## **Blood Coagulation Basics**



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### **Basic Coagulation Profile**

- Bleeding Time
- Platelet Count
- Prothrombin Time (PT)
- Activated partial Thromboplastin Time (APTT)
- Thrombin Time
- Fibrinogen Assays





#### Hemostasis

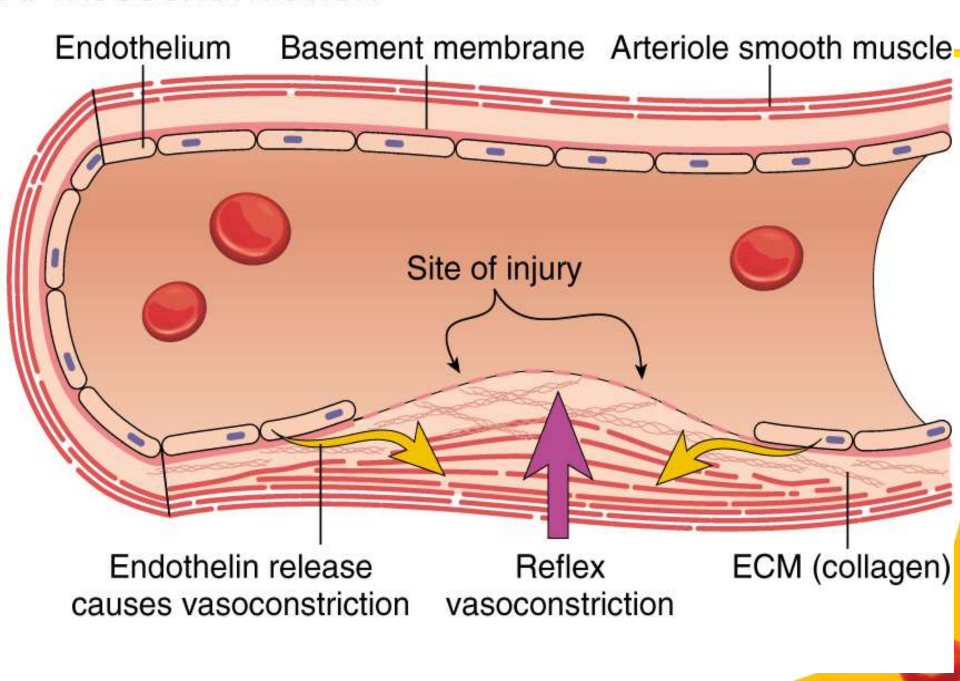
 Hemostasis ("hemo"=blood; stasis= remain") is the stoppage of bleeding, which is vitally important when blood vessels are damaged.

- Three steps:
  - Vasoconstriction
  - Primary Hemostasis- Platelet Plug formation
  - Secondary Hemostasis- Coagulation cascade activation leading to stable Clot formation

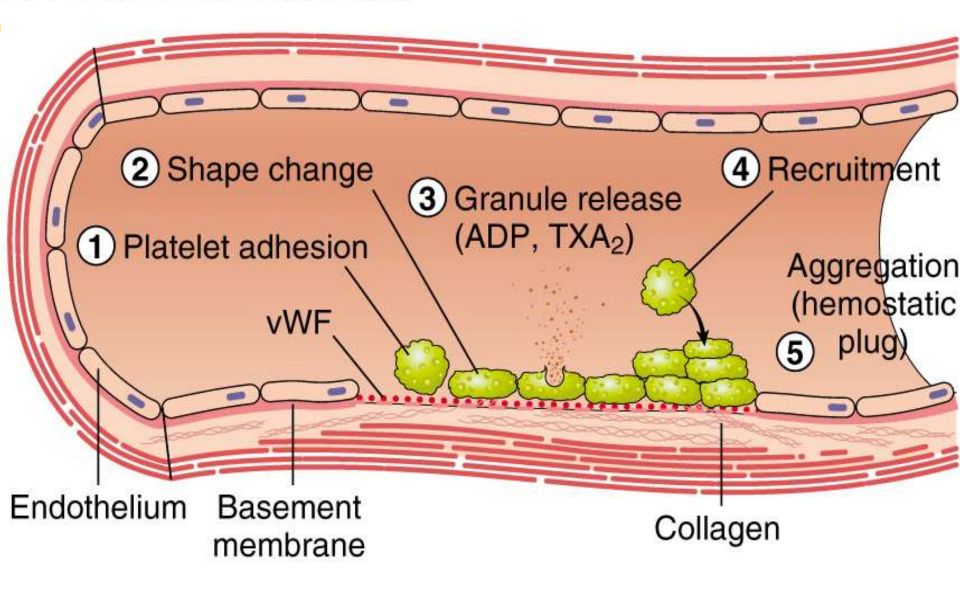




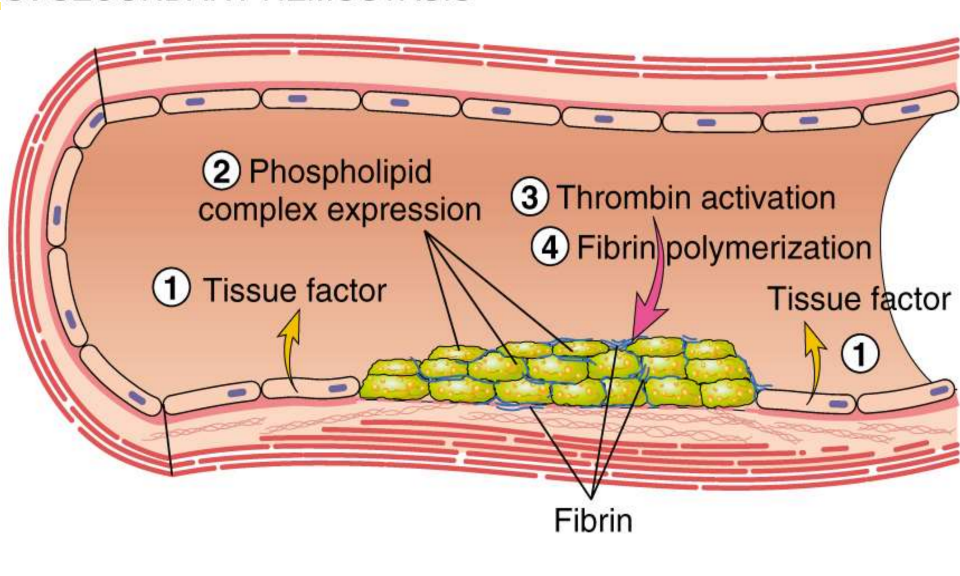
#### A. VASOCONSTRICTION



#### B. PRIMARY HEMOSTASIS



#### C. SECONDARY HEMOSTASIS



#### **CLOTTING FACTORS**

- I (fibrinogen
- II (prothrombin)
- Tissue factor
- Calcium (Factor IV)
- V (proaccelerin, labile factor)
- VI
- VII (stable factor)
- VIII (antihemophilic factor)
- IX (Christmas factor)
- X (Stuart-Prower factor)
- XI (plasma thromboplastin antecedent)
- XII (Hageman factor)
- XIII (fibrin-stabilizing factor)
- von Willebrand factor



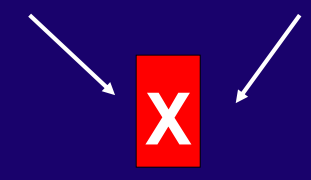


**Intrinsic Pathway** 

**Extrinsic Pathway** 

The PTT Pathway Intrinsic

The PT Pathway Extrinsic



The PT and the PTT pathway meet at factor X, because "X" marks common pathway

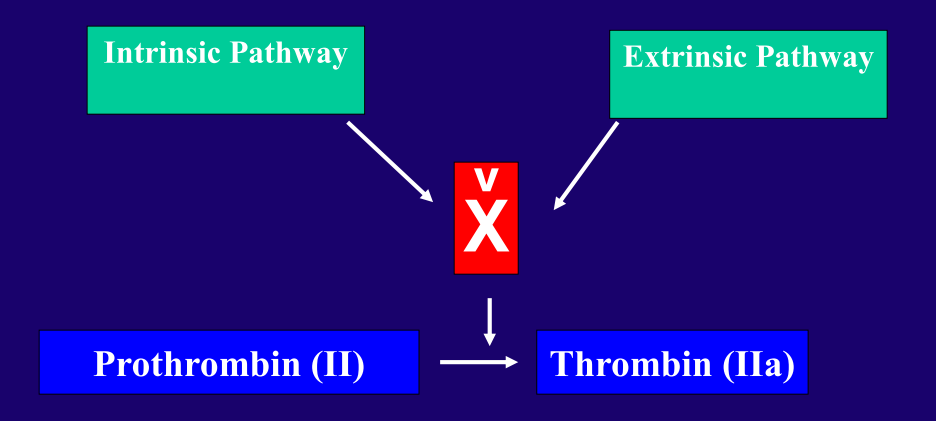
The PTT Pathway
Intrinsic

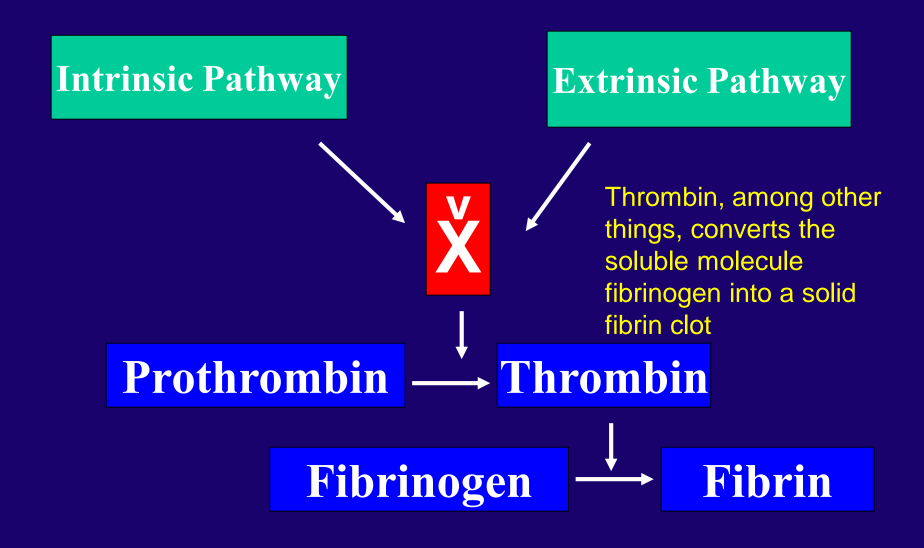
The PT Pathway
Extrinsic





Ca++, Phospholipid



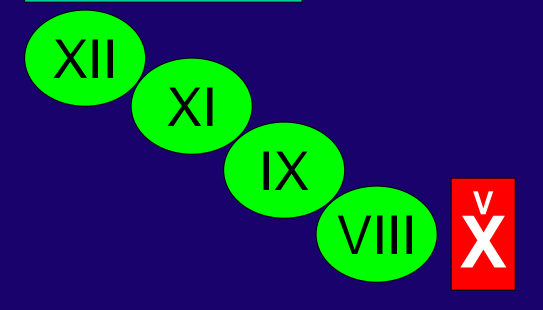


#### Coagulation made easy - the PT

The PT Pathway WEPT 7 Warfarin **Extrinsic Pathway** PT Factor VII Prothrombin (II) Thrombin (IIa) **Fibrinogen** 

#### Coagulation made easy - the aPTT

#### The PTT Pathway



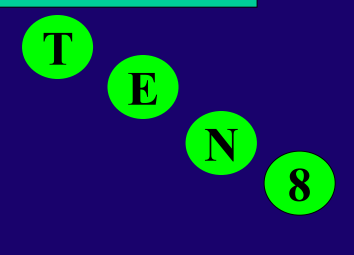
The PTT pathway has all those hideous roman numerals. . . How are we going to remember them? Hmmmmm. . . . . . .

Prothrombin — Thrombin

Fibrinogen Fibrin

#### Coagulation made easy - the aPTT

#### The PTT Pathway



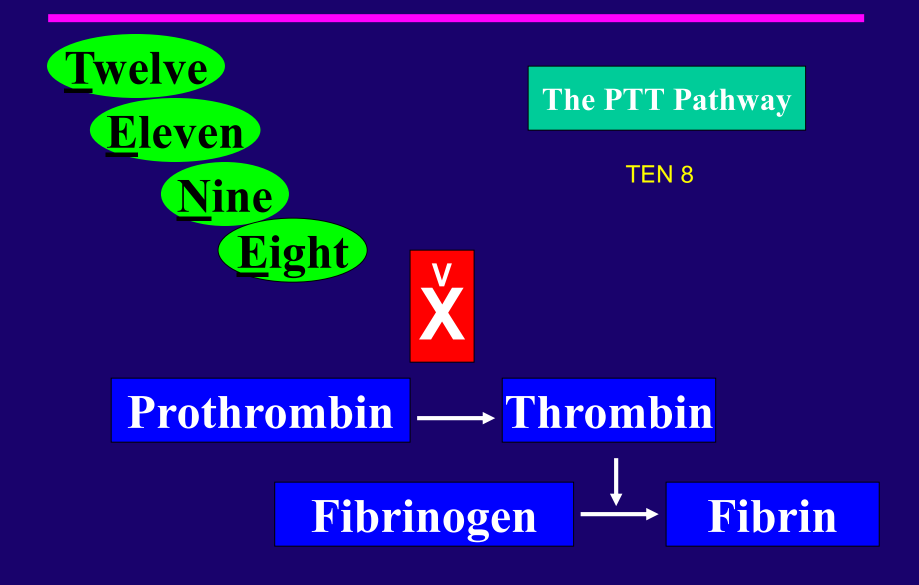


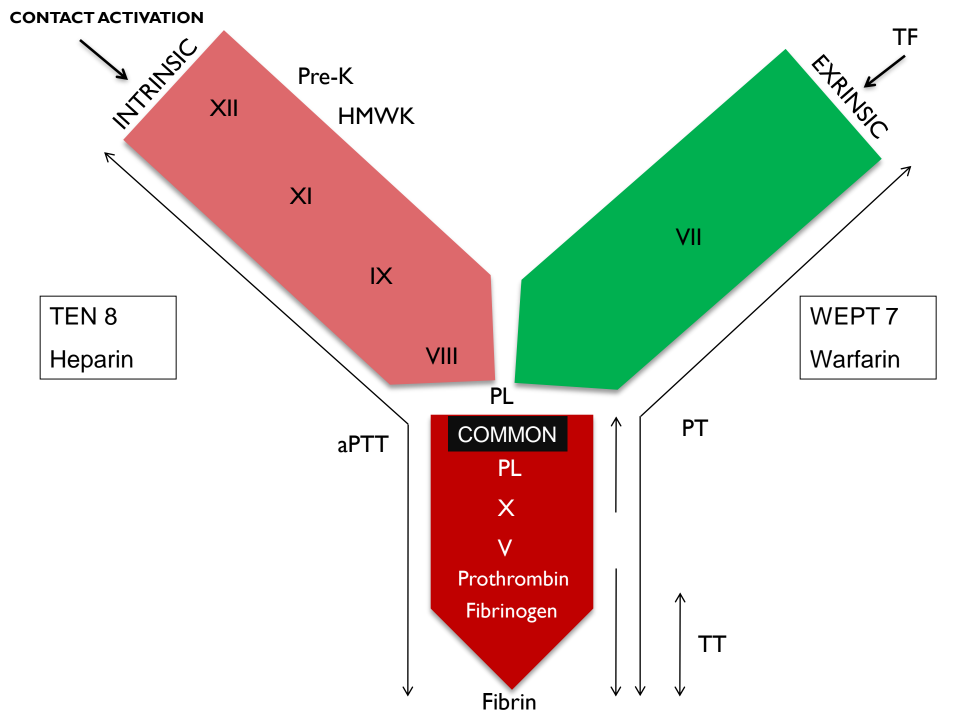


Prothrombin — Thrombin

Fibrinogen Fibrin

#### Coagulation made easy - the aPTT





#### PT

Platelet Poor Plasma Sample (Containing all Factors)

+

PT Reagent

(Tissue Factor, PL, Ca<sup>++</sup>)





Isolated Prolonged PT Prolonged PT in association with other coagulation abnormalities  Vitamin K deficiency Vitamin K antagonists e.g. warfarin, phenindione, rodenticides Liver disease Malabsorption (leading to vitamin K deficiency) High concentrations of unfractionated heparin Direct thrombin inhibitors e.g. Lepirudin, argatroban Afibrinogenaemia and dysfibrinogenemia Dilutional coagulopathy e.g. massive blood transfusion Multiple clotting factor deficiencies e.g. FV and FVIII deficiency Abnormalities of the vitamin K cycle e.g. mutations within the VKORC1 gene	Abnormality	Interpretation	
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	with other coagulation	Vitamin K antagonists e.g. warfarin, phenindione, rodenticides Liver disease Malabsorption (leading to vitamin K deficiency) High concentrations of unfractionated heparin Direct thrombin inhibitors e.g. Lepirudin, argatroban Afibrinogenaemia and dysfibrinogenemia Dilutional coagulopathy e.g. massive blood transfusion Multiple clotting factor deficiencies e.g. FV and FVIII deficiency Abnormalities of the vitamin K cycle e.g.	



#### **aPTT**

Platelet Poor Plasma Sample (Containing all Factors)

+
aPTT Reagent
(PL, Silica)

Incubation
+

Calcium Chloride





Abnormality	Interpretation
Isolated Prolonged APTT	Deficiencies of either XII, XI, IX & VIII.
	Acquired clotting factor inhibitors - these are most commonly directed against FVIII.
	Lupus anticoagulant [LA]
Prolonged APTT + Prolonged PT	Vitamin K deficiency
	Liver disease
	Direct thrombin inhibitors including Hirudin, Argatroban and Dabigatran.
	DIC - due to the consumption of clotting factors
	Massive blood transfusion leading to a dilutional coagulopathy



#### 1

#### All are components of hemostasis except-

- a) Blood Platelets
- b) Red blood cells
- b) Endothelial Cells
- c) Plasma Coagulation Factors





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#### Steps involved in hemostasis include all except-

- a) Vasodilatation
- b. Formation of a platelet plug
- c. Blood Coagulation
- d. Clot retraction and thrombus dissolution





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# Which organ is primarily responsible for the formation of coagulation factors?

- a) Kidney
- b) lungs
- c) liver
- d) Brain





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#### Which test evaluates the extrinsic pathway?

- a). PT
  - b). PTT
  - c). TT
  - d). Closure time
  - e). Bleeding time





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- a). PT WEPT 7
- b). PTT
  - c). TT
  - d). Closure time
  - e). Bleeding time





#### 5

#### Heparin is monitored by the following test-

- a). PT
  - b). PTT
- c). TT
- d). Closure time
- e). Bleeding time





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  - b). PTT
- c). TT
- d). Closure time
- e). Bleeding time





## Which of the following these patients may have a normal PTT?

- a) Thrombocytopenia
- b) Hemophilia A
- c) Hemophilia B
- d) Patient on heparin





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# Prolongation of Bleeding time seen in the following disorders <u>except-</u>

- a) Thrombasthenia
- b) Thrombocytopenia
- c) Von Willebrand ds.
- d) Hemophilia





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## Platelet poor plasma means platelet count less than---

- a) 50,000/cmm
- b) 10,000/cmm
- c) 1,00,000/cmm
- d) 15,000/cmm





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2/Female child – Easy Bruising

•	Test Patient	Reference Range
PT	<b>13</b> s	<b>11-14s</b>
APTT	<b>105</b> s	<b>23-35</b> s
Fibrinogen(Clauss	) 2.7g/L	1.5-4.0g/L
<b>Thrombin Time</b>	<b>13</b> s	<b>10-13s</b>



#### What Next

- Repeat Assay
- Mixing studies: The prolonged APTT corrects in a mix with normal plasma
- What factor assays would you request and why?

FVIII, IX, XI assays.

The FIX assay was normal but the FVIII assay was <5 IU/dl.



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#### **Thank You**

