrombin 20210 genetic test	Reliable	Reliable	Reliable	Reliable
Protein C activity or antigen	???‡	Reliable	Reliable	Low
Protein S activity or antigen	May be low	Reliable	Reliable	Low
Antithrombin activity	May be low	May be low	May be low	Reliable
Lupus anticoagulant	Reliable [§]	????	???ll	May be false positive
Anticardiolipin antibodies	Reliable [§]	Reliable	Reliable	Reliable
Anti-β ₂ -glycoprotein	Reliable [§]	Reliable	Reliable	Reliable
I antibodies				
Homocysteine	Reliable	Reliable	Reliable	Reliable

^{*}Reliable if the assay is performed with factor V-depleted plasma; thus, clinician needs to inquire how the individual laboratory performs the

Although many test kits used for lupus anticoagulant testing contain a heparin neutralizer, making these tests reliable on unfractionated heparin (UF) and possibly low molecular weight heparin (LMWH), clinicians need to ask their laboratory how their individual test kit performs in samples with UF and LMWH.

APC = activated protein C resistance. Must be off VKAs for 2-3 weeks prior to testing PrC, PrS

Tests to never send

- MTHFR gene analysis/polymorphism (33% of population, no increase in VTE risk)
- Homocysteine level (except for pt <30yo to eval for homocytsinuria)

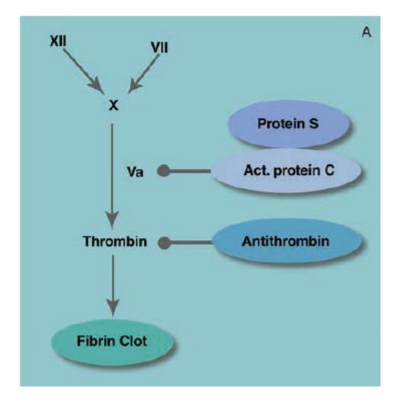
[†]Depending on the way the assay is performed, results may be unreliable; the health care provider needs to contact the laboratory and ask how the specific test performs on heparin.

[‡]Probably reliable, but limited data are available in literature.

⁶Test is often positive or elevated at time of acute thrombosis, but subsequently negative.

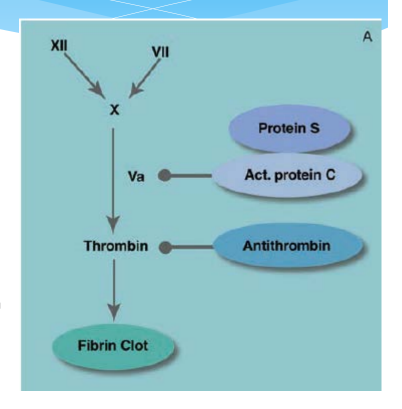
Thrombophilia

Incidence	Venous	Arterial
5%	Factor V leiden Heterozy -2.7x risk Homozy – 18x risk	No significant change
2%	Prothrombin G20210A Heterozy – 3x risk	Possible slight risk in young patients



Thrombophilia

Incidence	Venous	Arterial
5%	Factor V leiden Heterozy -2.7x risk Homozy – 18x risk	No significant change
2%	Prothrombin G20210A Heterozy – 3x risk	Possible in younger patients
0.2%	Protein C deficiency -24x risk	Risk in younger pts
0.1%	Protein S deficiency -31x risk	Risk in younger pts (<55yo)
0.1%	Antithrombin deficiency -30x risk	unclear



Absolute 10yr risk of VTE in FacV Leiden is 1-10% (population risk is 0.1% per year) Protein C and Protein S deficiency has 1% per year risk

Anti-phospholipid antibody

- * Risk for VTE AND arterial events (and pregnancy loss)
- * Diagnose with:
 - Thrombotic event or late pregnancy loss AND
 - * Lab evidence confirmed at least 12 weeks apart (not IgA)
 - * High rate of false-positive, especially in ICU
- * 5-15% rate of 'warfarin failure' (though may be partially due to misleading INR)



Case 2 - Presentation

- * 35yo female presents with abdominal pain and jaundice
- * She has no history of liver disease, heavy EtOH intake, or thrombosis.
- * Exam reveals ascites and RUQ pain, icteric sclerae



Case 2 - Presentation

- 35yo female presents with abdominal pain and jaundice
- * She has no history of liver disease, heavy EtOH intake, or thrombosis. No recent surgery, immobility, trauma, or plane flights.
- * Exam reveals ascites and RUQ pain, icteric sclerae
- * T Bili = 12
- * RUQ ultrasound with doppler reveals portal vein thrombosis.



Additional tests to consider

- Mesenteric/portal vein thrombosis without risk factor (cirrhosis):
 - * JAK2 V617F mutation (~32% of all splanchnic vein thromboses associated with this mutation) (Dentali, Blood 2009, 113:5617)
 - * ***about half of these patients will have abnormal blood counts at time of clot
 - * Flow cytometry to evaluate for PNH (paroxysmal nocturnal hemoglobinuria) (*rare*)
 - * Most of these patients will have intermittent 'hematuria'/hemolysis
 - May also present with cerebral thromboses
 - May also have cytopenias (aplastic anemia, MDS assoc)



Case 3 - Presentation

- * 65yo man admitted to the hospital for pneumonia
- * Hospital day 7 severe increase in respiratory distress
- * Chest CT reveals saddle pulmonary embolism
 - Developed in spite of heparin SC prophylaxis since time of admission



Case 3 - Labs

- * CBC: 13>42%<52k ANC 6.8 Cr 1.0 T Bili 0.2
- * Next test?



Case 3 - Labs

* CBC: 13>42%<52k (platelets 140k on admission)

* Next test?



Case 3 - Labs

- * CBC: 13>42%<52k (platelets 140k on admission)
- * Anti-PF4 antibody: 2.40
- * Interpretation:
 - * Weak-positive OD 0.40-<1.00 low probability (≤5%) of a strong-positive SRA</p>
 - * Strong positive OD ≥ 2.00 units >90% with positive SRA (J of Thromb Hemost 2008. 6(8):1304)
 - * High rate of mild false-positives, especially in setting of acute illness



HIT

- * 4T rule
 - * Timing (within 5-14 days of heparin (~24hrs if recent exposure within 100 days)
 - * Depth of thrombocytopenia <50% baseline (rare to get below 20K)
 - * Thrombosis
 - * No other causes of thrombocytopenia
- * Treatment
 - * Stop heparin
 - * If heparin is stopped without other anticoagulant (in true HIT), ~50% of patients develop VTE within 30 days of diagnosis
 - Start bivalirudin or argatroban (direct thrombin inhibitor)



HIT

- * After stopping heparin, platelets should increase
- * When plt >150k, can transition to warfarin
 - * Must use chromogenic Factor X for transition or stop/start if on argatroban (since it elevates INR)
- * For future prophylaxis fondapariunx is an option (1 case report of HIT)
- * Maturing data on the oral direct anticoagulants (Kunk, PR et al. <u>J Thrombolysis.</u> 2016 Sep 8.)
- If antibody-negative, heparin may be used in the future with close monitoring



Case 4

- 70yo presents with LLE edema and pain after total knee replacement
- Ultrasound confirms L popliteal DVT
- ★ Started on enoxaparin → warfarin
- * Do you recommend ambulation?
- * How long do you recommend anticoagulation?
- * Additional testing?



Provoked DVT

- Following provoked DVT 3 months anticoagulation is adequate (as long as provoking factor no longer present)
- No hypercoaguable testing recommended
- * Ambulation after DVT has not been shown to increase risk of embolization, and decreases risk of post-thrombotic syndrome



Case 5

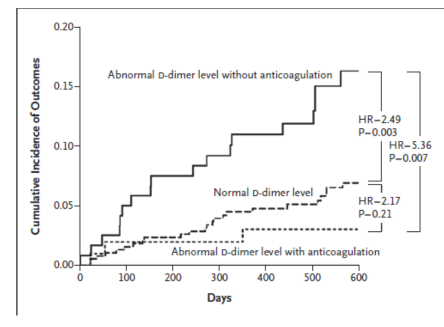
- * 40yo man presents with LLE edema and pain
- * Ultrasound confirms L popliteal DVT
- No recent surgeries, no personal or family history of thrombosis.
- * He drove from Gilbert to Phoenix the day before the event.
- * No chest pain, dyspnea, or palpitations
- He is started on enoxaparin
- * Additional testing at this time?
- * How long do you anticoagulate?



Unprovoked DVT

- * No clear consensus!!
 - But with second event always indefinite
- Two options for first event
 - * Indefinite
 - Attempt to come off at three months for first event
 - * 1 month after stopping anticoagulation perform D-dimer
 - * Elevated: 15% risk of recurrence
 - * Decreased to 2.9% if warfarin is restarted

Normal: 6% risk of recurrence



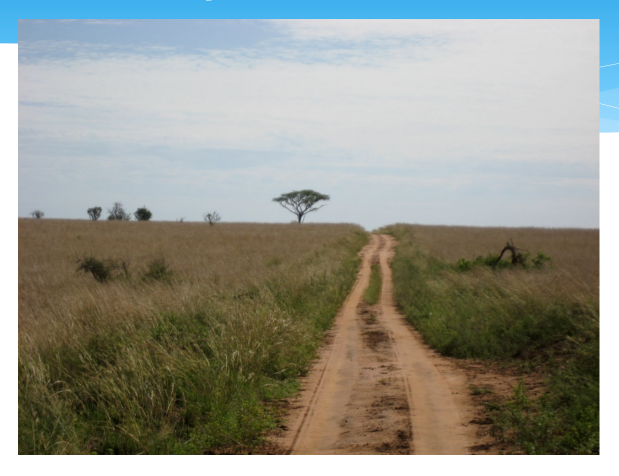
Palareti NEJM 2006



Questions



Questions



American Society of Hematology www.hematology.org



Crossing the Bridge

- Patients with Atrial Fibrillation (did not include DVT)
- * LMWH vs placebo: start 3 days before procedure until 24 hours before procedure and continue for 5 -10 days after the procedure.
- * Warfarin treatment stopped 5 days before procedure and resumed within 24 hours after procedure
- * Incidence of arterial thromboembolism
 - * 0.4% in the no-bridging group
 - * 0.3% in the bridging group
- * Incidence of major bleeding
 - * 1.3% in no-bridging group
 - * 3.2% in the bridging group

Line-associated DVT

- * Incidence of line-associated DVT 6-13%
 - Usually within first 6 weeks after placement
 - * Usually suggested by difficulty drawing and/or infusing through the catheter.
 - *Inability to draw blood alone (i.e. "ball valve effect") is a nonspecific finding and does not predict thrombosis of the catheter lumen or the vessel.



Line-associated DVT

Incidence of line-associated DVT 6-13%

- Usually within first 6 weeks after placement
- Usually suggested by difficulty drawing and/or infusing through the catheter.
 - * Inability to draw blood alone (i.e. "ball valve effect") is a nonspecific finding and does not predict thrombosis of the catheter lumen or the vesse
- Additional risk factors for CVC-associated DVT include:
 - Prior catheter placement and/or upper extremity DVT
 - Catheter malposition (e.g. tip is high in the SVC rather than at the cavalatrial junction
 - Stiffer catheter (e.g. polyethylene vs silastic)
 - * Larger diameter catheter (e.g. indwelling tunneled pheresis catheter)
 - * Line-associated infection
 - Infusion of sclerosing chemotherapy
 - Use of a thrombogenic agent (e.g. thalidomide)
 - * Heparin-induced thrombocytopenia
 - Regional bulky lymphadenopathy
 - * Procoagulant states (Fac V Leiden, PT G20210A)
- * Ultrasound may not detect thrombus in SVC/proximal vessels



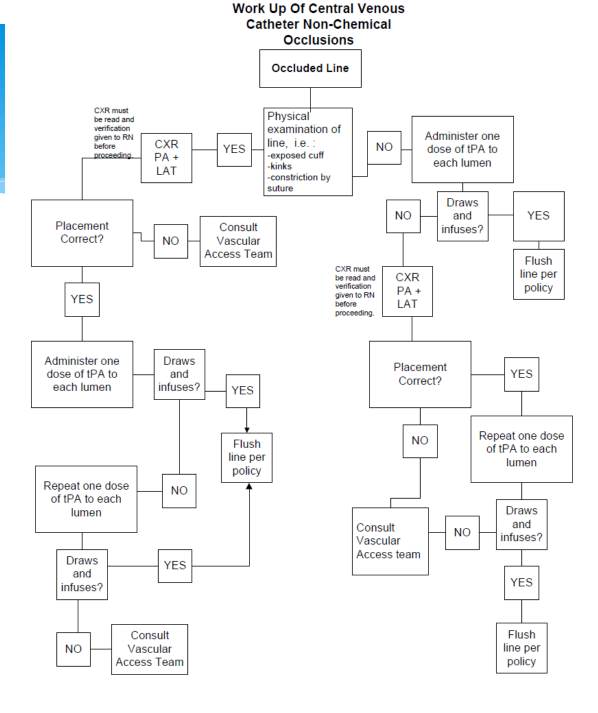
Management of CVC-associated DVT

- * May remove line
 - preferred especially if patient expected to have thrombocytopenia or central vessels affected
 - * If no thrombocytopenia, anticoagulate x 3 months after line removal
- * May treat with anticoagulation without removal if non-occlusive thrombus
 - * Usually 3 month duration



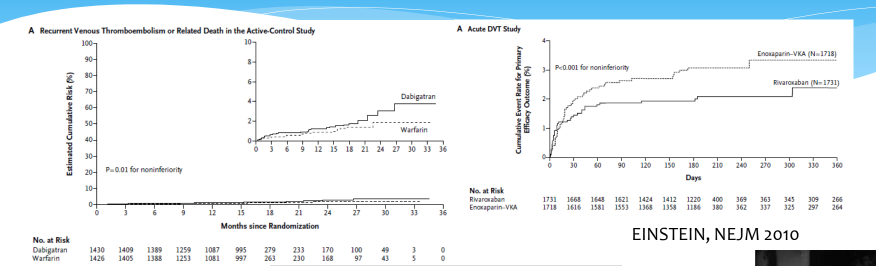
* COOL-2 Trial supports use of tPA in occluded lines

- * JCO 2002. 20:317
- * Restores flow in 87% of lines at 120min following up to 2 doses of tPA

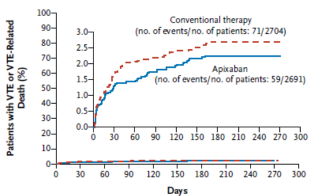




EINSTEIN, RE-MEDY/RE-SONATE, AMPLIFY



Schulman NEJM 2013





Agnelli, NEJM 2013

Warfarin and Cancer Patients



- * More drug interactions
- Less consistent oral intake
- * More variable INR
 - * More bleeding events
 - * More VTE recurrence



Meliotus alba "Sweet Clover"

- * 1920 Bleeding cattle N USA, sweet clover implicated
- * 1940 Karl Link and H Campbell discovered coumarin
- 1948 Warfarin synthesized by Link
- 1952 Approved as rodenticide
- 1954 Approved for human use

Wisconsin Alumni Research Foundation - arin

VTE: Other Anticoagulants

- Dabigatran, anti-thrombin
- Rivaroxaban, anti-FXa only one approved by FDA for DVT/PE treatment
- Apixaban, anti-Fxa
- vs warfarin
 - More rapid onset
 - Uniform dosing (no INR checks) caution with renal dysfunction or morbid obesity
 - No reversal agent
 - Higher cost

Case 4: Presentation

- 23 yo woman, aeronautical engineer
- cc = rash on ankles & shins, easy bruising ~ 10 days rash is not pruritic or painful
- Denies recent contact with new soaps or detergents
- Bruises on her arms & sides, unrelated to trauma
- Also has nosebleeds, gum bleeding with flossing and unusually heavy menses last week
- URI 3 weeks ago, now resolved.
- Exam: no lymphadenopathy, no hepatosplenomegaly stool is guaiac positive

Case 4: Skin Rash



Type of bleeding disorder?

Her signs and symptoms suggest what type of bleeding disorder?

Type of bleeding disorder?

Her signs and symptoms suggest what type of bleeding disorder?

Abnormality of primary hemostasis

Additional History

- Bleeding problems in the past? Procedures or trauma? (include wisdom tooth extracted) None
- What medications are you taking? None
- Do you drink alcohol? If so, how often? No
- Do you use intravenous drugs? No
- Do you have unprotected sex?
- Anyone in your family have a bleeding problem?
- Any recent unexpected loss of weight? No

Laboratory Evaluation

What laboratory tests would you order?

- CBC
- PT & PTT
- TT (Thrombin time)
- Peripheral smear

Laboratory Results

WBC (× 10³/mm³)

 $6.0 \quad (4.3-10)$

• Hgb (gm%)

13.1 (12-16)

Hct (%)

39 (38-50)

MCV (fL)

86 (78-96)

Plt Ct (× 10³/mm³)

3 (150-450)

• PT (sec)

11.6 (10.4 - 12.8)

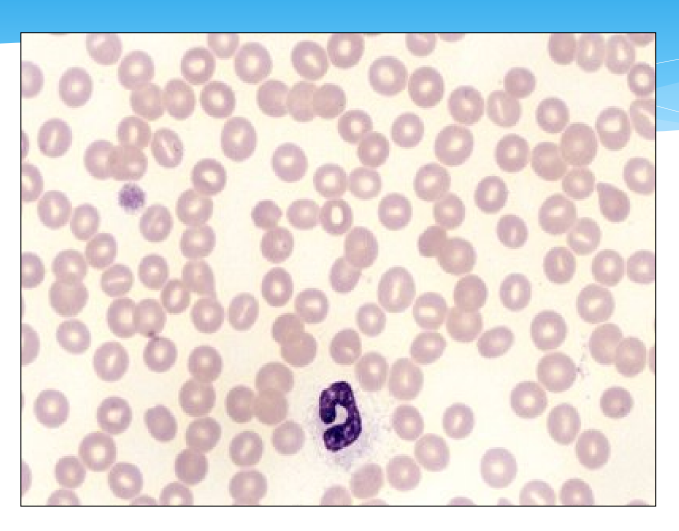
PTT (sec)

32 (24 - 36)

• TT (sec)

22 (18 - 28)

Peripheral smear



Thrombocytopenic Mechanisms

- Decreased production
 - decreased thrombopoietin (liver disease)
 - toxins (e.g. alcohol, radiation, drugs)
 - vitamin B12 or folate deficiency
 - marrow infiltration (malignancy, fibrosis/granuloma)
 - primary marrow disorders (aplastic anemia, myelodysplasia)
 - viral infections (e.g. HIV, HCV)
- Accelerated destruction
 - immune mediated
 - non-immune mediated (DIC, TTP, etc)
- Sequestration
 - hypersplenism

Differential Diagnosis

- Acute leukemia
- Aplastic anemia
- Hepatitis
- HIV
- Auto-immune thrombocytopenic purpura (ITP)
- Systemic Lupus Erythematosus (SLE)

Differential Diagnosis

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Treatment Options

- Platelet transfusion (life-threatining bleed)
 - Since platelets will be consumed as soon as transfused, only do so in setting of active bleeding
- Prednisone
- IV IgG & prednisone
- Anti-RhD immuneglobulin (WinRho)
- Cyclophosphamide
- Splenectomy

One Month Follow Up

- On prednisone 10mg/day
- Difficulty sleeping, marked irritability
- Exam: gained 10 kg, Cushingoid, facial acne
- Bruises anterior tibial legs, few palatal petechiae
- Platelet count = 12,000
- Liver function normal; HIV antibody, negative

What are your next step(s)?

- Increase steroid dose
- Immunization against encapsulated organisms

Second-Line Therapies

- Splenectomy
- Pulse dexamethasone
- Cyclophosphamide
- Anti-CD20 antibody (Rituximab)
- Thrombopoietin mimetic agents
- Other Rx options if above fail:
 - MMF
 - Azathioprine
 - Danazol

Questions?

Case 1 Treatment of Hemophilias

<u>A</u> <u>B</u> .

Therapeutic plasma derived or concentrates rHu FVIII rHu FIX

- recovery (%) 90 35

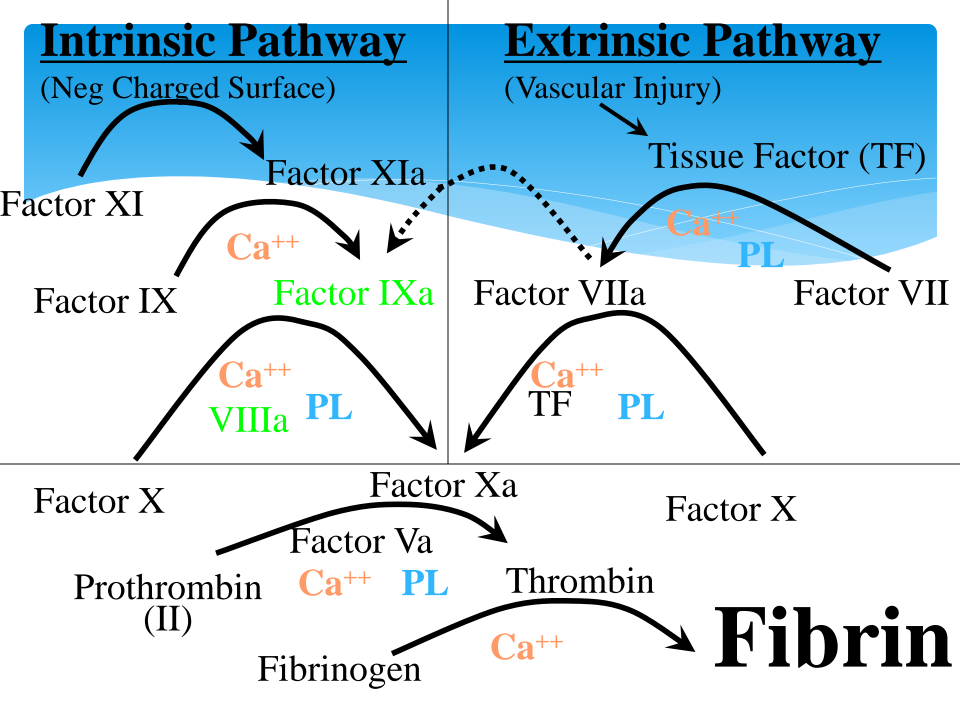
- t_{1/2} (hrs) 8-10 16-24

DDAVP (response) + if mild none

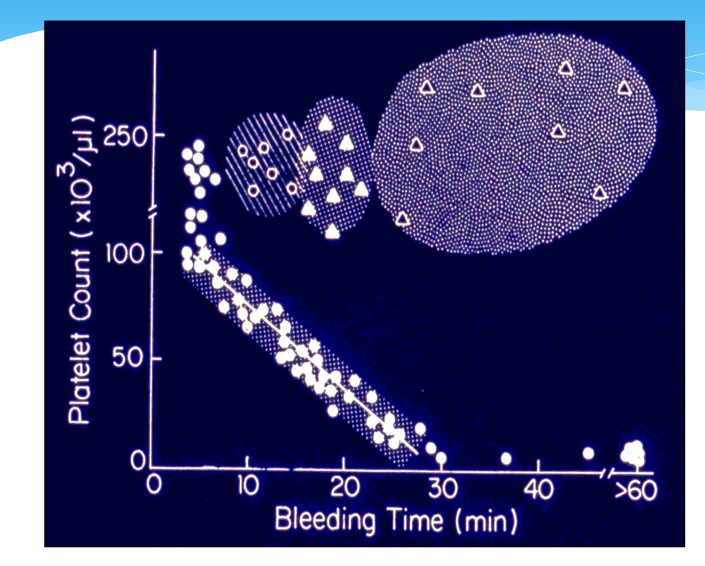
• E-aminocaproic acid minor procedures (EACA, Amicar®) (eg, dental extractions)

None of the above cryoprecipitate FFP

available



H4-4. Relationship Between Platelet Count & Bleeding Time



ASAMildWDsevVWD

H6-7. Laboratory Results

- PT, PTT, TT, fibrinogen = normal
- Antithrombin III, Protein C & Protein S all = normal
- Prothrombin gene = GG20210, homozygous normal
- PTT not prolonged with activated PC → APC resistance
- Factor V gene homozygous 506QQ (Leiden alleles)

H6-8. Recurrent VTE: Congenital Risks

congenital disorder	frequency (%)
Activated Protein C (APC) resistance	20-50
Prothrombin mutation (PT20210A)	10-20
Protein C deficiency	<5
Protein S deficiency	<5
Antithrombin III (ATIII) deficiency	<3
Other (Plasminogen, Dysfibrinogenem	ia) <1

H4-9. VWD Subtypes

inheritance

deficiency

Type 1 Autosomal dominant

Quantitative

2. Type 2 Autosomal dominant

Qualitative

3. Type 3 Autosomal recessive Severe/absent

Dabigatran reversal agent

BRIDGE trial (NEJM 2015)

CT for occult cancer after VTE (NEJM 2015)



Chromogenic Factor X Assay

Chromogenic Factor X	INR
40-25%	2-3
35-20%	3-4

- Chromogenic Factor X levels
 - * >40% indicate a likely sub-therapeutic anticoagulant effect (INR < 2)</p>
 - * <20% indicate a likely supra-therapeutic effect (INR > 3).

Case 5

- * 83yo man with atrial fibrillation presents after a fall. His wife reports that he is on dabigatran.
- * He is confused and has an ecchymosis on the R forehead
- * CT scan reveals an 8mm subdural hemorrhage
- * CBC 5.7>12.5<140
- * Cr 1.5 INR 1.1 PTT 38 sec

What additional testing do you recommend?

Target Specific Oral Anticoagulants

* Target-specific oral Anticoagulant bleeding

Anticoagulant	Mechanism	Laboratory testing
Dabigatran	Direct thrombin inhibitor	Thrombin time elevated
Rivaroxaban	Factor Xa inhibitor	Anti-Xa activity
Apixaban	Factor Xa inhibitor	Anti-Xa activity

Anticoagulant Reversal

GENERIC (BRAND) NAMES	ELIMINATION HALF-LIFE	REMOVED BY HD	STRATEGIES TO REVERSE OR MINIMIZE DRUG EFFECT	
apixaban	8-15 hours	NO	Drug activity can be assessed with anti-factor Xa activity assay	
(Eliquis)	(longer in renal		If ingested within 2 hours, administer activated charcoal	
	impairment)		• Adexanet increase drug clearance; correlation of shortening PT/aPTT with reduction in ble	its) d will not eeding risk is unknown
argatroban	40 – 50 minutes	~ 20%	 Turn off infusion Degree of reversal can be assessed with PTT and/or plasma-diluted thrombin time 	
bivalirudin (Angiomax)	25 minutes (up to 1 hr in severe renal impairment)	~ 25%	 Turn off infusion Degree of reversal can be assessed with plasma-diluted thrombin time 	
dabigatran	14-17 hours		 Drug activity can be assessed with aPTT and/or plasma-dilutime 	ited thrombin
(Pradaxa)	(up to 34 hrs in severe renal impairment)	~ 65%	• Idarucizumab NOTE. FOR may partially correct art trains plasma-unisted thrombin time but we clearance; correlation of lab results with reduction in bleeding risk is unknown	nits) ബ സം: increase drug
Rivaroxaban	Healthy: 5-9 hrs Elderly: 11-13 hrs		Drug activity can be assessed with anti-factor Xa activity	
(Xarelto)	(longer in renal impairment)	NO	Adexanet Increase drug clearance; correlation of shortening PT/aPTT with reduction in) units) y and will not bleeding risk is unknow

Siegal et al. N Engl J Med. 2015;373(25):2413-24