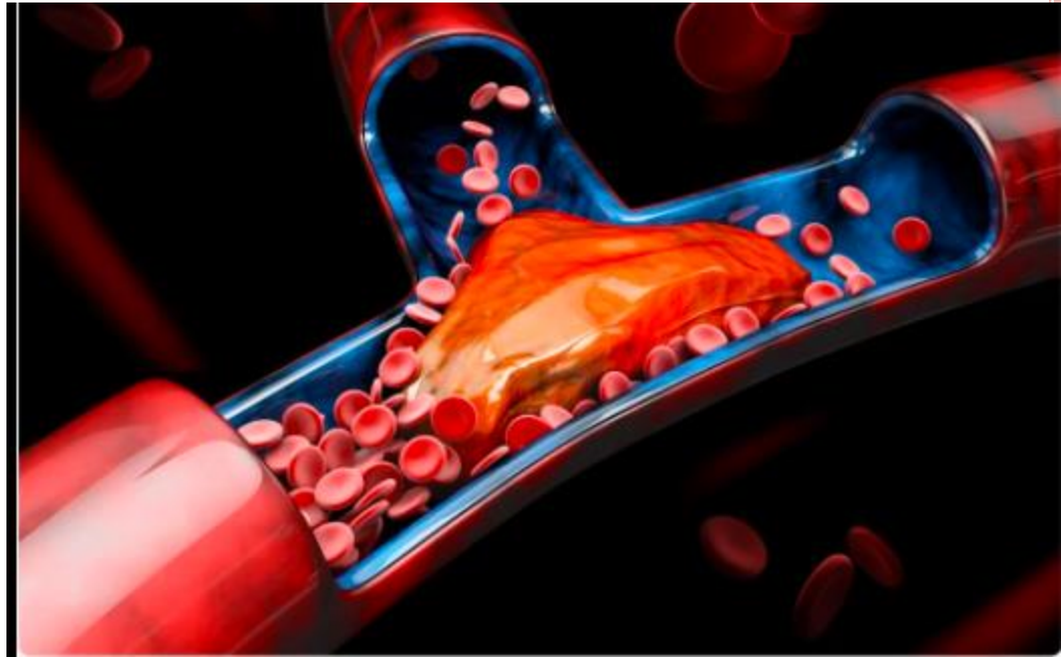


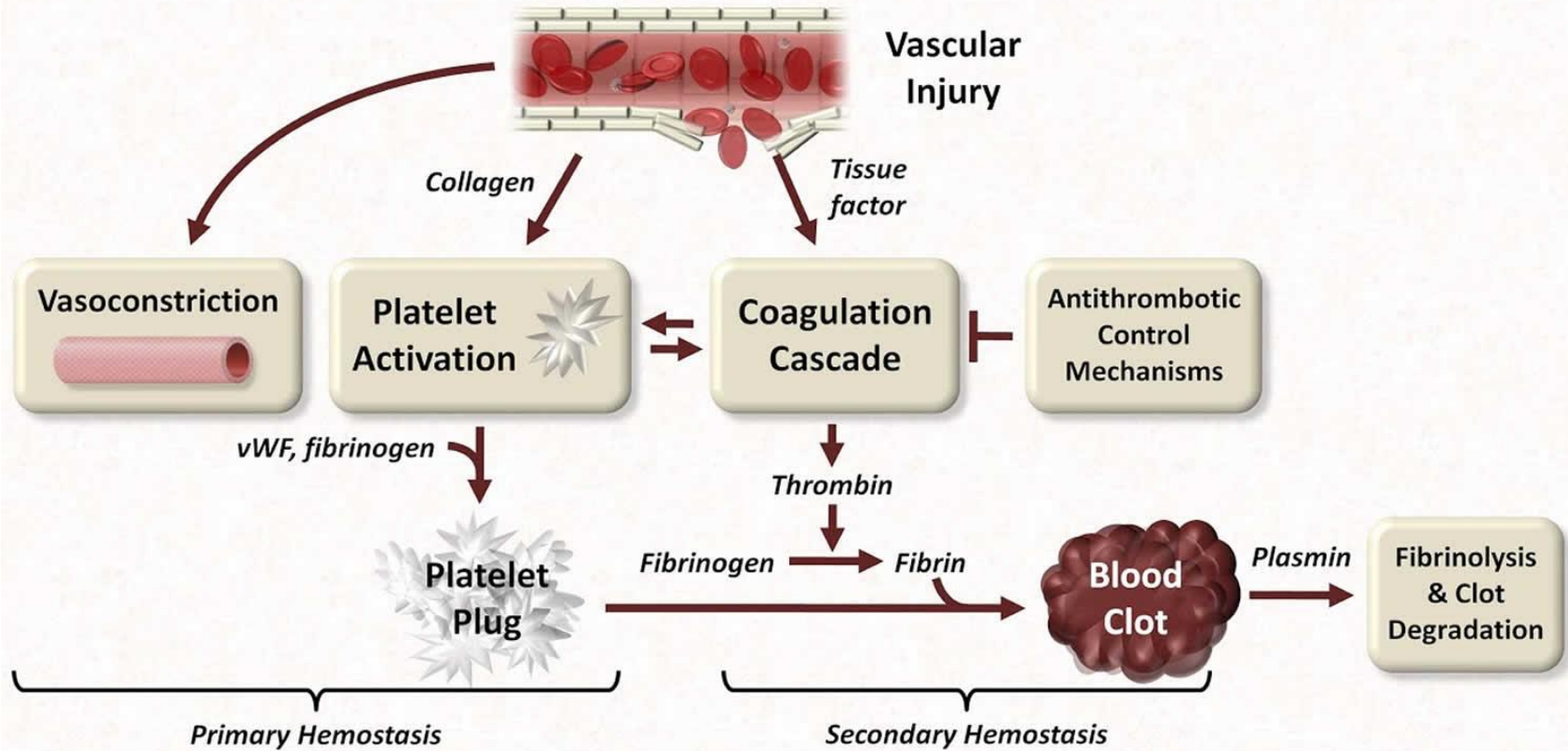
# THROMBOSIS

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21-12-2022.



# Major Components of Hemostasis

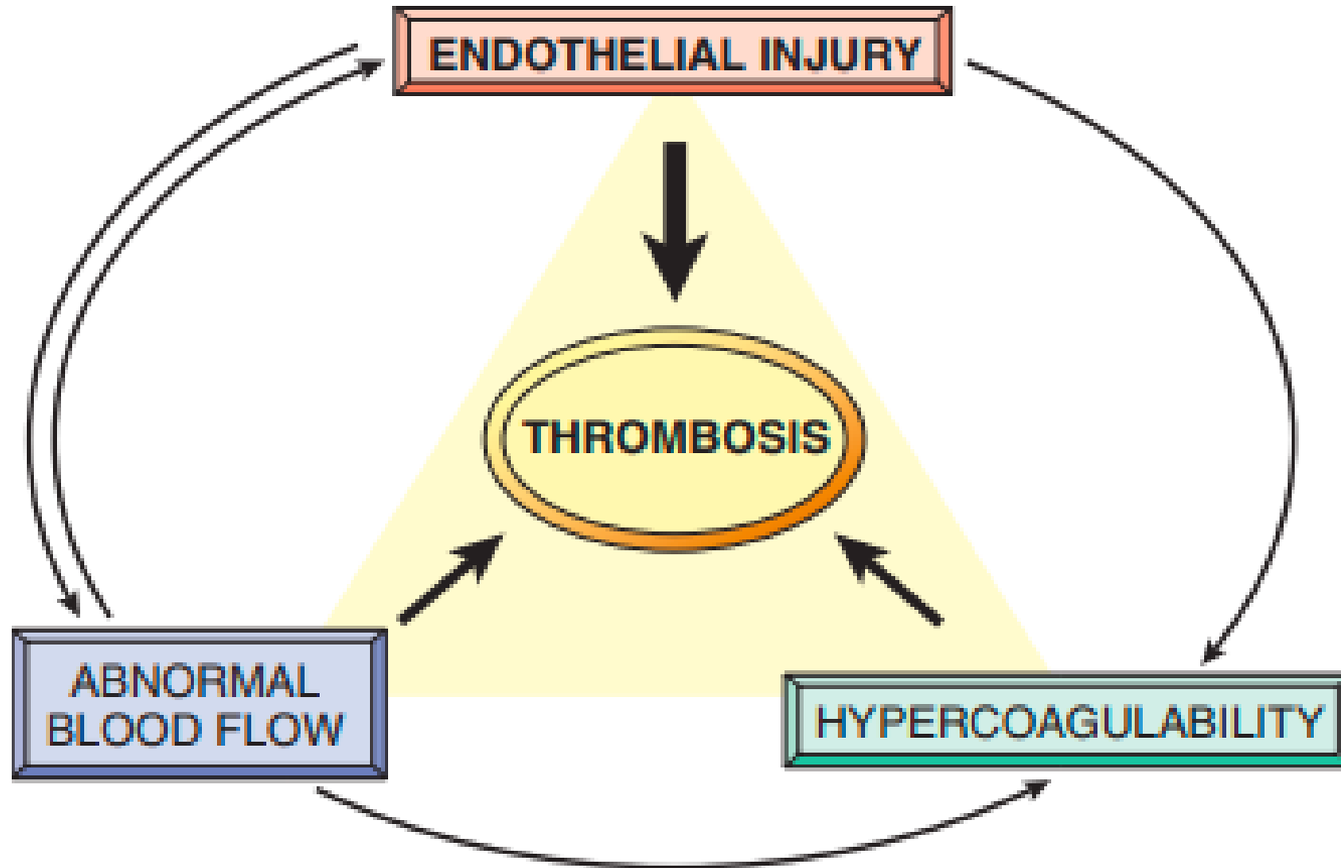


# THROMBOSIS

- **Thrombosis** is the formation of a blood clot, known as a thrombus, within a blood vessel.
- The primary abnormalities that lead to intravascular thrombosis are:
  - ❖ (1) endothelial injury.
  - ❖ (2) stasis or turbulent blood flow.
  - ❖ (3) hypercoagulability of the blood



➤ (the so-called “Virchow triad”)

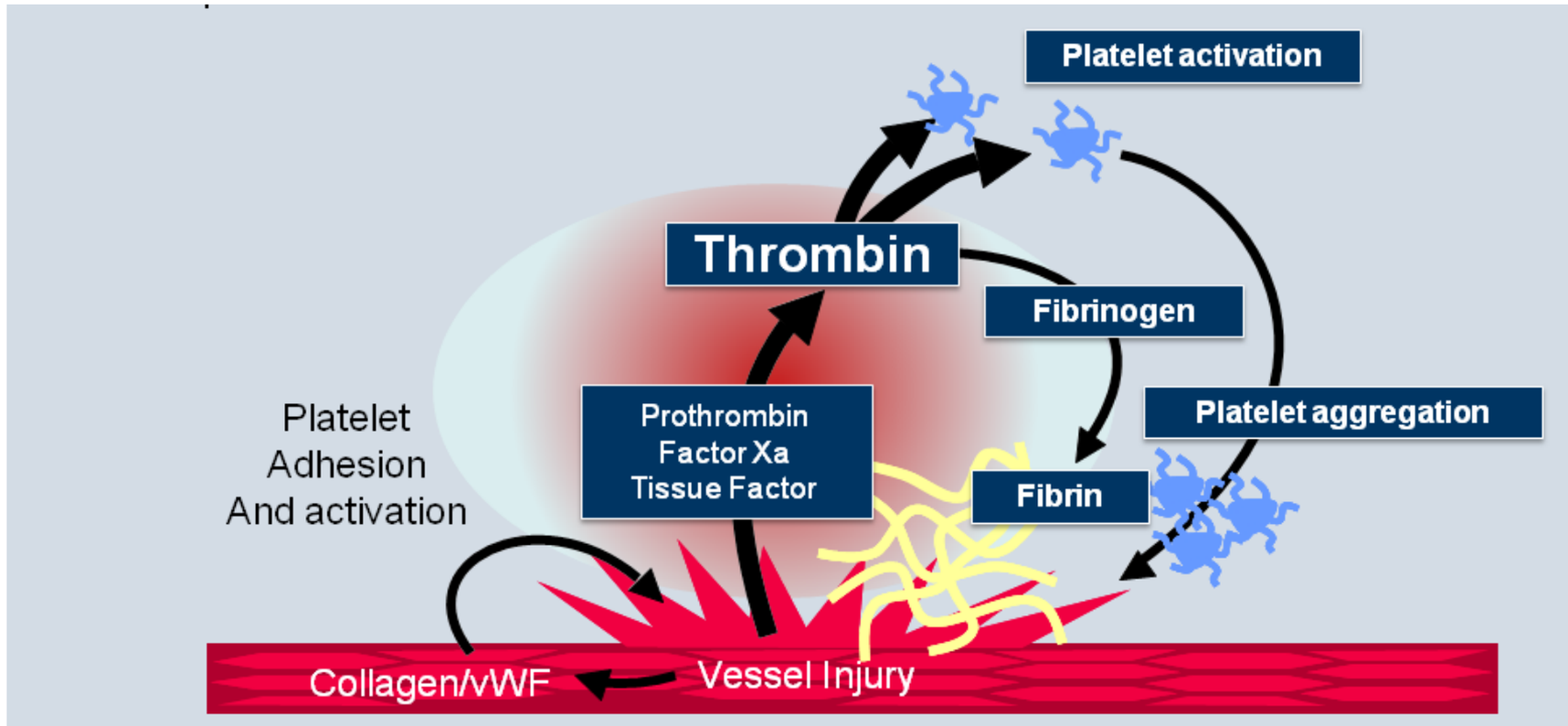


# 1. ENDOTHELIAL INJURY

- Endothelial injury leading to platelet activation.
- cardiac and arterial clots are typically rich in platelets.
- Endothelial injury may be caused by:
  - physical injury.
  - infectious agents.
  - inflammatory mediators.
  - metabolic abnormalities, such as hypercholesterolemia or homocystinemia,
  - toxins absorbed from cigarette smoking.



severe endothelial injury may trigger thrombosis by exposing VWF and tissue factor



- Endothelial injury .....that's mean:
  - platelet activation: exposed VWF, TF.
  - Endothelial dysfunction : prothrombotic endothelium.
  
  - Procoagulant changes:
    - ↓thrombomodulin, ↑thrombin
  
  - Anti-fibrinolytic effects:
    - ↑ Plasminogen activator inhibitors (PAI).
    - ↓t-PA.



## Plasminogen activators

Tissue plasminogen activator (tPA),  
urokinase, streptokinase  
staphylokinase, vampire bat PA

## PAI-1

(inhibits tPA,  
present in blood  
in small concentration)

Cleave  
plasminogen into  
active plasmin

**Plasminogen** → **Plasmin**

Degrades  
fibrin clot,  
core of  
thrombus

$\alpha_2$ -antiplasmin  
(physiological inhibitor of  
plasmin present in blood in concentration  
6-8X exceeding therapeutic dose of plasmin)

**Fibrin**

**Fibrin degradation  
products**

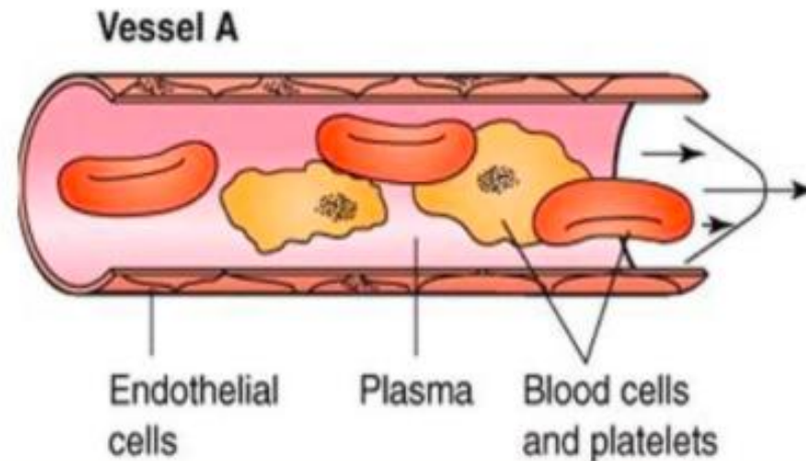


## 2. ABNORMAL BLOOD FLOW

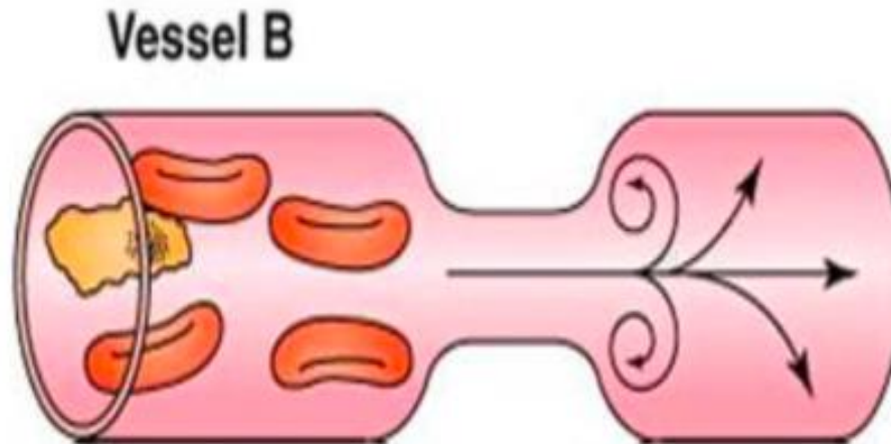
- Stasis is a major factor in the development of venous thrombus.
- stasis and turbulence have the following effects:
  - Stasis allows platelets and leukocytes to come into contact with the endothelium when the flow is sluggish.
  - Stasis also slows the washout of activated clotting factors and impedes the inflow of clotting factor inhibitors.



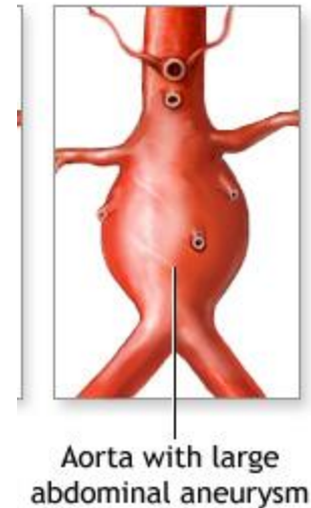
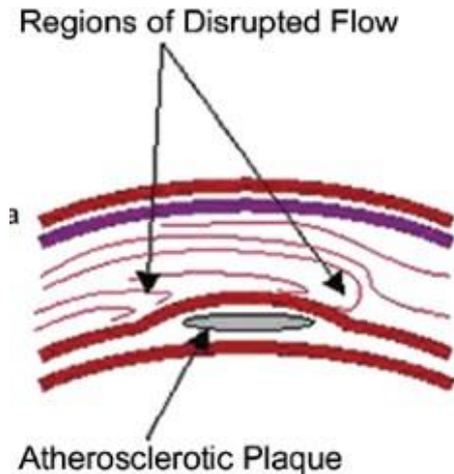
- Normally **laminar**, with the blood components arranged in layers
- The plasma forms the outer layer & slides smoothly along the endothelium
- Blood cells form the 'axial' layer in the centre of the blood stream
- This allows the blood to flow smoothly, layers slide over each other, axial part moves fastest.



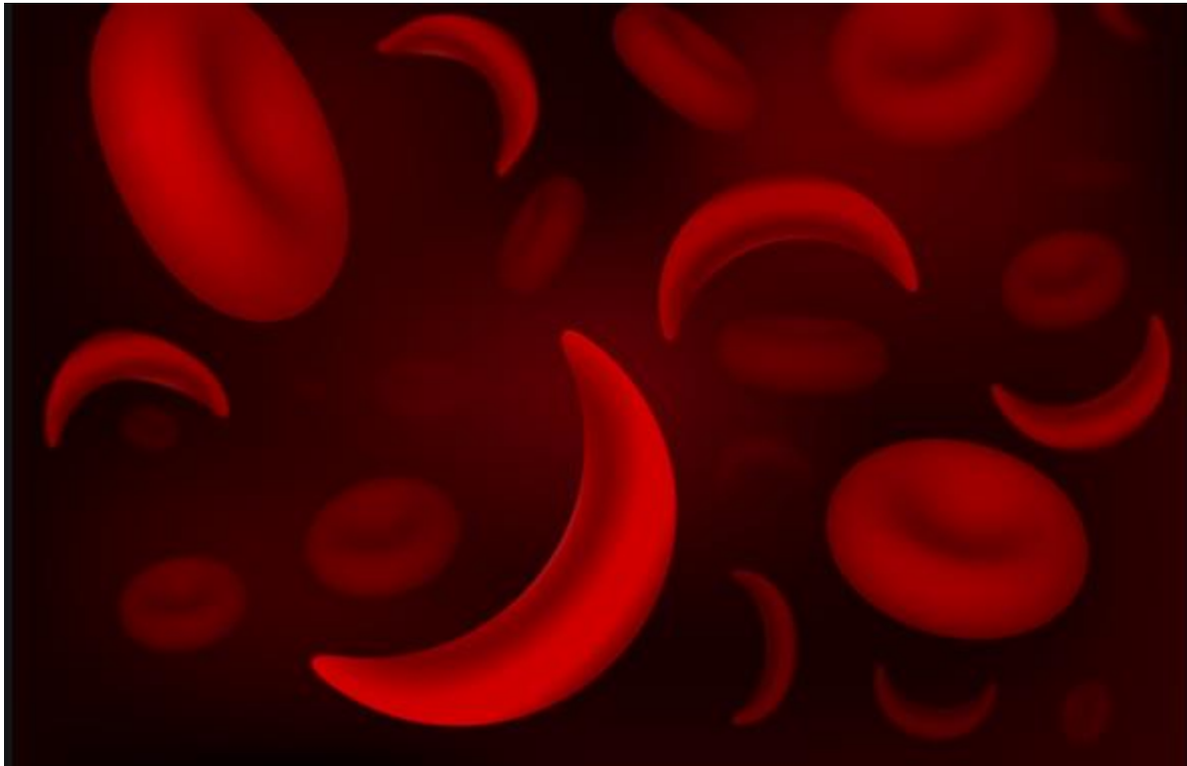
- When we take a blood pressure the sounds we here are caused by **turbulent flow** of blood
- **Turbulent flow** -caused by change in vessel diameter, increase in velocity, & low blood viscosity



- Turbulent and static blood flow contributes to thrombosis in a number of clinical setting:
- 1. Ulcerated atherosclerotic plaques.
- 2. aortic aneurysms create local stasis and consequently are fertile sites for thrombosis



- 3. Hyperviscosity syndromes (such as polycythemia vera )increase resistance to flow and cause small vessel stasis.
- 4. sickle cell anemia:
  - The deformed red cells in cause vascular occlusions.



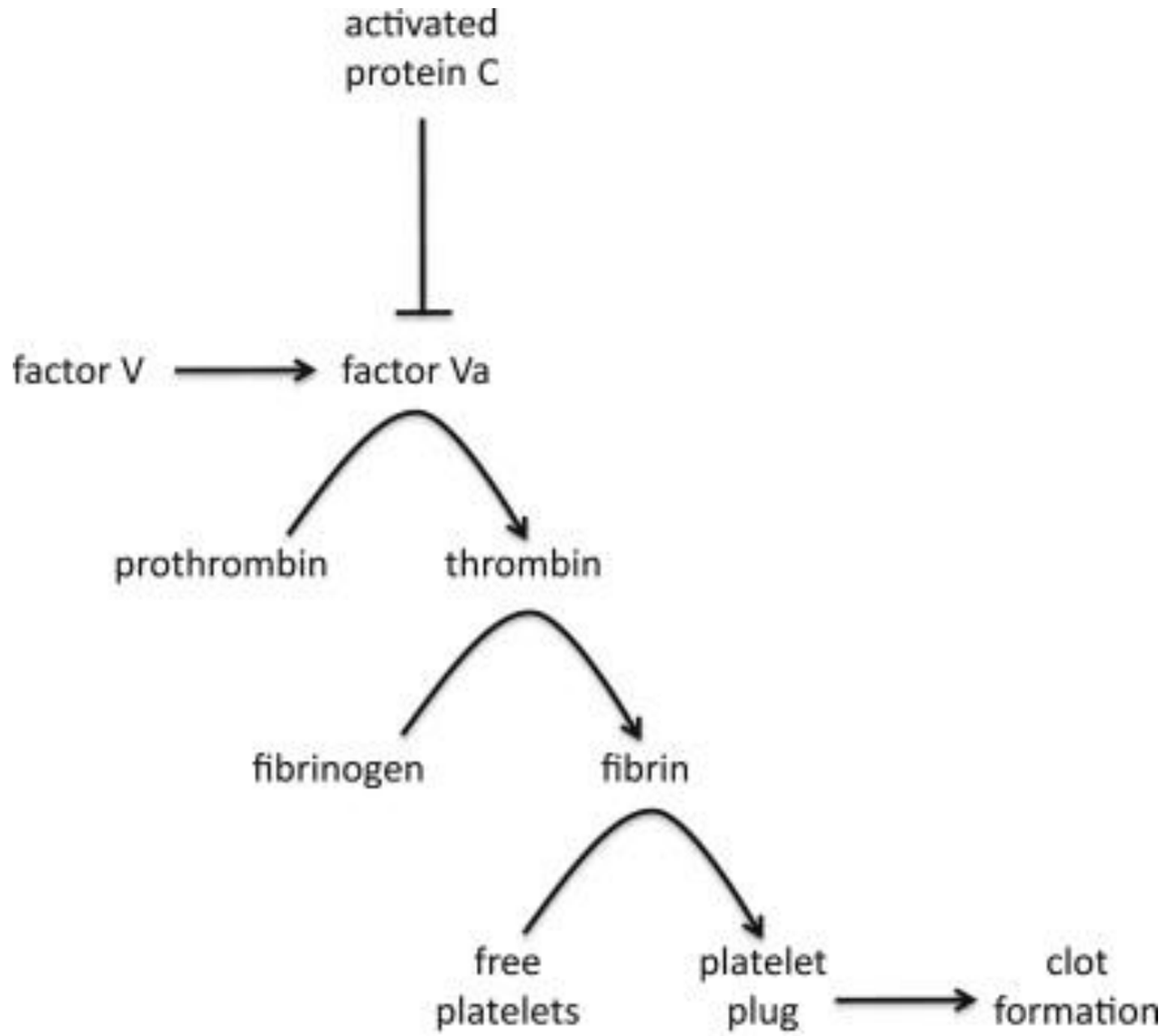
### 3. HYPERCOAGULABILITY

- Hypercoagulability refers to an abnormally high tendency of the blood to clot, and is typically caused by alterations in coagulation factors.
- Is an important underlying risk factor for venous thrombosis



- The alterations of the coagulation pathways that predispose affected persons to thrombosis can be divided into:
  - Primary (genetic):
    - ❖ mutations in the factor V .
    - ❖ elevated levels of homocysteine \*
  - secondary (acquired) disorders:
    - ❖ oral contraceptive.
    - ❖ Pregnancy\*\*.
    - ❖ Cancers\*\*\*.



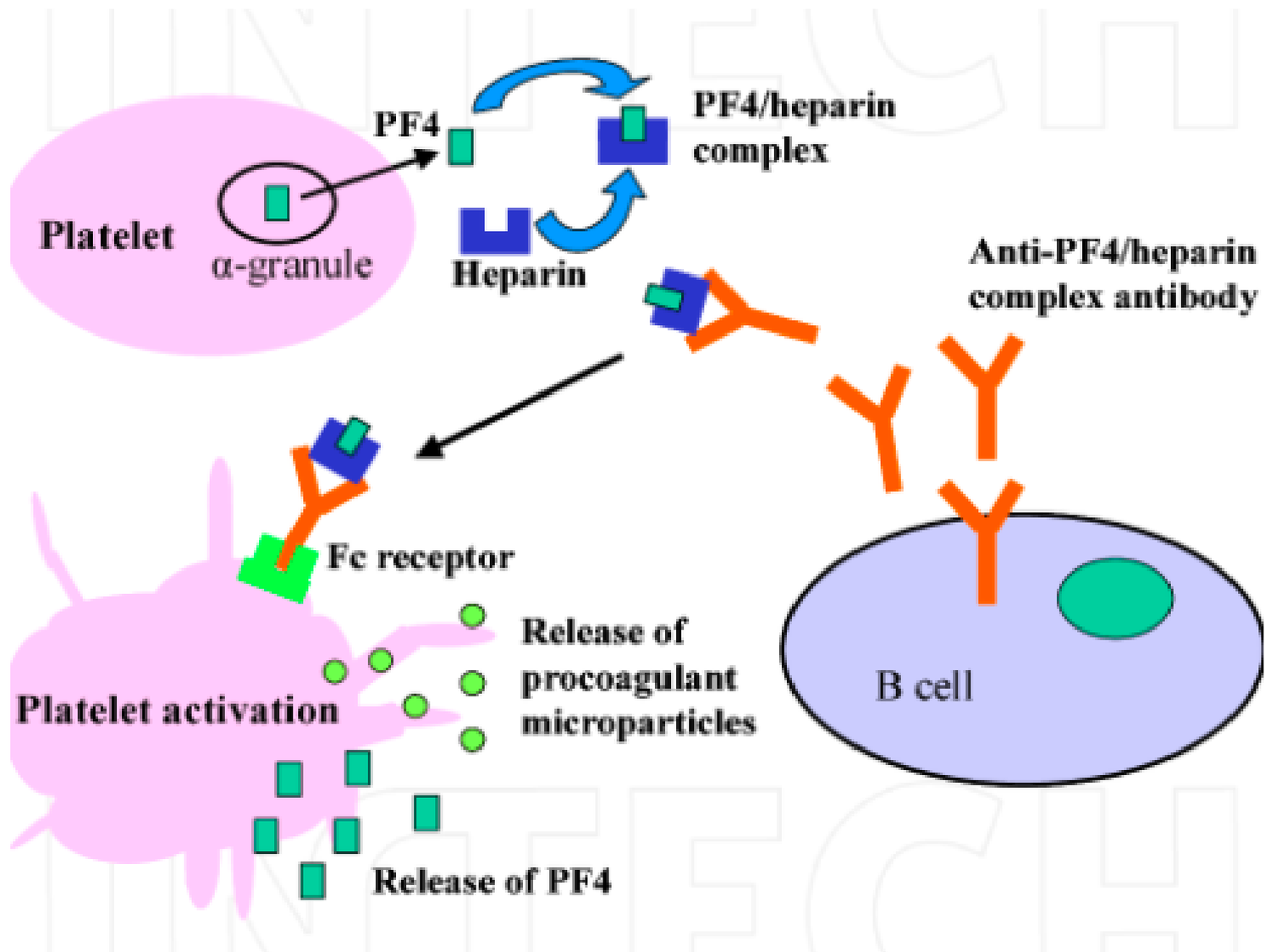


- Among the acquired thrombophilic states, two are particularly important clinical problems and deserve special mention:



- 1. Heparin-induced thrombocytopenia (HIT) syndrome:
  - occurs in patients treated with unfractionated heparin.
  - It is marked by the development of autoantibodies that bind complexes of heparin and platelet membrane protein (platelet factor-4).
  - Its resulting in platelet activation, aggregation, and consumption (hence thrombocytopenia), as well as causing endothelial cell injury.





- 2. Anti-phospholipid antibody syndrome:
- Acquired antibodies against phospholipid - protein complexes.
- Suspected antibody targets include  $\beta$ 2-glycoprotein I, a plasma protein that associates with the surfaces of endothelial cells, trophoblasts, and prothrombin
- clinical manifestations, including :
  - recurrent thromboses.
  - repeated miscarriages.
  - cardiac valve vegetations.
  - thrombocytopenia



- Depending on the vascular bed involved, the clinical presentations can include:
- pulmonary embolism (following lower extremity venous thrombosis).
- pulmonary hypertension (from recurrent subclinical pulmonary emboli)
- Stroke.
- bowel infarction.
- renovascular hypertension.

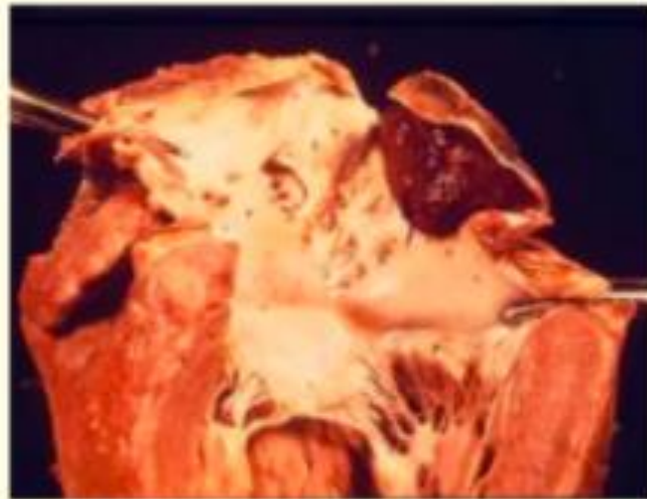
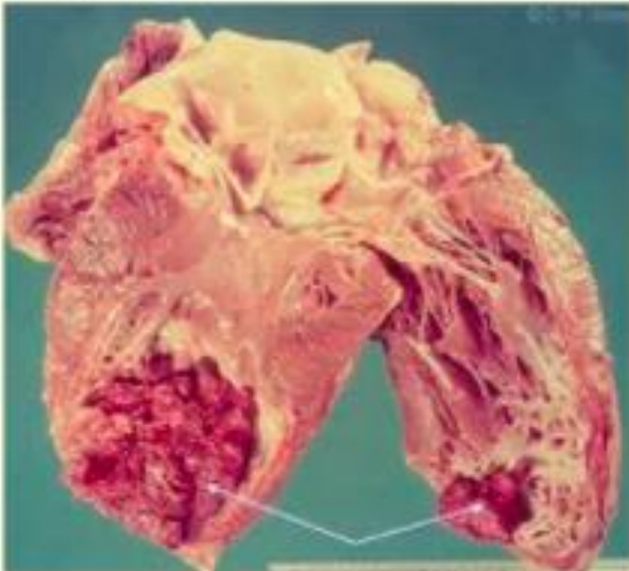


# MORPHOLOGY

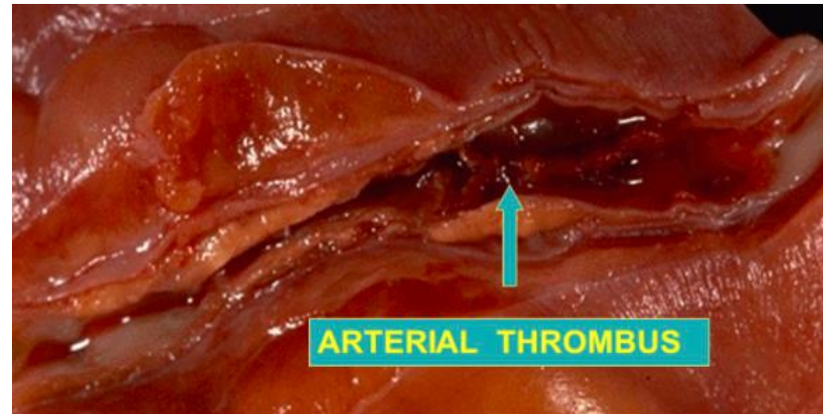
- Thrombi can develop anywhere in the cardiovascular system.
- Arterial or cardiac thrombi typically arise at sites of endothelial injury or turbulence.
- venous thrombi characteristically occur at sites of stasis.



- Mural thrombi:
- Thrombi occurring in heart chambers or in the aortic lumen



\*Arterial thrombi are frequently occlusive. They are typically rich in platelets

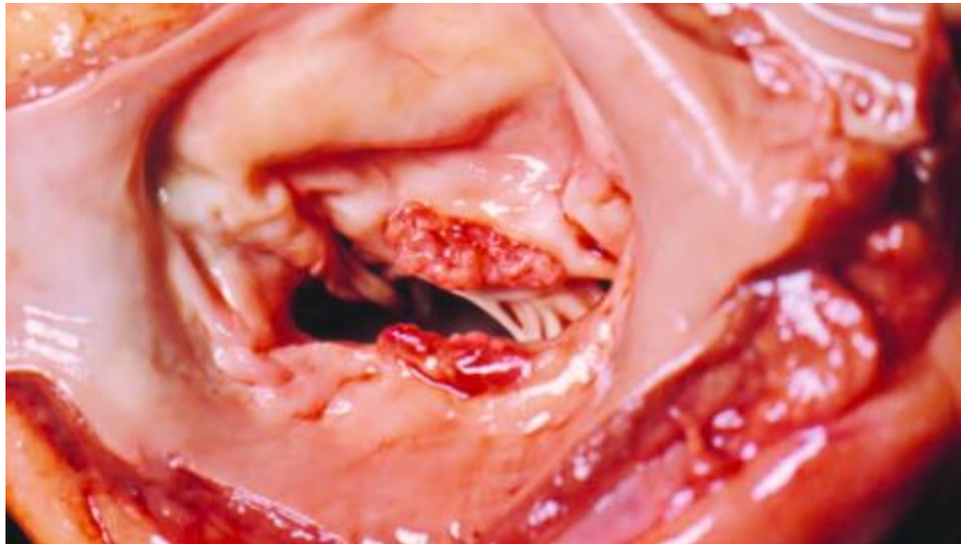


## VENOUS THROMBI (PHLEBOTHROMBOSIS):

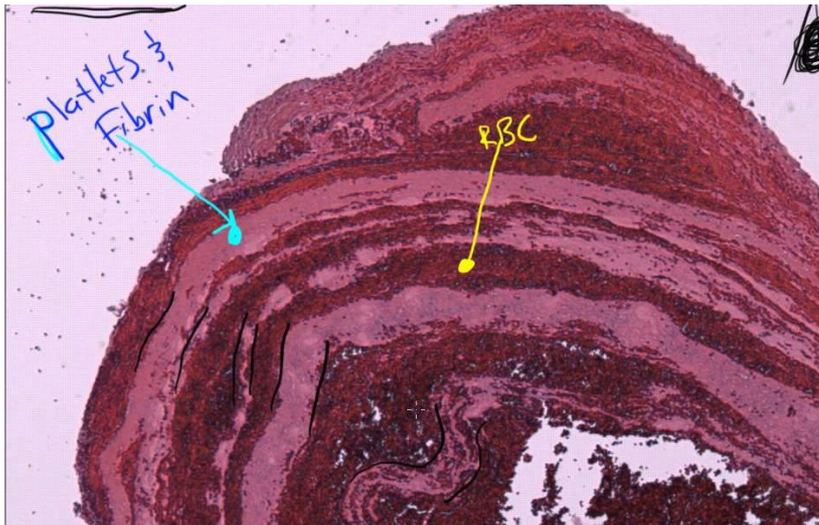
- they frequently propagate some distance toward the heart, forming a long cast within the vessel lumen that is prone to give rise to emboli.
- they tend to contain more red cells.

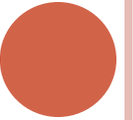


- Vegetations: Thrombi on heart valves , divided into :
  - infective endocarditis: Infective thrombotic masses
  - nonbacterial thrombotic endocarditis: Sterile vegetations.
  - LibmanSacks endocarditis:
    - Sterile, occur in the systemic lupus erythematosus.



- Thrombi can have grossly (and microscopically) apparent laminations called lines of Zahn; these represent
- pale platelet and fibrin layers alternating with darker red cell-rich layers.





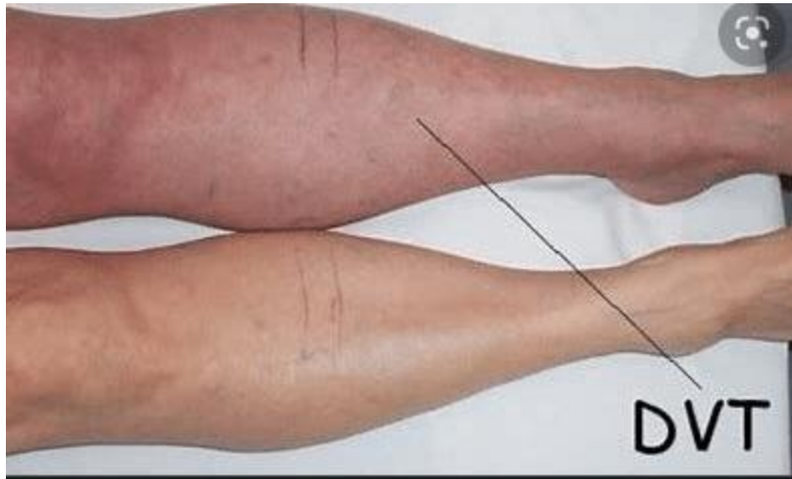
# FATE OF THE THROMBUS

- 1. Propagation: The thrombus enlargement.
- 2. Embolization: transported in the vasculature.
- 3. Dissolution: shrinkage and complete dissolution.
- 4. Organization
  - ingrowth of endothelial cells, smooth muscle cells, and fibroblasts.
- 5. Recanalization: capillary channels are formed create canal along the length of the thrombus, thereby reestablishing the continuity of the original lumen.

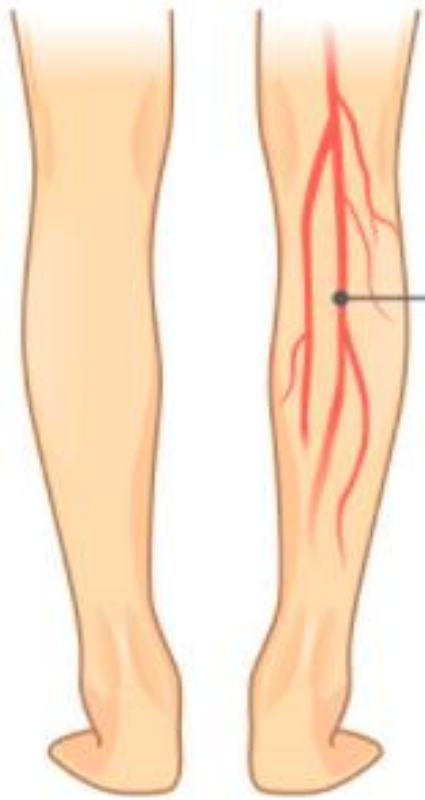


# CLINICAL FEATURES

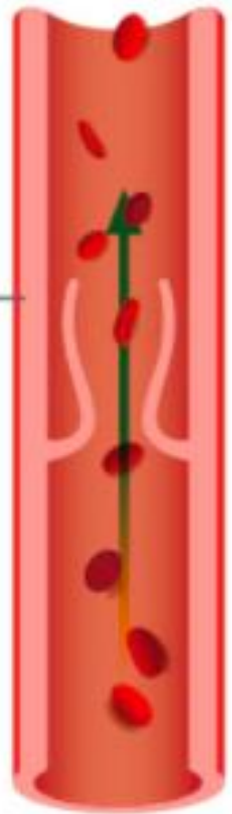
- 1. Venous Thrombosis (Phlebothrombosis)
- Pain.
- local congestion and swelling from impaired venous outflow.
- varicose ulcers.
- rarely embolize.



Deep Veins of the Leg



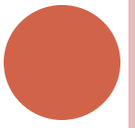
Normal Blood Flow



Deep Vein Thrombosis



Embolus



- 2. Arterial and Cardiac Thrombosis;
- Atherosclerosis is a major cause of arterial thromboses because it is associated with the loss of endothelial integrity and with abnormal blood flow.
- Both cardiac and aortic mural thrombi are prone to embolization. . The brain, kidneys, and spleen are particularly likely targets because of their rich blood supply.





**thrombosis** of a coronary artery

Massive thrombosis (*arrows*) from distal portion of left main coronary artery



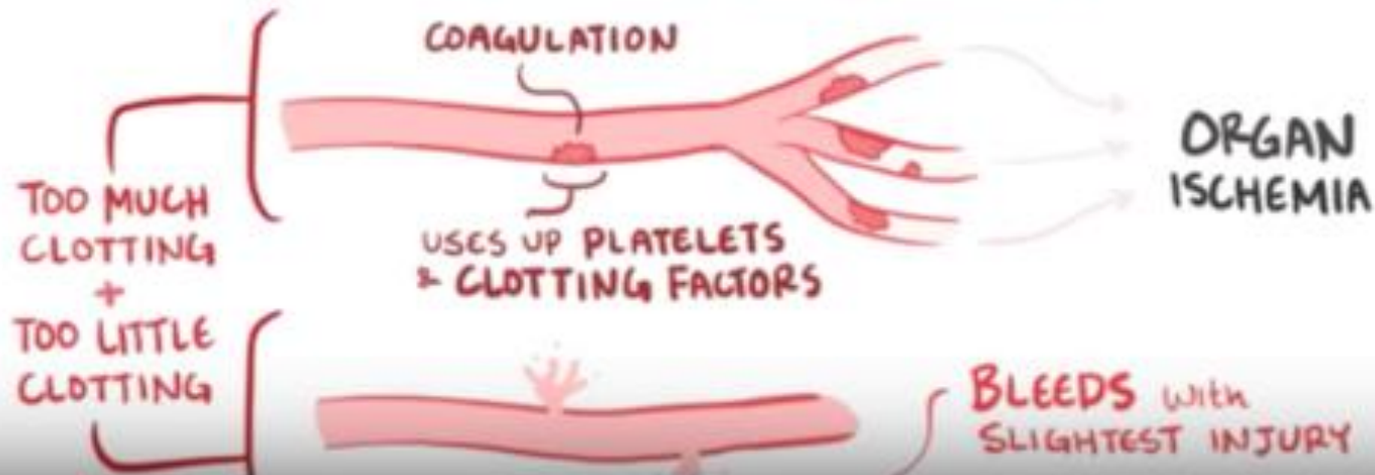
# DISSEMINATED INTRAVASCULAR COAGULATION (DIC)

- DIC is widespread thrombosis within the microcirculation that may be of sudden or insidious onset.
- DIC = widespread microvascular thrombosis + fibrinolytic mechanisms activation.
- DIC consumes platelets and coagulation proteins (hence the synonym consumptive coagulopathy).



# DISSEMINATED INTRAVASCULAR COAGULATION (DIC) ( CONSUMPTION COAGULOPATHY )

\* HEMOSTASIS ~ OUT OF CONTROL





Excessive hemorrhage (Bruising, petechiae).

