

## HEMODYNAMIC DISORDERS, THROMBOEMBOLISM, AND SHOCK 2

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• Normal hemostasis comprises a series of <u>regulated</u> processes that <u>culminate</u> in the <u>formation</u> of a <u>blood clot</u> that <u>limits bleeding</u> from an <u>injured vessel</u>

• The pathologic counterpart of hemostasis is thrombosis, the formation of blood clot (thrombus) within non-traumatized context versel.

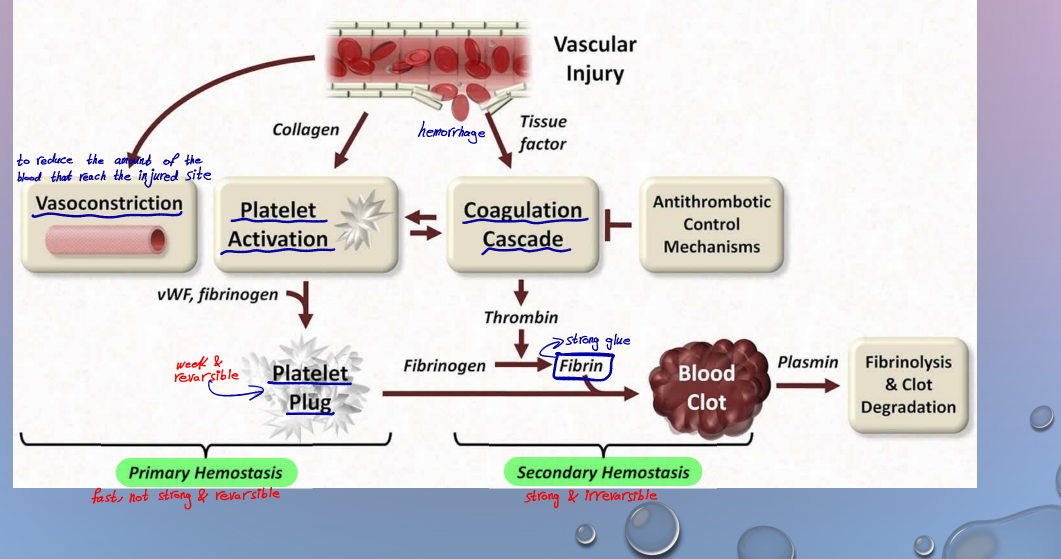


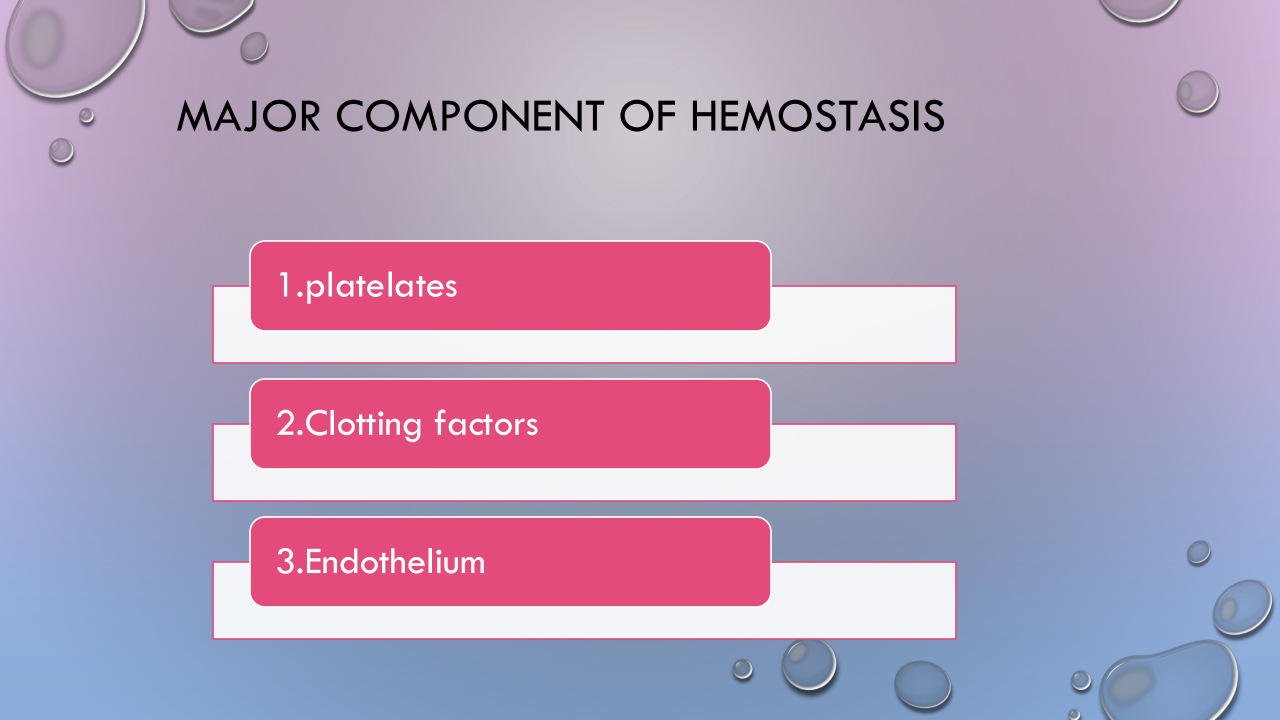


 Hemostasis is process involving <u>platelets</u>, <u>clotting factors</u>, and <u>endothelium</u> that occurs at the site of vascular injury and culminates in the formation of a blood clot, which serves to prevent or limit the extent of bleeding.

### **Major Components of Hemostasis**

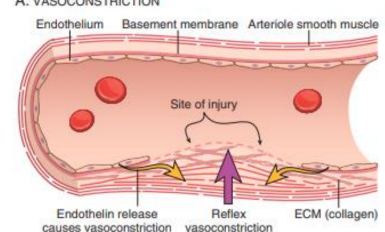
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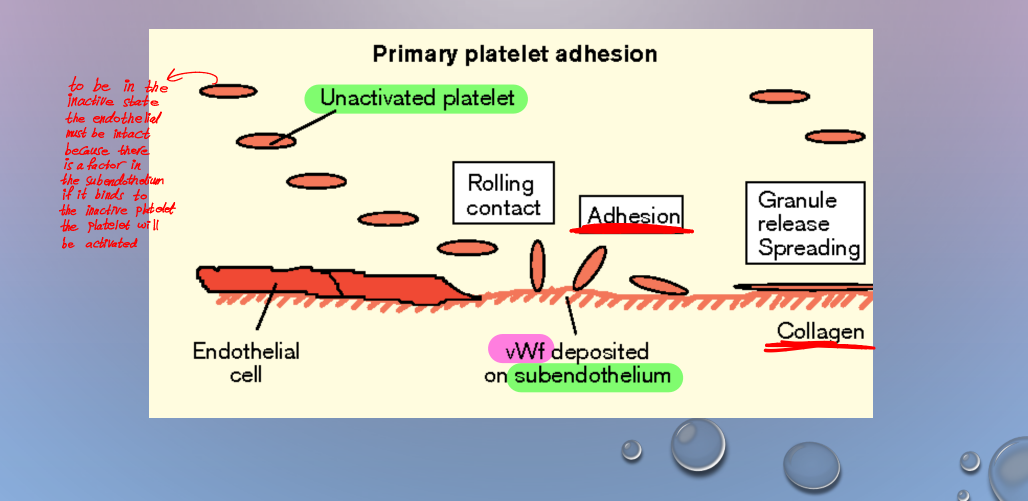
THE GENERAL SEQUENCE OF EVENTS LEADING TO HEMOSTASIS AT A SITE OF VASCULAR INJURY INCLUDE: I. PRIMARY HEMOSTASIS

- <u>1. ARTERIOLAR VASOCONSTRICTION</u>:
- occurs immediately and markedly reduces blood flow to the injured area.
- it is mediated by reflex neurogenic mechanisms.
- it is augmented by endothelin, a potent endothelium-derived vasoconstrictor.
- this <u>effect is transient</u>, however, <u>bleeding would resume</u> if <u>not followed by</u> activation of <u>platelets</u> and <u>coagulation</u> factors.

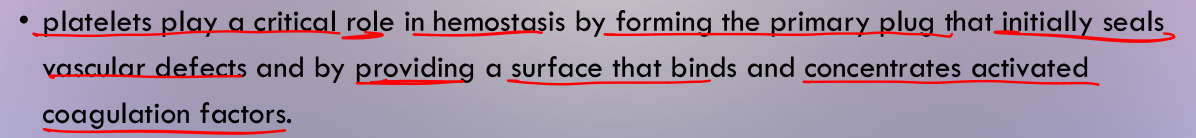




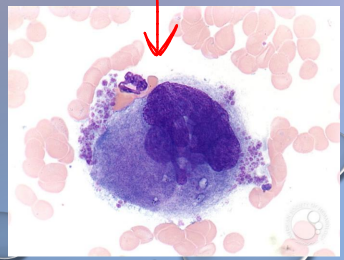
• THE FORMATION OF THE PLATELET PLUG.







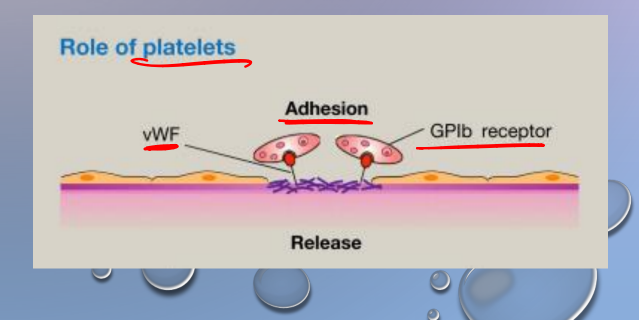
 platelets are disc-shaped anucleate cell fragments that are shed from megakaryocytes in the bone marrow into the bloodstream.



### PLATELETS UNDERGO A SEQUENCE OF REACTIONS AFTER A TRAUMATIC VASCULAR INJURY THAT CULMINATE IN THE FORMATION OF A PLATELET PLUG

### • <u>1. PLATELET ADHESION</u>:

• is mediated via interactions with vwf, which acts as a bridge between the platelet surface receptor glycoprotein ib (gpib) and exposed collagen.





- <u>A. CHANGES IN SHAPE</u> from smooth discs to "spiky "with greatly increased surface area.
- alterations in glycoprotein iib/iiia that increase its affinity for fibrinogen
- the <u>translocation</u> of <u>negatively charged</u> phospholipids to the platelet surface *I need this surface to be negatively Evel charged for the Precipitation of Gagulation system*
- <u>B. SECRETION OF GRANULE CONTENTS, e.g.</u>
- ✓ THROMBIN: activates platelets
- ✓ <u>ADP</u>: create an <u>additional</u> rounds of platelet activation.

✓ THROMBOXANE A2 (TXA2): a potent inducer of platelet aggregation.

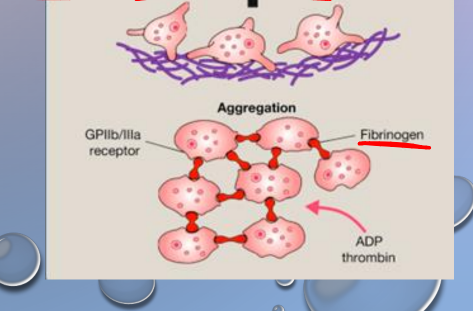
3. PLATELET AGGREGATION FOLLOWS THEIR ACTIVATION.

• The conformational change in glycoprotein iib/iiia allows binding of fibrinogen that forms bridges between adjacent platelets, leading to their aggregation.

ادابا مما من المراع من المبترة fibrinogen cause <u>reversible aggregation</u> & Unstable

✓ thrombin cause irreversible aggregation (converts fibrinogen into insoluble fibrin).

cytoskeleton cause contraction of the plug.
 to make this network stronger

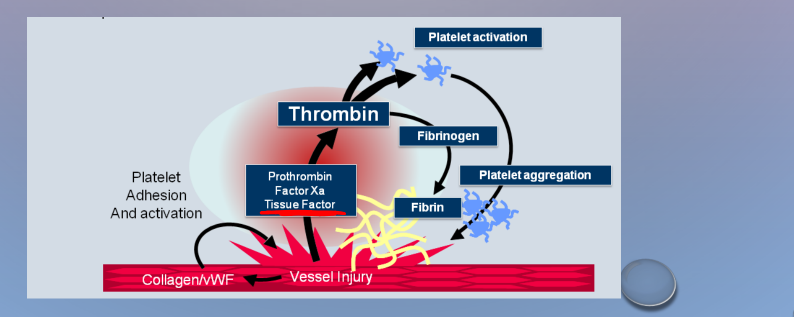




- DEPOSITION OF FIBRIN.
  - VASCULAR INJURY EXPOSES TISSUE FACTOR AT THE SITE OF INJURY.

from subendothelial

• TISSUE FACTOR BINDS AND ACTIVATES FACTOR VIL, SETTING IN MOTION A CASCADE OF REACTIONS THAT CULIMINATES IN THROMBIN GENERATION.

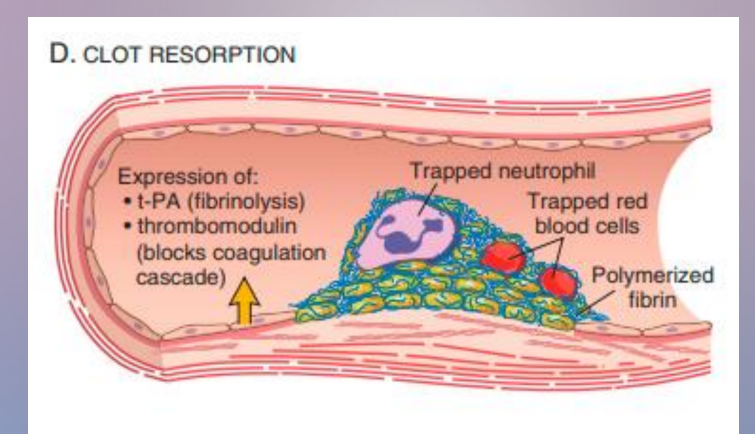


I. CLOT STABILIZATION AND RESORPTION:

• polymerized fibrin and platelet aggregates undergo <u>contraction</u> to form a <u>solid</u>, permanent plug that prevents further hemorrhage.

• entrapped red cells and leukocytes are also found in hemostatic plugs, in part due to adherence of leukocytes to p-selectin expressed on activated platelet

 at this stage, counterregulatory mechanisms (e.g., tissue plasminogen activator, t-pa made by endothelial cells) are set into motion that limit clotting to the site of injury, and eventually lead to clot resorption and tissue repair.



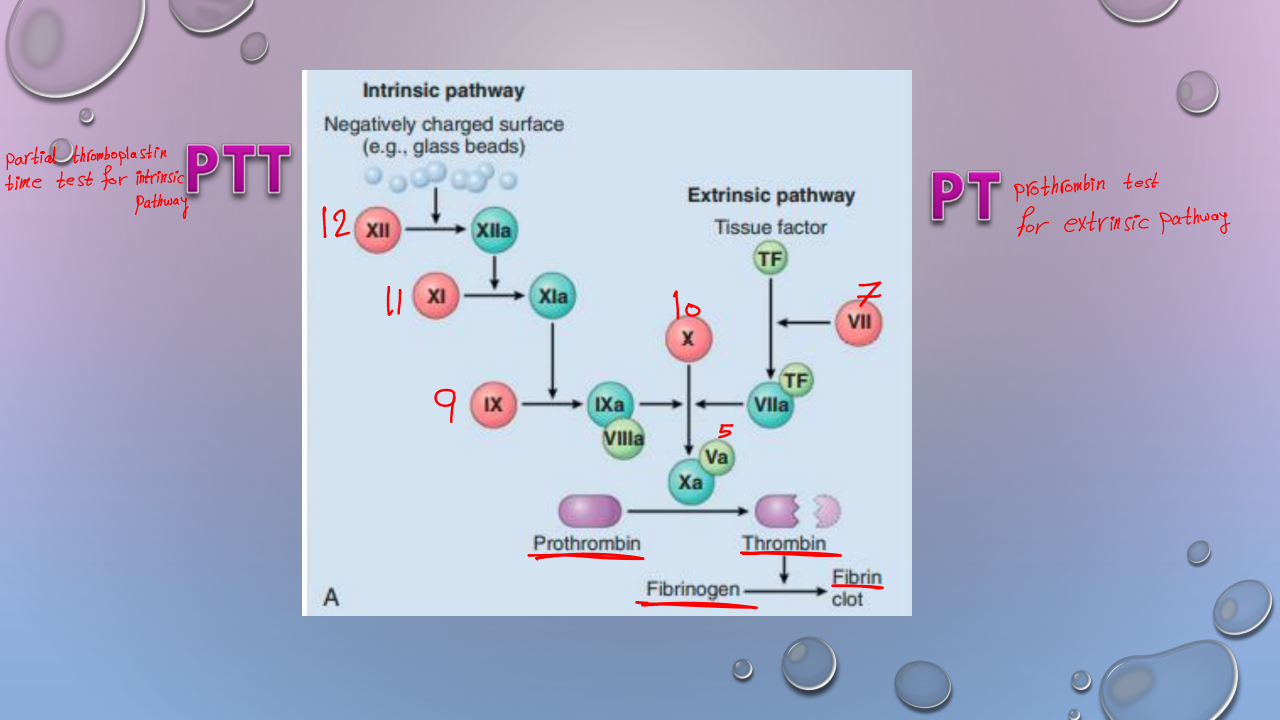


## COAGULATION CASCADE

 the coagulation cascade is a series of amplifying enzymatic reactions that lead to the deposition of <u>an insoluble</u> fibrin clot.

 each reaction step involves an <u>enzyme</u> (an activated coagulation factor), a <u>substrate</u> (an <u>inactive proenzyme form of a coagulation factor</u>), and a <u>cofactor (a</u> reaction accelerator).

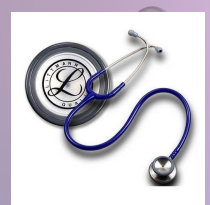
 these components are assembled on a negatively charged phospholipid surface, which is provided by activated platelets. assembly of reaction complexes also depends on calcium





### vitamin K antagonists

Mnemonic for Vitamin K Dependent Clotting Factors "Two plus seven is nine NOT ten!" 2 7 9 10 1972 Coagulation cascade has traditionally been divided into the extrinsic and intrinsic pathways

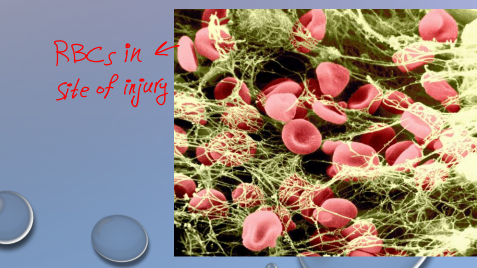


- <u>1.THE PROTHROMBIN TIME (PT)</u>
- ASSAY ASSESSES THE FUNCTION OF THE PROTEINS IN THE <u>EXTRINSIC PATHWAY (FACTORS VII,</u> X, V, II (PROTHROMBIN), AND FIBRINOGEN).
- THE PARTIAL THROMBOPLASTIN TIME (PTT)
- ASSAY SCREENS THE FUNCTION OF THE PROTEINS IN THE INTRINSIC PATHWAY (FACTORS XII, XI, IX, VIII, X, V, II, AND FIBRINOGEN.

# MONG PHROMBIN'S MOST IMPORTANT ACTIVITIES ARE THE FOLLOWING:

- O CONVERSION OF FIBRINOGEN INTO CROSSLINKED FIBRIN.
- THROMBIN DIRECTLY CONVERTS SOLUBLE FIBRINOGEN INTO FIBRIN MONOMERS THAT
   POLYMERIZE INTO AN INSOLUBLE FIBRIL.
- <u>2.PLATELET ACTIVATION.</u>
- <u>4. ANTI-COAGULANT EFFECTS.</u>
- ENCOUNTERING NORMAL ENDOTHELIUM, THROMBIN CHANGES FROM A PROCOAGULANT TO AN ANTICOAGULANT.

thrombin \_ proGagulant -> site of injury > anti Gagulant -> intact endothelial

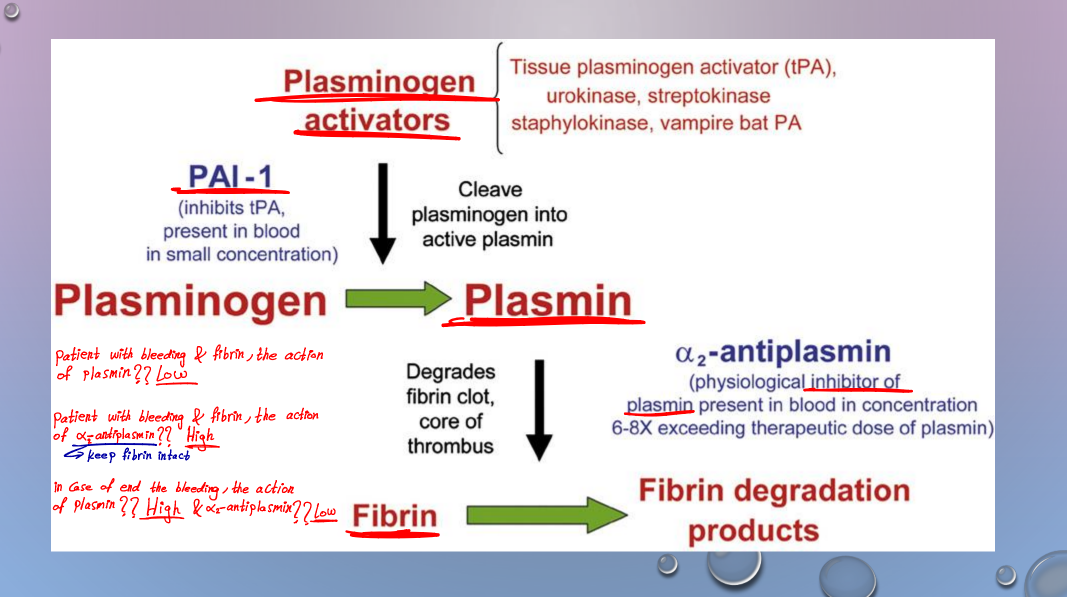


## FACTORS THAT LIMIT COAGULATION.

 coagulation must be restricted to the site of vascular injury to prevent dangerous consequences through:

- <u>1.simple dilution</u>:
- blood flowing at the site of injury washes out activated coagulation factors, which are rapidly removed by the liver.
- 2.requirement for negatively charged phospholipids
- <u>3. fibrinolytic cascade</u>:
- through the enzymatic activity of plasmin, which breaks down fibrin and interferes with its polymerization.

### fibrinolytic cascade



- - AN ELEVATED LEVEL OF BREAKDOWN PRODUCTS OF FIBRINOGEN (D-DIMERS) ARE A USEFUL
    CLINICAL MARKERS OF SEVERAL THROMBOTIC STATES

# **D-Dimer test**

Venous thromboembolism (VTE)

Patient & it's right leg swelling hotness & tenderness (unilateral)



• The balance between the anticoagulant and procoagulant activities of endothelium often determines whether clot formation, propagation, or dissolution occurs.

#### .PLATELET INHIBITORY EFFECTS:

serve as a barrier that shields platelets from subendothelial vwf and collagen.

✓ releases a number of factors that inhibit platelet activation and aggregation. among the most important are prostacyclin (pgi2), nitric oxide (no).

- endothelial cells bind and alter the activity of thrombin, which is one of the most potent activators of platelets.
- <u>2. ANTICOAGULANT EFFECTS.</u>
- normal endothelium shields coagulation factors from tissue factor in vessel walls and expresses multiple factors that actively oppose coagulation:
- MOST NOTABLY thrombomodulin, endothelial protein c receptor, heparin-like molecules, and tissue factor pathway inhibitor.

