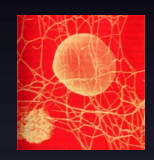


PLATELETS STRUCTURE AND FUNCTIONS COAGULATION MECHANISMS

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Vessel injury



Antithrombogenic

(Favors fluid blood)

Thrombogenic

(Favors clotting)

OBJECTIVES

- *At the end of this lecture you should be able to describe.....
 - *Hemostasis and its steps?
 - *Platelets Structure & Functions
 - *The 2 pathways of coagulation
 - * Factors affecting coagulation
 - *Bleeding & clothing disorders

HEMOSTASIS

From an injured blood vessel is the

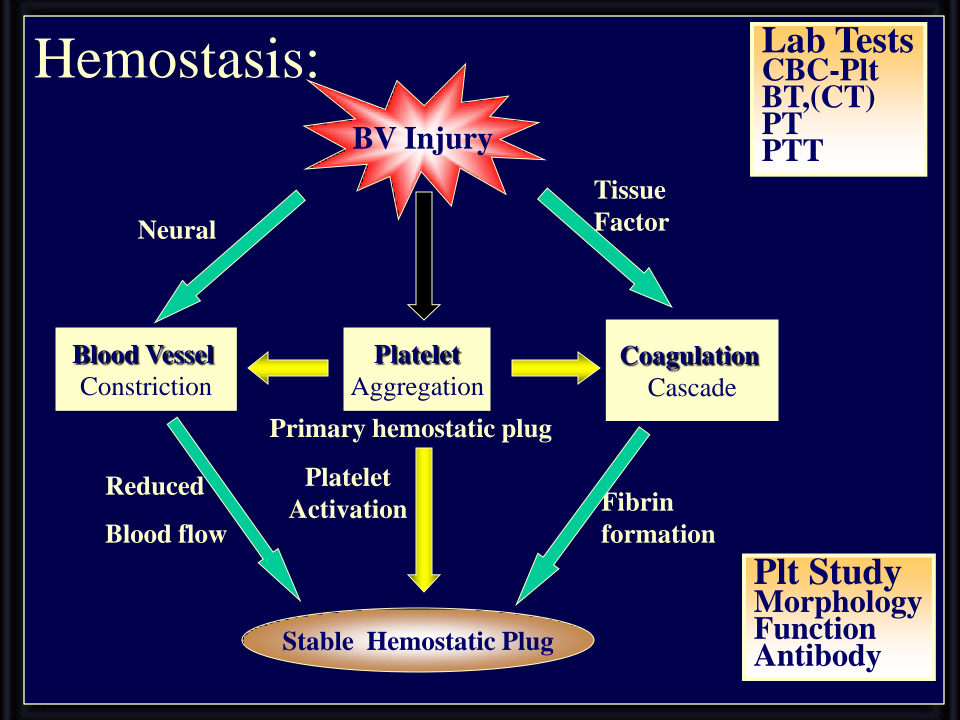
Prevention of blood loss

Or

Stoppage of bleeding

Or

Arrest of bleeding from a broken blood vessel



STEPS OF HEMOSTASIS

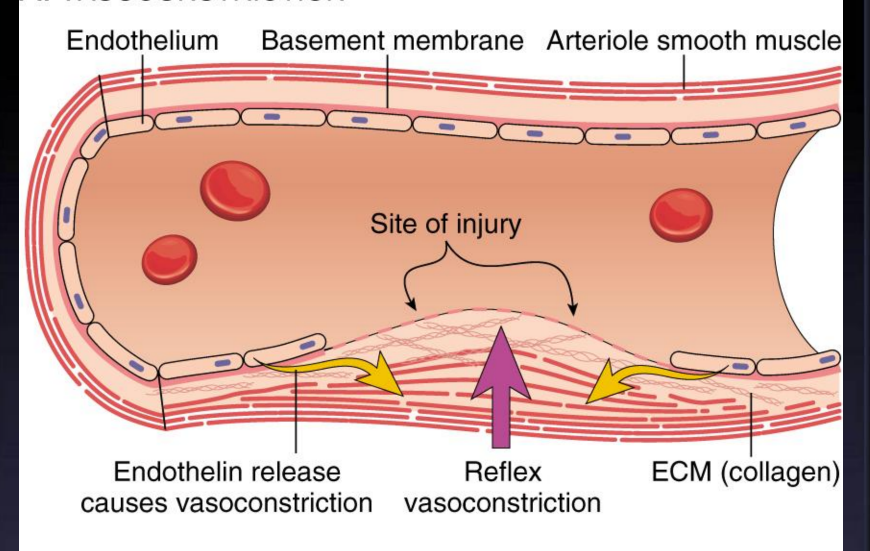
- 1. Vascular Spasm
- 2. Formation of platelet plug
- 3. Blood Coagulation
- 4. Clot Retraction

1-VASCULAR SPASM (Vascular Constriction)

* Factors

- * Nervous reflexes
- Local myogenic spasm
- * Local humoral factor
- For smaller vessels
 - **❖** Platelets → Thrombokanc A_2 (Vasoconstrictor)
- Importance
 - ❖ Crushing injuries → Intense spasm → No lethal loss
 of blood

A. VASOCONSTRICTION

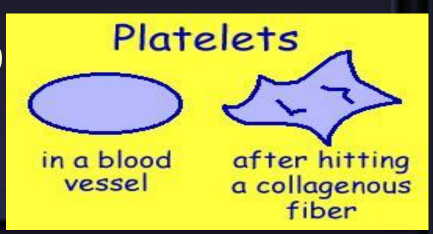


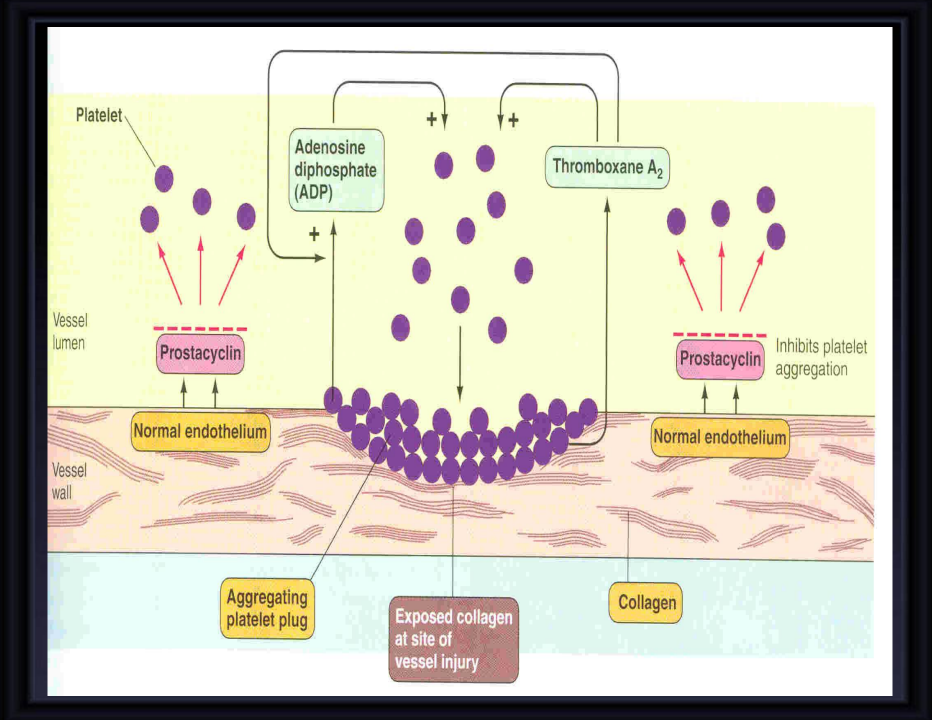
2-FORMATION OF PLATELET PLUG

***** Importance of platelet plug → enough to stop bleeding from small vascular damage

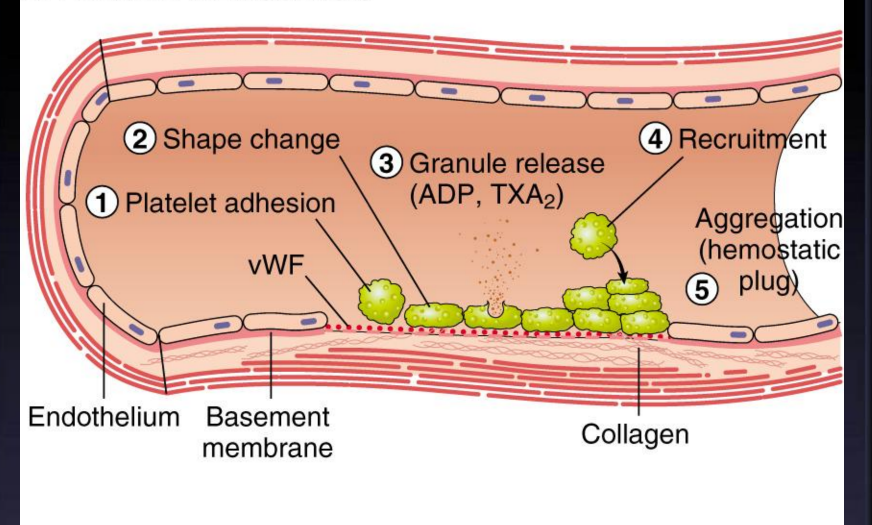
When platelets come in contact with a damaged vascular Surface they;

- Swell & assume irregular forms
- Contarct
- Release granules (ADP,TxA2)
- Become sticky





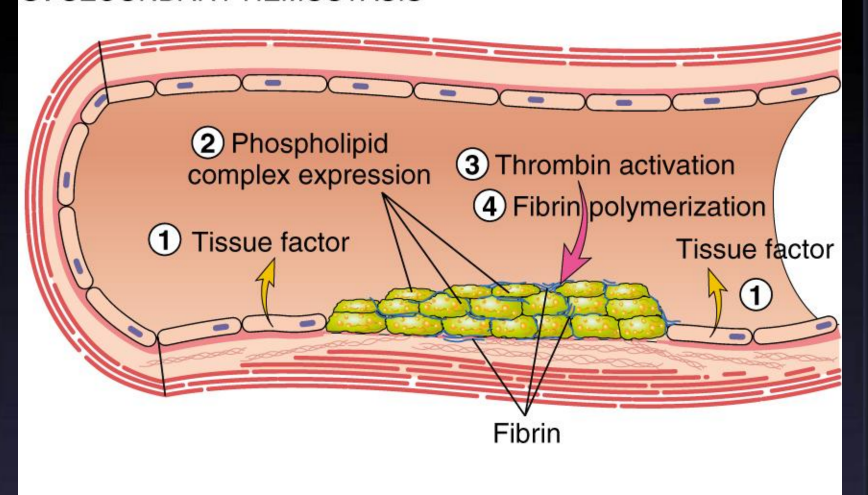
B. PRIMARY HEMOSTASIS



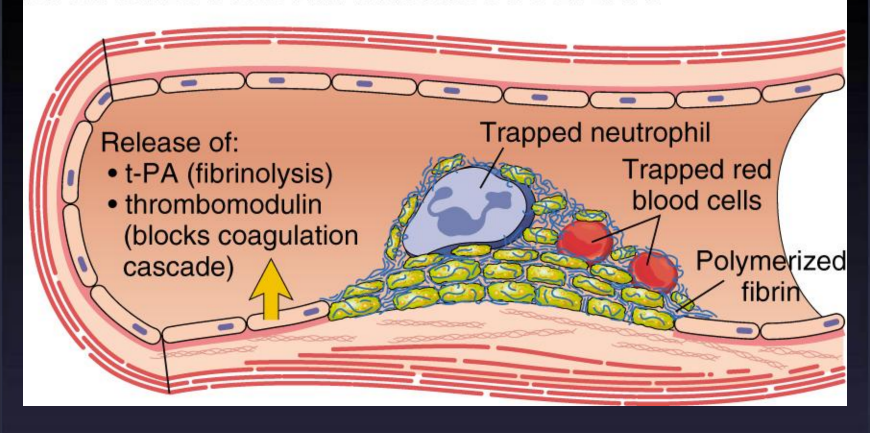
3-BLOOD COAGULATION Formation Of Clot

- * <u>Blood clotting</u> is the transformation of blood from a liquid into a solid gel form
- Pathways
 - * Intrinsic
 - * Extrinsic
- Initiated by: Activator substances from traumatized vascular wall, plts & blood proteins
- * Begins to develop in
 - * 15-20 sec \rightarrow Minor trauma
 - \star 1-2 min. \rightarrow Severe trauma

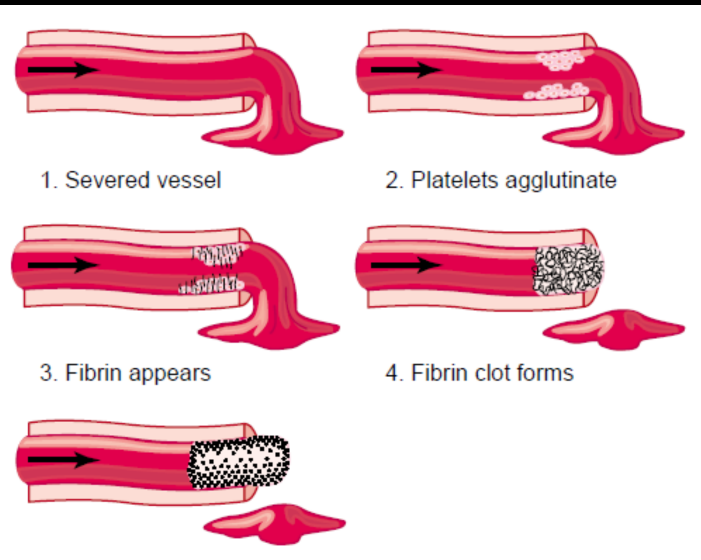
C. SECONDARY HEMOSTASIS



D. THROMBUS AND ANTITHROMBOTIC EVENTS



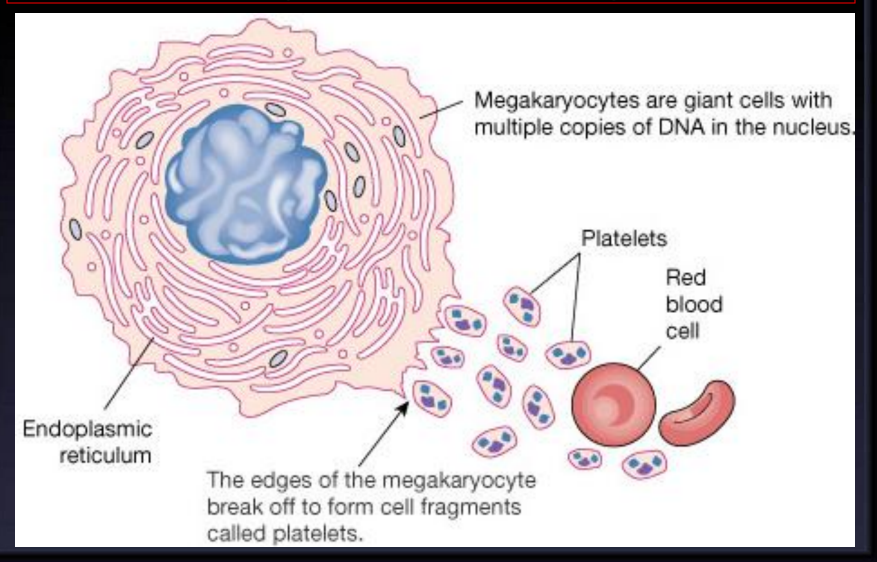
Physical Events Of Clotting Process



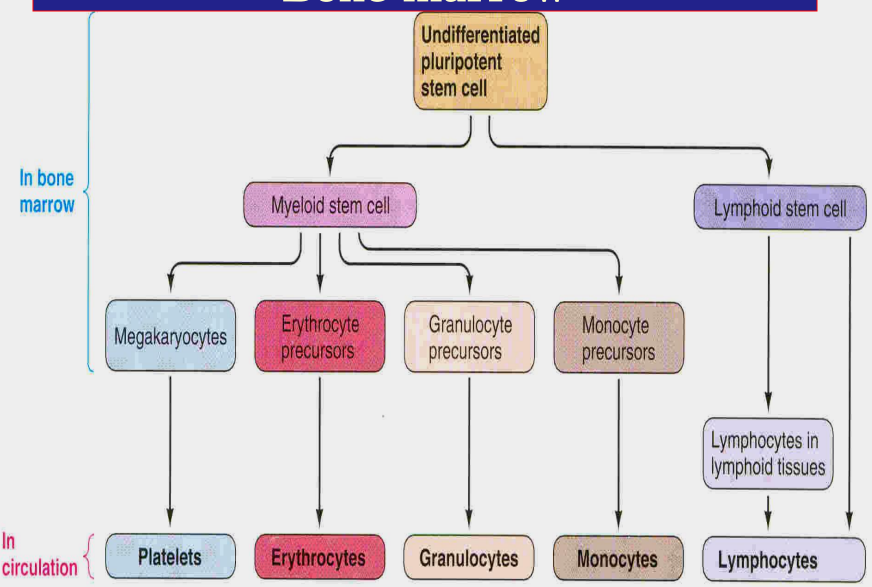
5. Clot retraction occurs

PLATELETS

Formed by fragmentation from megakaryoctyes



SITE OF FORMATION Bone-marrow



PLATELETS (Characteristics)

SHAPE: MINUTE ROUND OR OVAL DISCS

SIZE: 1-4 um IN DIAMETER

HALF LIFE: 8-12 DAYS

COUNT: 150,000 – 300,000/ microlitrer

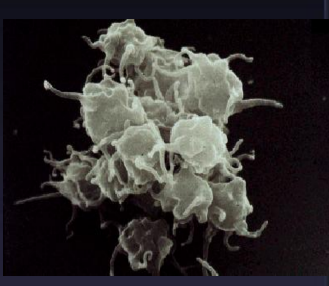
LOCATION: 80% in blood & 20% in spleen

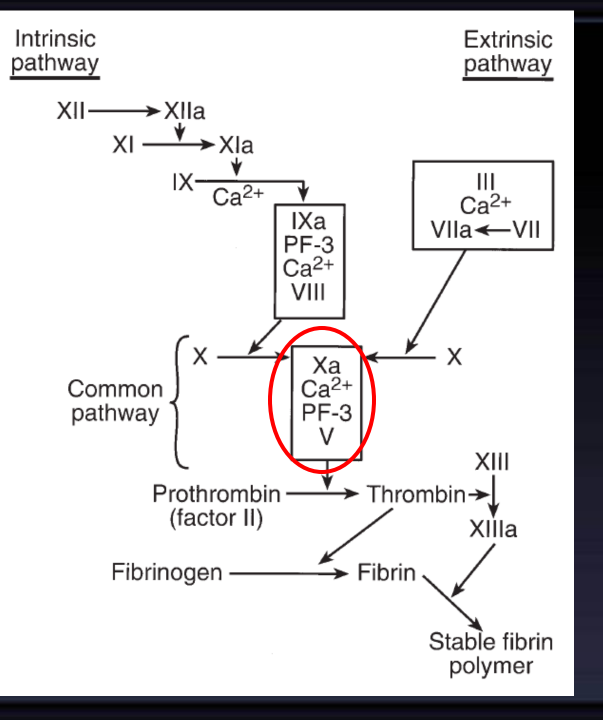
- Contractile, adhesive, cell fragments.
- * Store coagulation factors & enzymes
- * Surface Binding sites for fibrinogen
- * Surface Glycoprotein Antigens-HPA1.

FUNCTIONAL CHARACTERISTICS

- ACTIN AND MYOSIN MOLECULES
- THROMBESTHENIN
- ENDOPLASMIC RETICULUM AND GOLGI APPARATUS
- MITOCHONDRIA
- ENZYME SYSTEMS FOR SYNTHESIS OF PROSTAGLANDINS
- FIBRIN STABILIZING FACTOR
- GROWTH FACTOR







MECHANISM OF CLOTTING

- 1. Formation of Prothrombin activator complex
- 2. Conversion of prothrombin into thrombin
- 3. Conversion of fibrinogen into fibrin

COAGULATION CASCADE

Formation Of Prothrombin Activator Complex is by:

*2 Ways

- ***By Extrinsic pathway** → trauma to vascular wall and surrounding tissues
- ♦ By Intrinsic pathway \rightarrow trauma to the blood \rightarrow Is the rate limiting factor

CONVERISON OF PRTHROMBIN TO THROMBIN

By Prothrombin Activator Complex

* Prothrombin

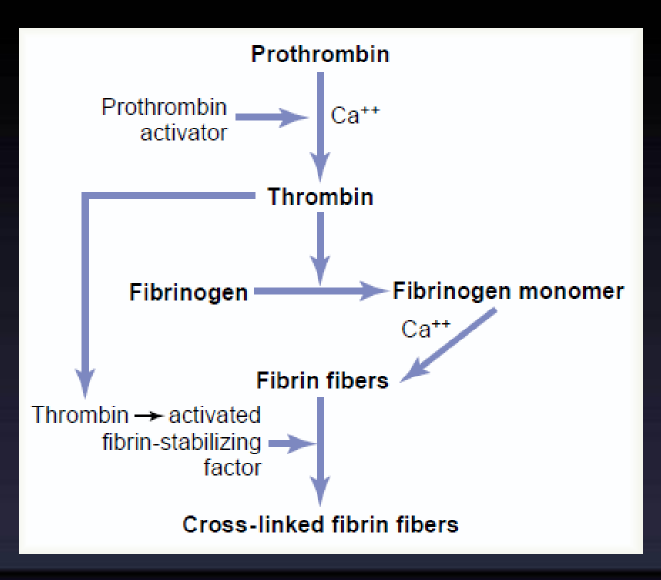
- ❖ Plasma protein (Alpha₂ globulin)
- **♦ Mol. Wt. 68,700**
- Plasma conc. 15 mg/dl
- Unstable protein
- Synthesized by liver
- ***** Vitamin-K is required for synthesis

CONVERSION OF FIBRINGEN TO FIBRIN Formation Of Clot

* Fibrinogen

- Mol. Wt. 340,000
- **❖ Plasma conc.** − 100 − 700 mg/dl
- Synthesized in liver

ACTION OF THROMBIN ON FIBRONOGEN TO FORM FIBRIN



BLOOD CLOT

A meshwork of fibrin fibres running in all directions entrapping blood cells, platelets and plasma

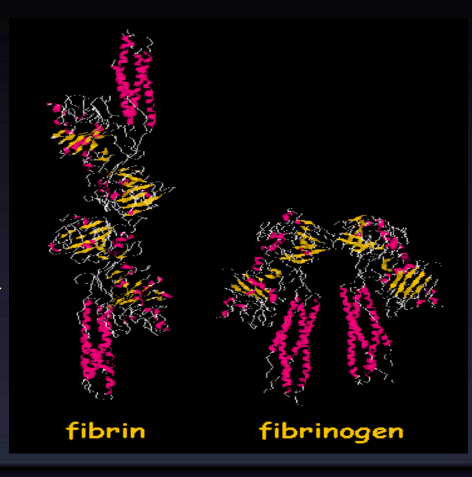


Table 36-1

Clotting Factors Guyton

Clotting Factors in Blood and Their Synonyms

Clotting Factor	Synonyms
Fibrinogen Prothrombin	Factor I Factor II
Tissue factor	
Calcium	Factor III; tissue thromboplastin
Factor V	Proaccelerin; labile factor; Ac-globulin (Ac-G)
Factor VII	Serum prothrombin conversion accelerator (SPCA); proconvertin; stable factor
Factor VIII	Antihemophilic factor (AHF); antihemophilic globulin (AHG); antihemophilic factor A
Factor IX	Plasma thromboplastin component (PTC); Christmas factor; antihemophilic factor B
Factor X	Stuart factor; Stuart-Prower factor
Factor XI	Plasma thromboplastin antecedent (PTA); antihemophilic factor C
Factor XII	Hageman factor
Factor XIII	Fibrin-stabilizing factor
Prekallikrein	Fletcher factor
High-molecular-weight	Fitzgerald factor; HMWK
kininogen	(high-molecular-weight) kininogen
Platelets	

Clotting Factors Ganong

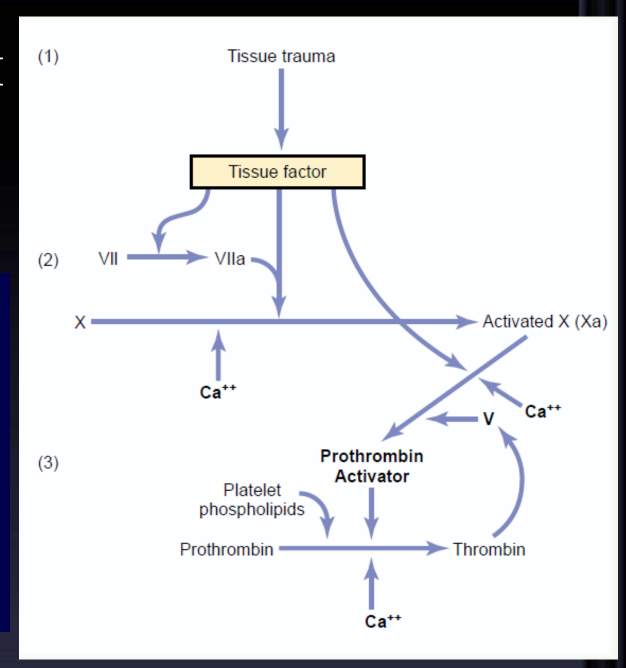
TABLE 31–5 System for naming blood-clotting factors.

Factora	Names
1	Fibrinogen
II	Prothrombin
III	Thromboplastin
IV	Calcium
V	Proaccelerin, labile factor, accelerator globulin
VII	Proconvertin, SPCA, stable factor
VIII	Antihemophilic factor (AHF), antihemophilic factor A, antihemophilic globulin (AHG)
IX	Plasma thromboplastic component (PTC), Christmas factor, antihemophilic factor B
X	Stuart–Prower factor
XI	Plasma thromboplastin antecedent (PTA), antihemophilic factor C
XII	Hageman factor, glass factor
XIII	Fibrin-stabilizing factor, Laki–Lorand factor
HMW-K	High-molecular-weight kininogen, Fitzgerald factor
Pre-Ka	Prekallikrein, Fletcher factor
Ka	Kallikrein
PL	Platelet phospholipid

^aFactor VI is not a separate entity and has been dropped.

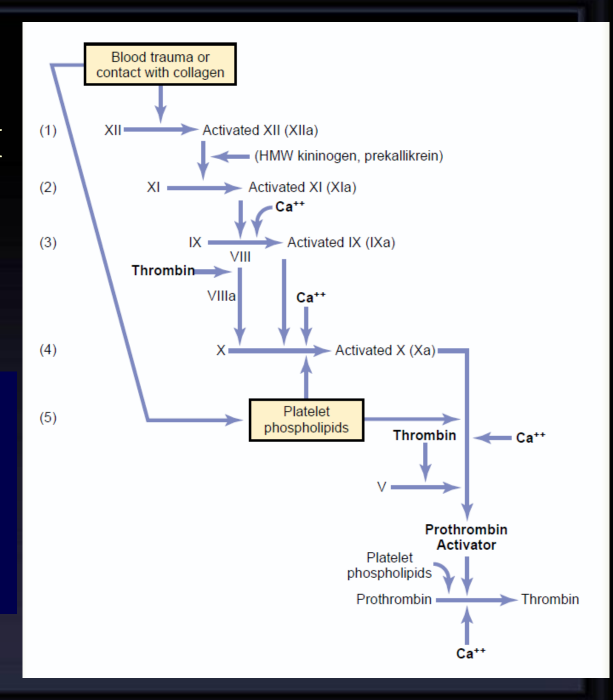
EXTRINSIC MECHNANISM FOR INITIATING CLOTTING

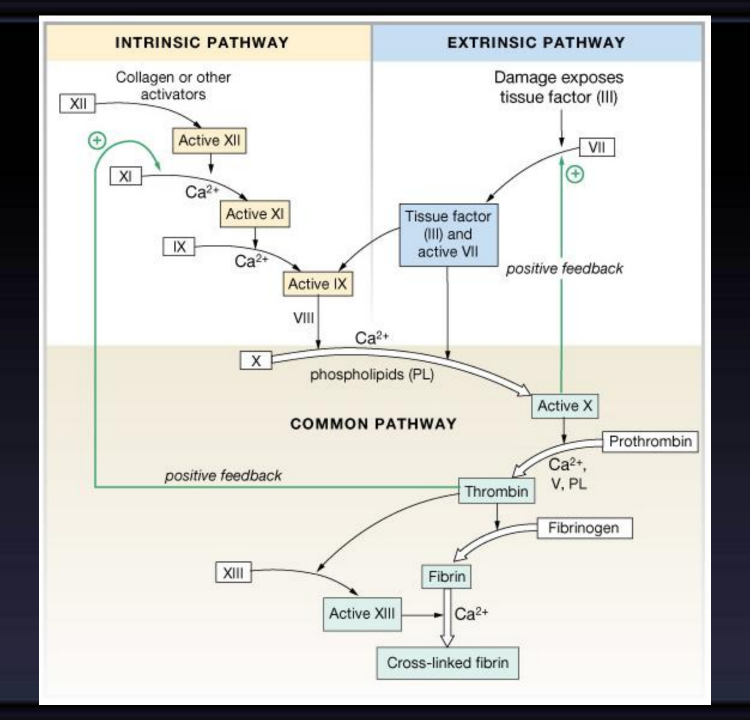
TFor tissue thromboplastin; includes phospholipids from the membranes of the tissue plus a lipoprotein complex that functions mainly as a proteolytic enzyme.



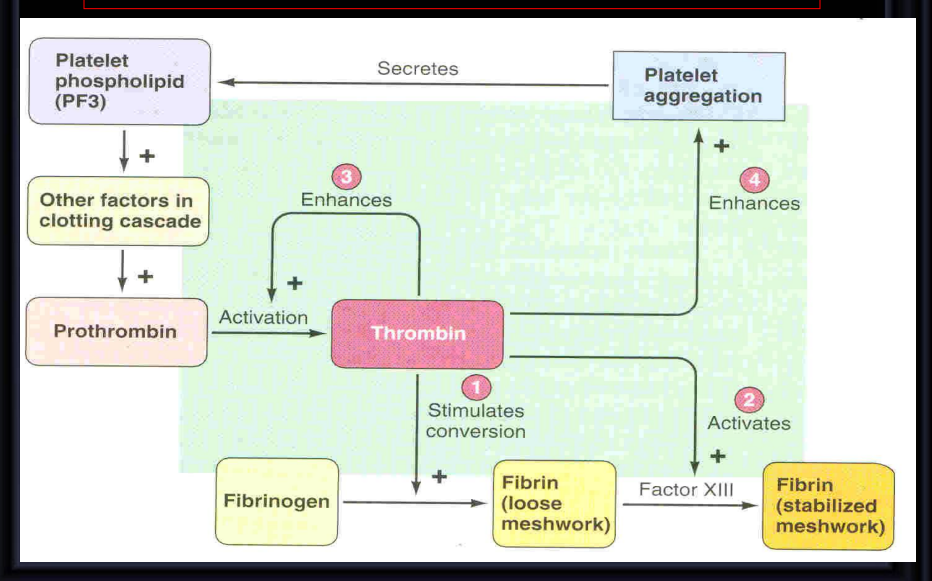
INTRINSIC MECHNANISM FOR INITIATING CLOTTING

Trauma to the blood itself or exposure of the blood to collagen from a traumatized blood vessel wall





ROLE OF THROMBIN IN HEMOSTASIS



CLOT RETRACTION

- * When clot contracts, it expresses most of the fluid from the clot within 20-60 min called → Serum
- * SERUM CANNOT CLOT
- * ROLE OF PLTS IN CLOT FORMATION & RETRACTION
- * VICIOUS CIRCLE OF CLOT FORMATION

ROLE OF CALCIUM IONS IN CLOTTING

 \bullet No Ca⁺⁺ \rightarrow No Clotting

Blood samples are prevented from clotting by:

- **♦ Citrate ions** → **Deionization of Ca**⁺⁺
- **♦ Oxalate ions** → precipitate the Ca⁺⁺
- ♦ Heparin → combines with antithrombin effectiveness increases by 100-1000 fold, Also remove Factors XII, XI, X, and IX
- *** Warfarin: ↓** production of Factors VII, IX and X by liver.
- **❖ EDTA** → strongly & irreversibly chelates Ca⁺⁺

LYSIS OF BLOOD CLOTS PLASMIN

Plasminogen / Profibrinolysin

T-PA (Tissue Plasminogen Activator)

Plasmin or Fibrinolysin

Lysis of clot

NATURAL INTRAVASCULAR ANTICOAGULANTS

1. Endothelial Surface Factors

- * Smoothness of Endothelium
- Glycocalyx Layers
- * Thrombomodulin Protein binds to thrombin → Activates
 Protein C (with ProtS) → inactivates factors V & VIII
 and inactivates an inhibitor of tPA → increasing the
 formation of plasmin.
- 2. Antithrombin action of Fibrin and Antithrombin III
 - * 85-90 % Thrombin binds with Fibrin
 - * 10-15 % Thrombin binds with Antithrombin III

Antithrombin III is a circulating protease blocking clot factors

NATURAL INTRAVASCULAR ANTICOAGULANTS

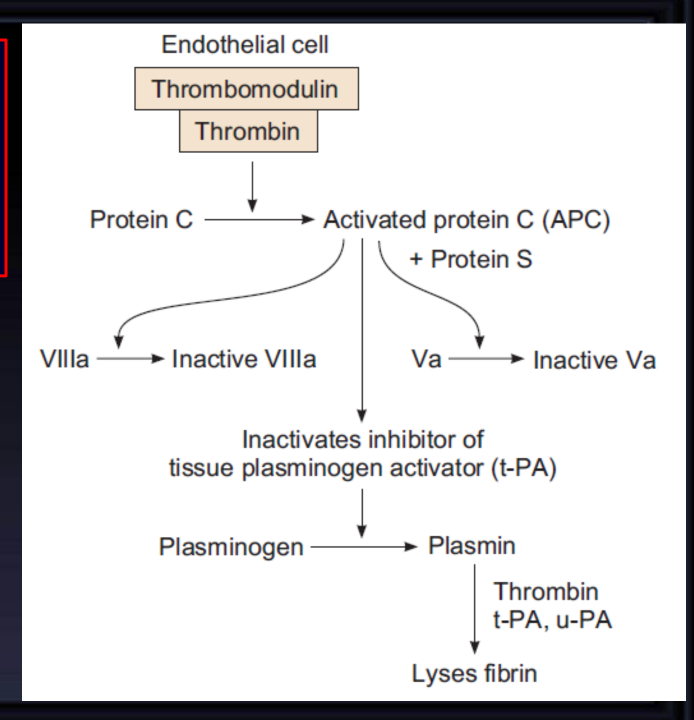
3. Heparin

- vely charged conjugated polysaccharide
 - * Increase the effectiveness of Antithrombin III
 - Produced by
 - Mast cells
 - * Basophil cells
- Most widely used anticoagulant clinically e.g. in stroke
- 4. Alpha₂ Macrogobulin
 - Acts as a binding agent for several coagulation factors

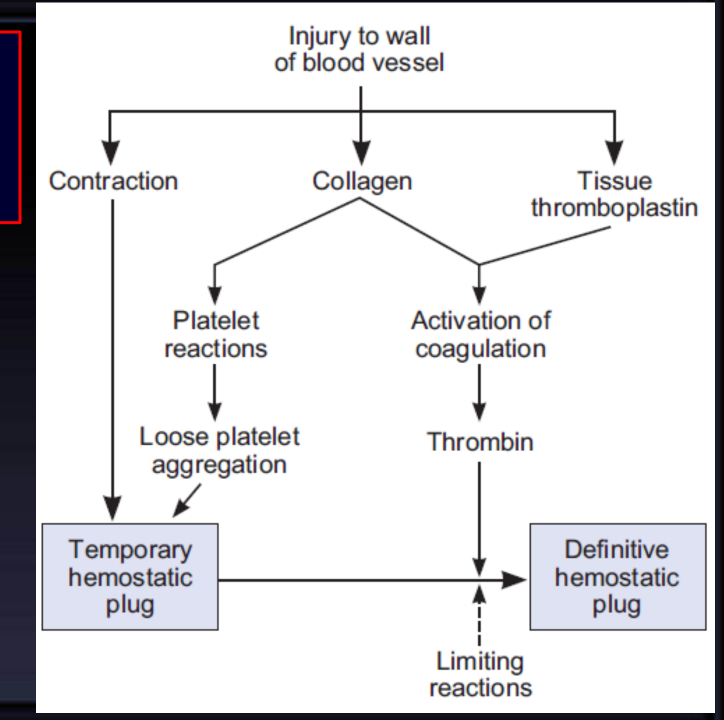
BLEEDING & CLOTTING DISORDERS

- A. Liver diseases & Vitamin-K deficiency
- B. Hemophilia
- c. Thrombocytopenia

The fibrinolytic system and its regulation by Protein C



Summary of reactions involved in hemostasis.



BLEEDING DISORDERS

- A. Liver diseases & Vitamin-K deficiency
- e.g. Hepatitis, Cirrhosis
 - Decreased formation of clotting factors
 - Icnreased clotting time
- Vitamin K dependent factors
 - Prothrombin, Factor VII, IX, X

BLEEDING DISORDERS

- A. Vitamin-K
- Fat soluble vitamin
- Required by liver for formation 4 clotting factors
- Sources
 - Diet
 - Sythesized in the intestinal tract by bacteria
- Deficiency
 - Malabsorption syndromes
 - Biliary obstruction
 - ***** Broad spectrum antibiotics
 - Dietary def (in Neonates)
 - **Rx.:** Treat the underlying cause Vit K injections

THROMBOCYTOPENIA

- * Platelet count upto 50,000 ul
- *** Less than 10,000 ----- Fatal**
- * ETIOLOGY
- * Decreased production
 - Aplastic anemia
 - * Leukemia
 - * Drugs
 - * Infections (HIV, Measles)

THROMBOCYTOPENIA (cont.)

* Increased destruction

- * ITP
- * Drugs
- * Infections

Clinical Features

- Easy brusability
- ***** Epistaxis
- Gum bleeding
- * Hemorrhage after minor trauma
- Petechiae/Ecchymosis

THROMBOCYTOPENIA (cont.)

- Diagnosis
 - *PLT decreased
 - *B.T increased
- * Rx
 - * Rx of the underlying cause
 - *** PLT concentrates**
 - Fresh whole blood transfusion
 - * Splenectomy



CLOTTING DISORDER HEMOPHILIA

* HEMOPHILIA A

- Classic Hemophilia
- *85 % cases, X Linked recessive trait, females are carriers
- Deficiency of smaller part of factor VIII

* HEMOPHILIA B

- Christmas disease
- *15 % cases
- Deficiency of factor IX

HEMOPHILIA

***** Clinical Features

- * Easy bruising, massive bleeding after trauma or operation, hemorrhages in joints
- Factor VIII
 - **♦ Small Comp.** → Hemophilia A
 - ***** Large Comp. → Von-Willebrand's disease
- * Rx
 - *Injection of factor VIII (Hemophilia A)
 - **❖ Injection of factor IX (Hemophilia B)**

FACTORS AFFECTING BLOOD PLATELET COUNT

- **♦** AGE: ↓ in Newborn
- * MENSTRUAL CYCLE:
 - **♦** ↓ Prior to menstruation
 - **♦** ↑ After menstruation
- ❖ PREGNANCY: ↓
- ❖ INJURY: ↑
- * ADRENALINE: ↑
- ♦ HYPOXIA: ↑
- ♦ SMOKING: ↓
- ♦ NUTRITIONAL DEFICIENCIES: ↓ eg; vitamin B12, folic acid and iron

LAB TESTS FOR HEMOSTASIS

- Blood Counts
- Platelets aggregation
- * PT (eg; Warfarin monitoring)
- * APTT (eg; Heparin monitoring)



- * CT
- Fibrinogen
- * Protein C
- * T-PA
- * PAI-1
- * TFPI-T & TFPI-F

