



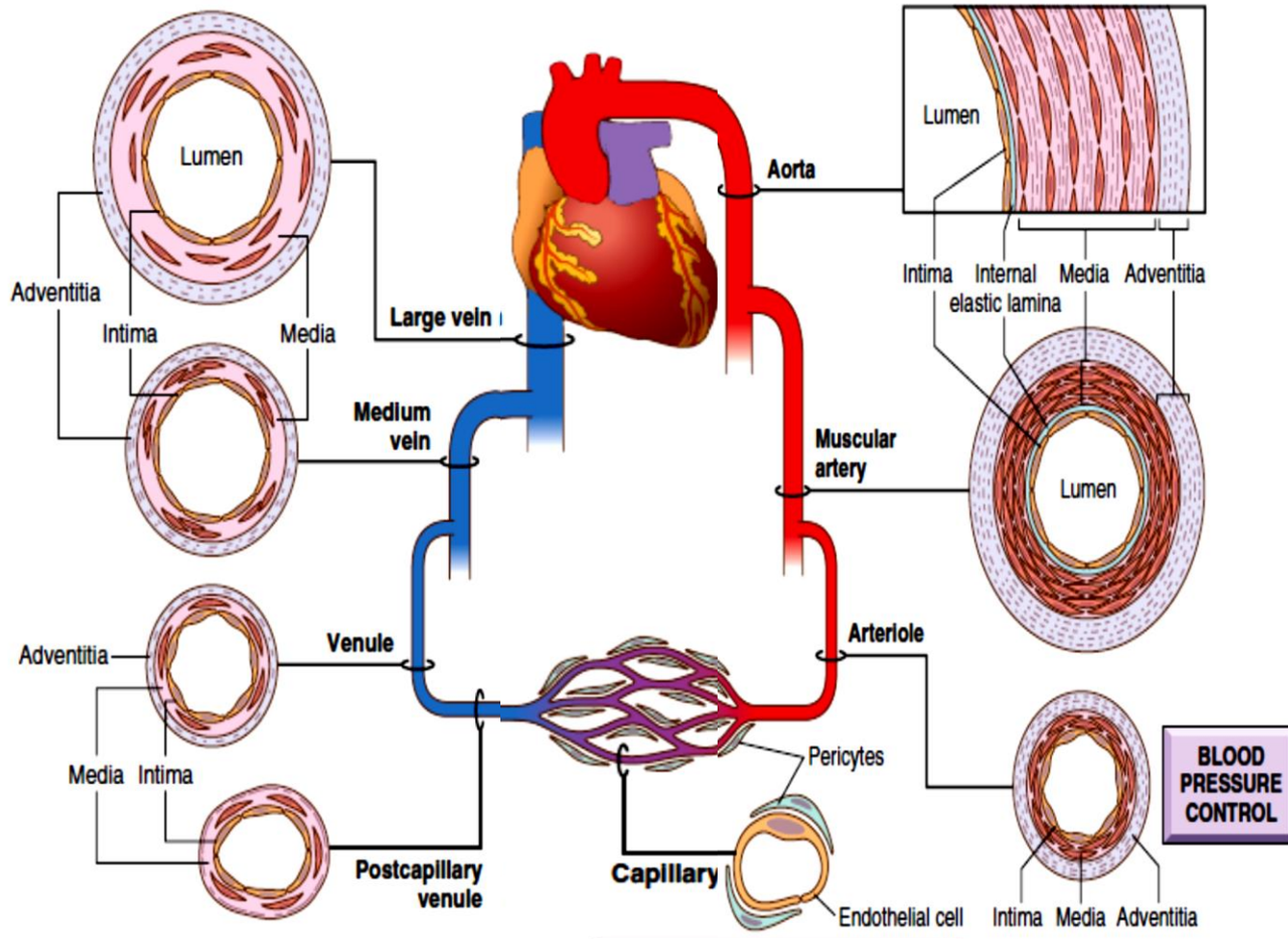
# **THROMBOSIS**

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**University of Jordan**

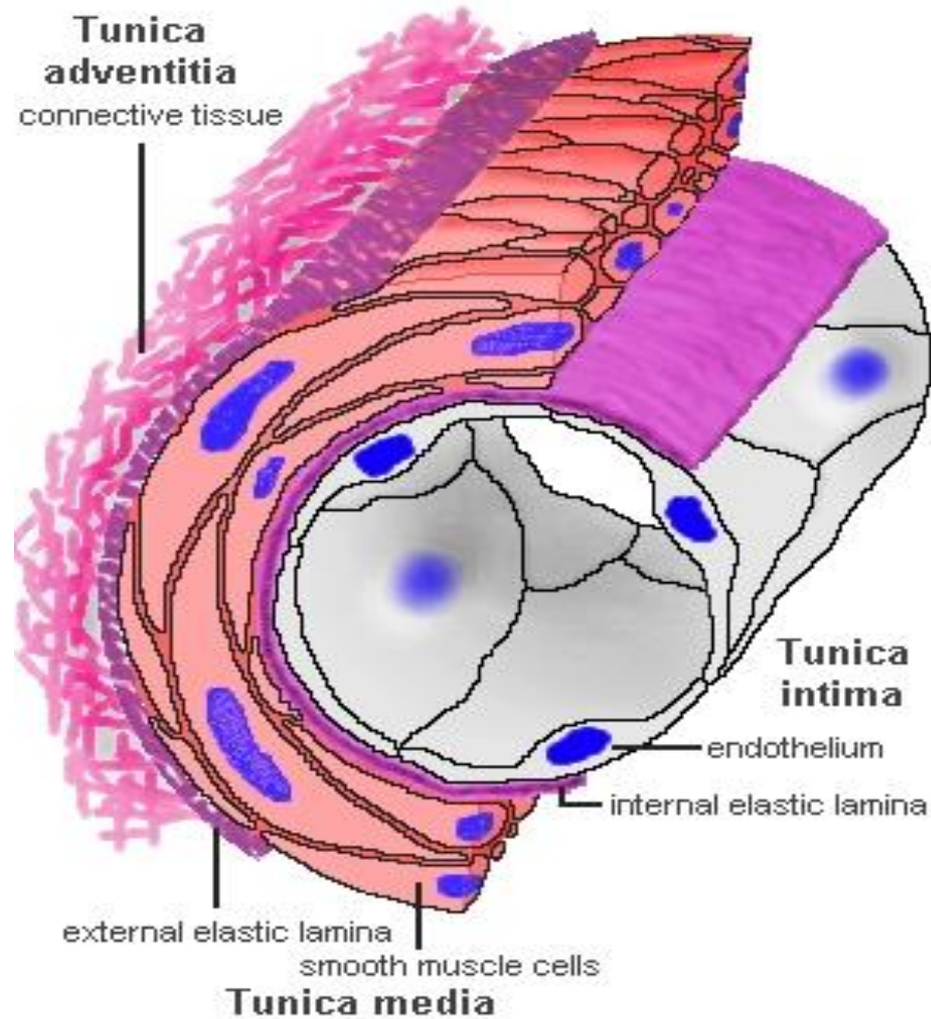
# CARDIOVASCULAR SYSTEM

## Venous circulation

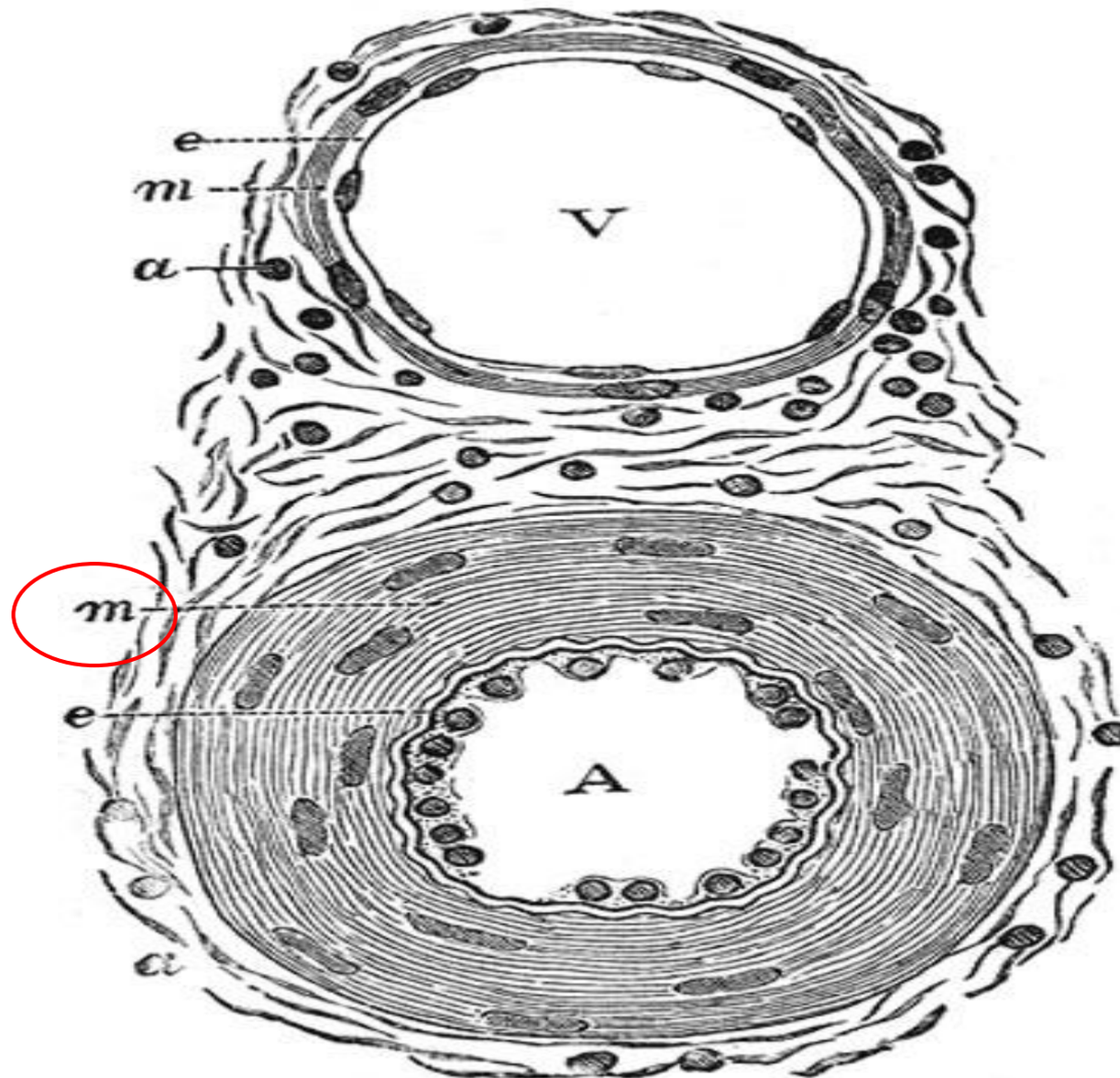
## Arterial circulation



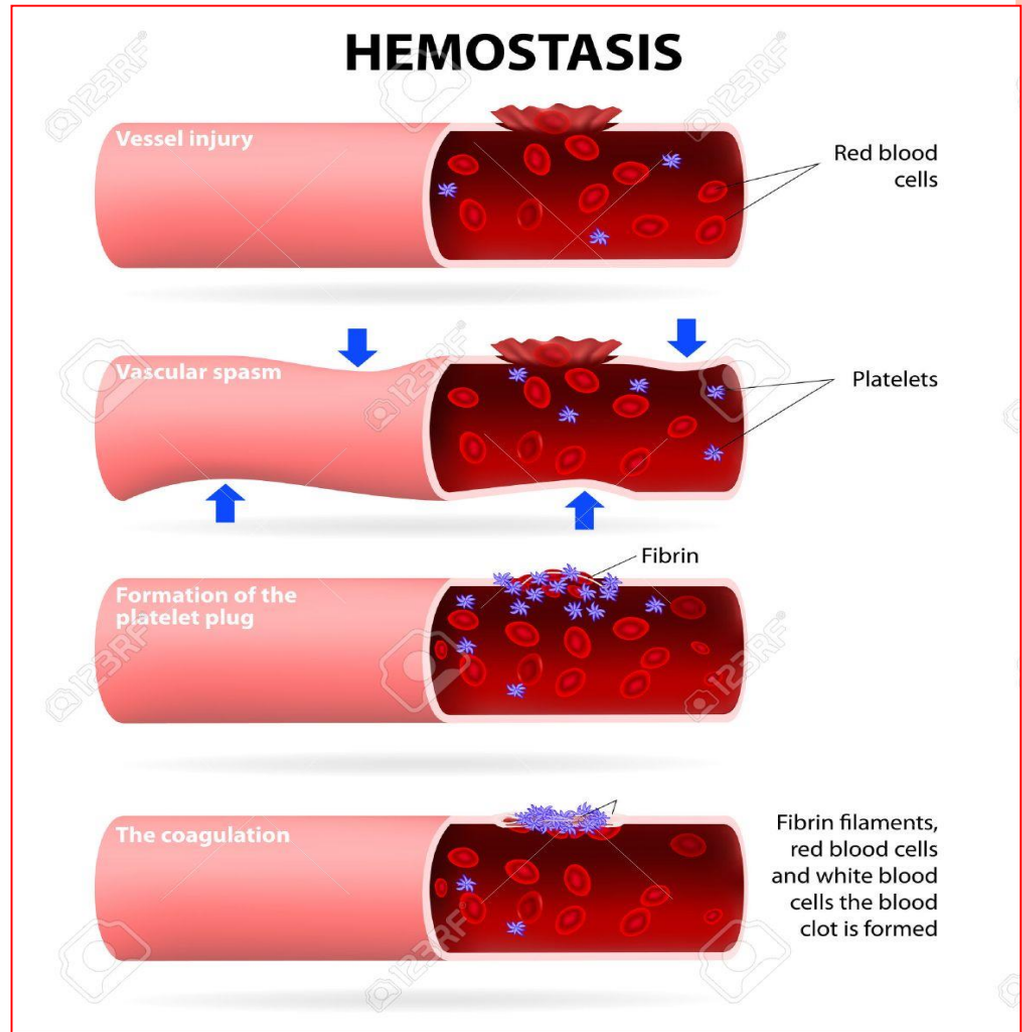
# NORMAL BLOOD VESSEL HISTOLOGY



# ARTERY (A) VS VEIN (V)



# PHYSIOLOGY OF THROMBOSIS

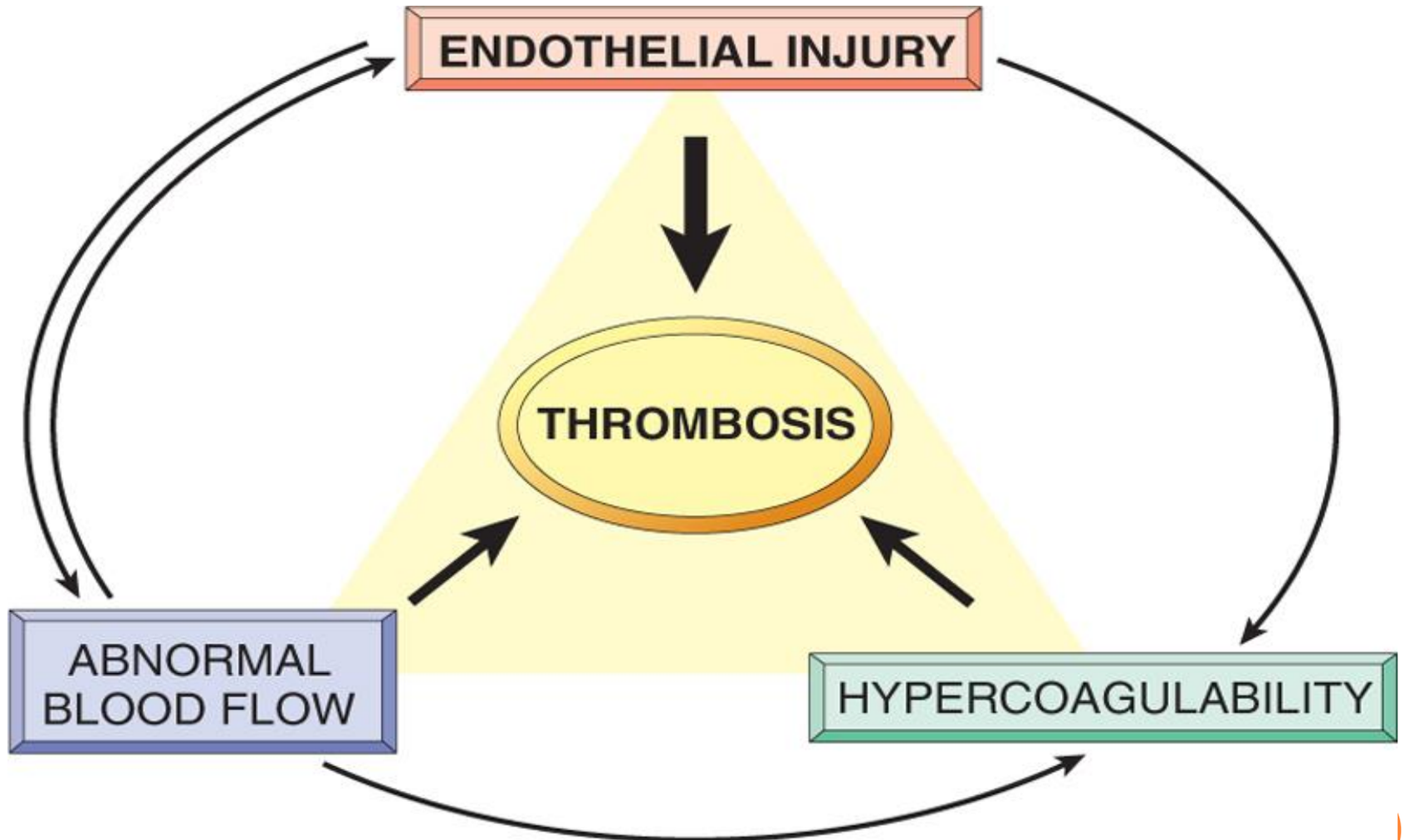


# THROMBOSIS- PATHOLOGICAL ASPECTS

- Blood coagulation is a very important physiological event to protect our hemostasis, and life
- **However**, at certain points, this process can be pathological that may endorse injury and cause harm to our body
- This happens whenever unnecessary blood clotting is activated
- **The “pathological” thrombosis is caused by the presence of at least one of 3 factors (together called Virchow’s triad):**



## Virchow's triad



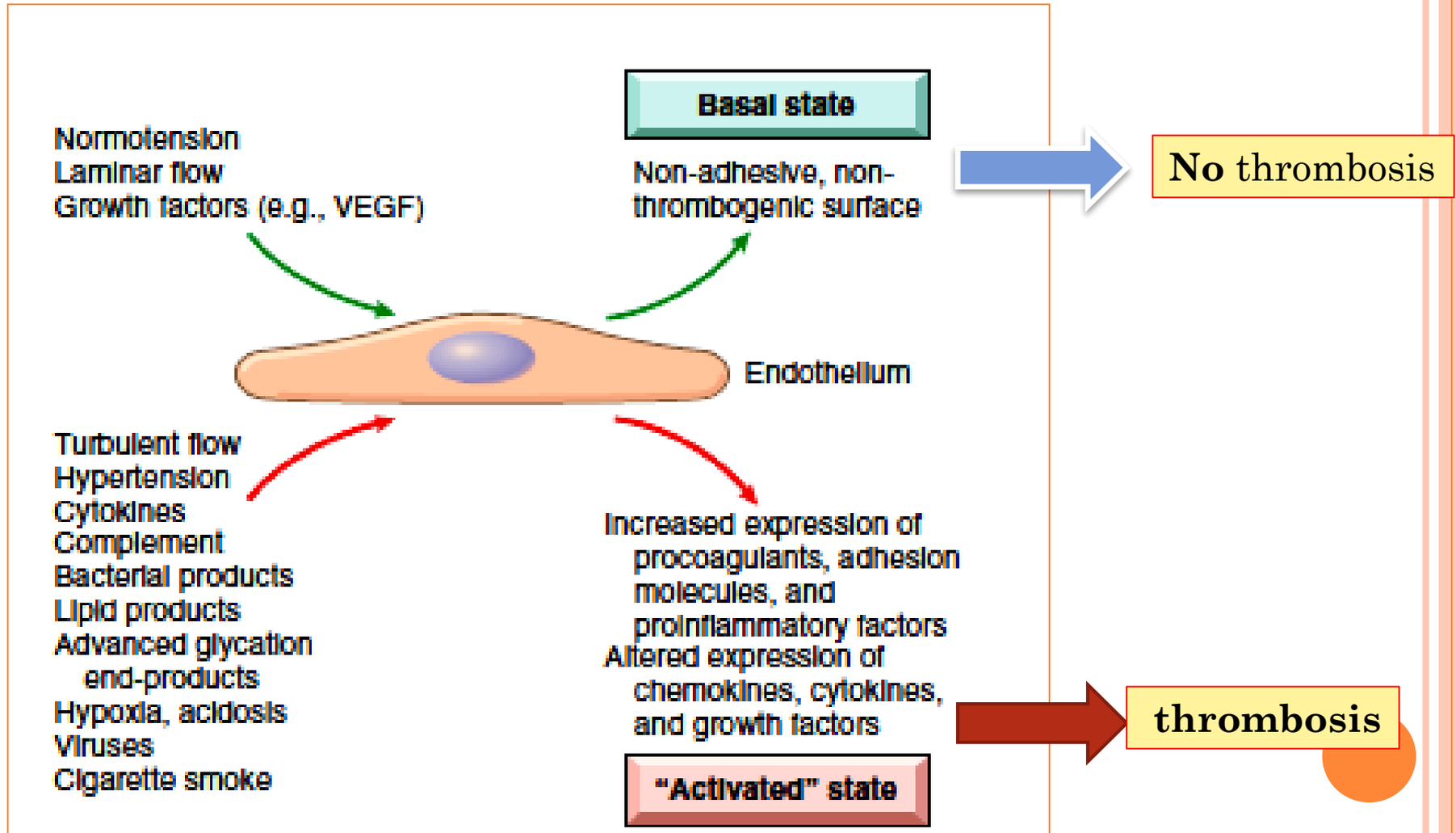
# THROMBOSIS- PATHOLOGICAL ASPECTS

- Pathogenesis (called *Virchow's triad*):
  1. *Endothelial\* Injury ( Heart, Arteries)*
  2. *Stasis (abnormal blood flow)*
  3. *Blood Hypercoagulability*

\* Endothelial cells are special type of cells that cover the inside surface of blood vessels and heart.



# CONTRIBUTION OF ENDOTHELIAL CELLS TO COAGULATION



Endothelial Cell Injury and exposure of subendothelial collagen



Adherence of platelets



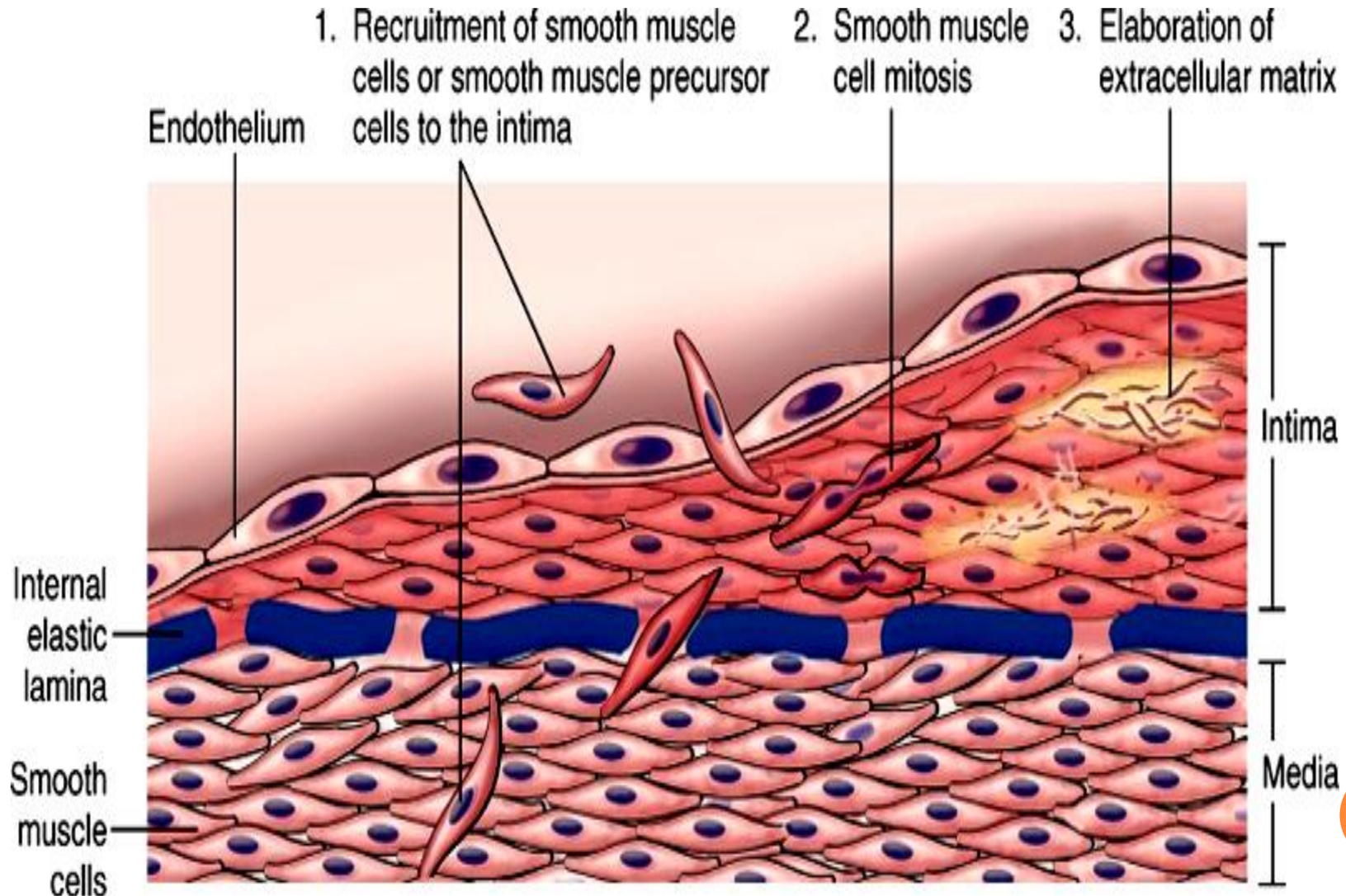
Release of tissue factor



Progression of coagulation event .....



# Response of Vascular Wall Cells to Injury



# RESPONSE OF VASCULAR WALL CELLS TO INJURY

- Injury results in a **healing response**
- Pathologic effect of **vascular healing**:

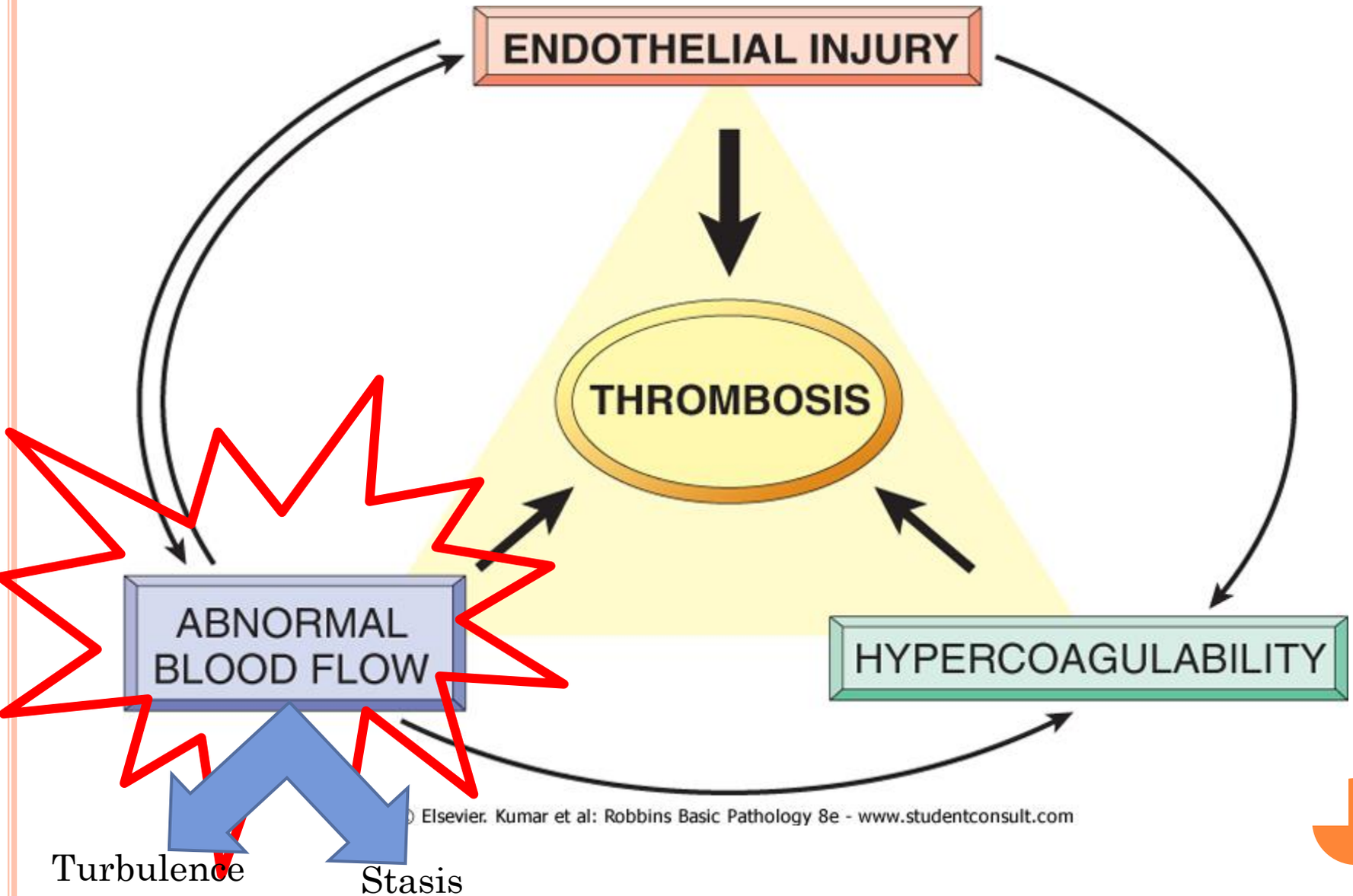
**Excessive thickening of the intima →→  
luminal stenosis & blockage of  
vascular flow**



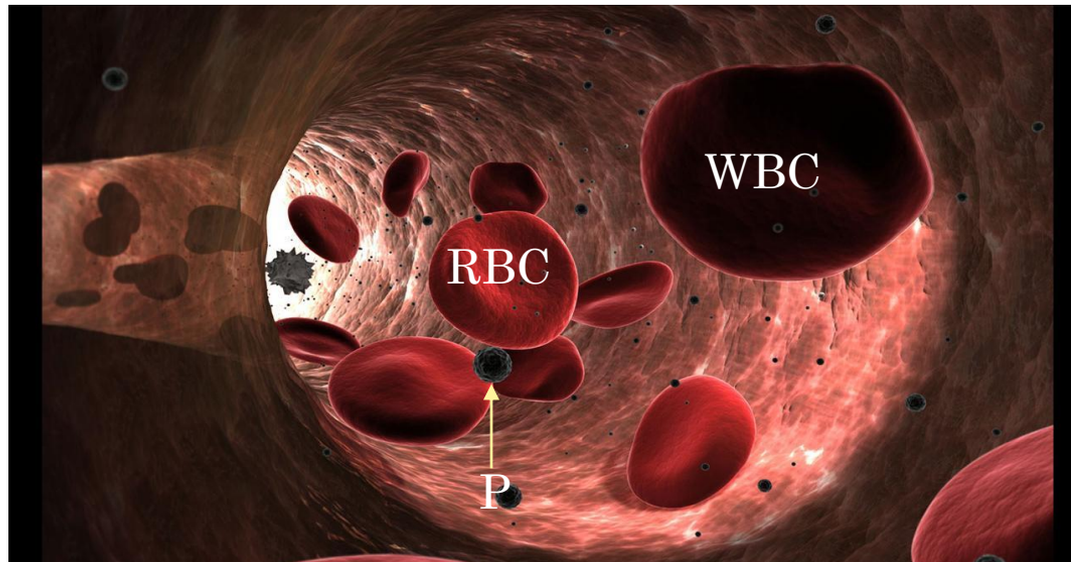
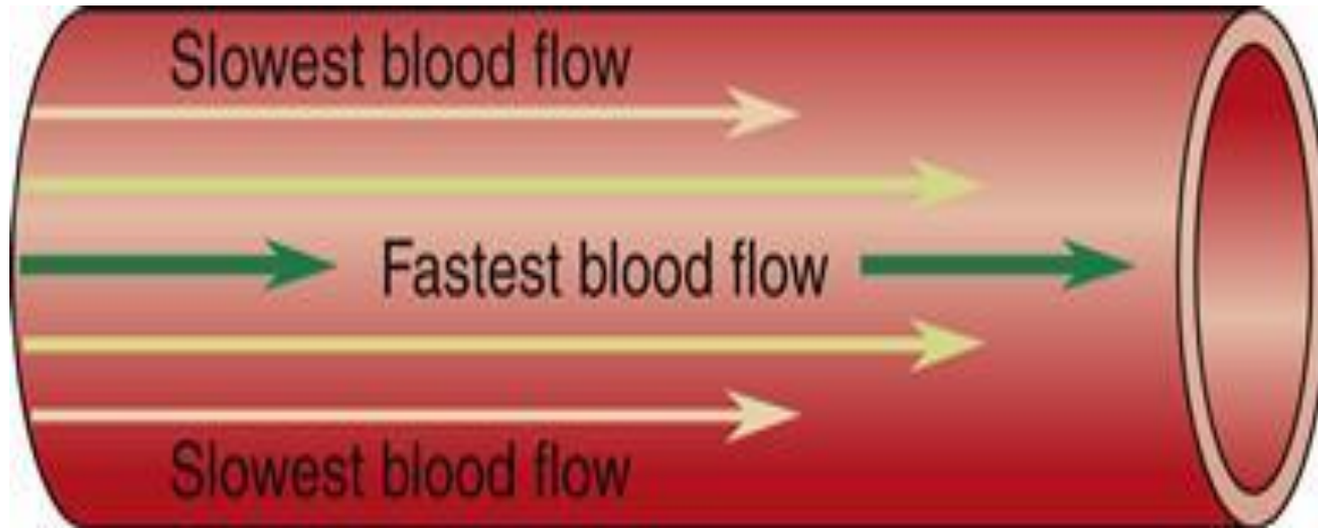
## ○ Causes of Endothelial injury

1. *Valvulitis*
2. *MI*
3. *Atherosclerosis*
4. *Traumatic or inflammatory conditions*
5. *Hypertension*
6. *Endotoxins*
7. *Hypercholesterolemia*
8. *Radiation*
9. *Smoking*
10. *.....etc.*

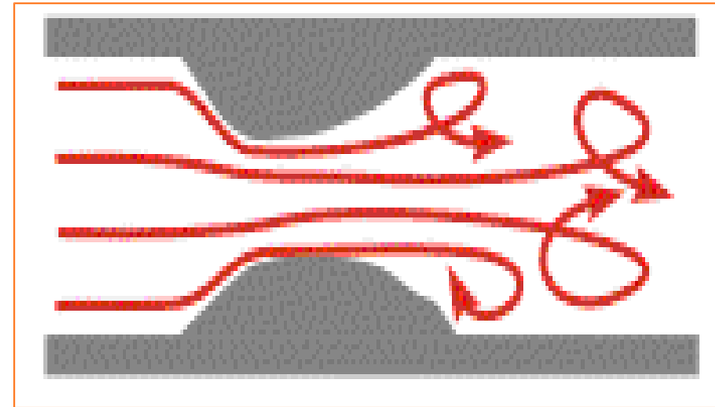
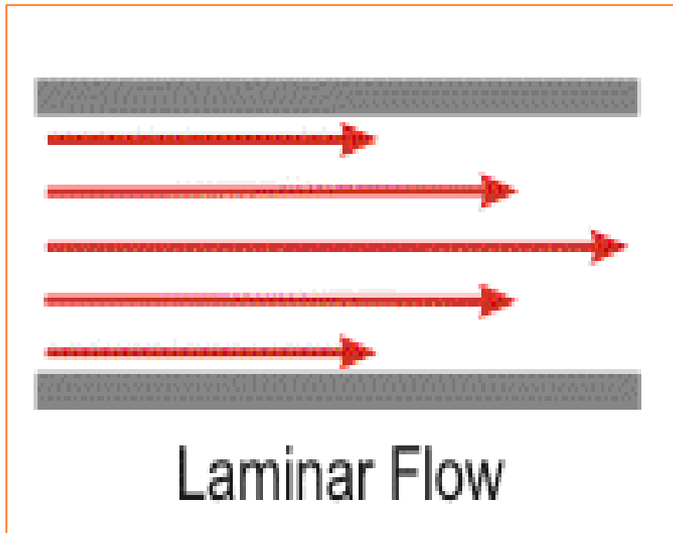




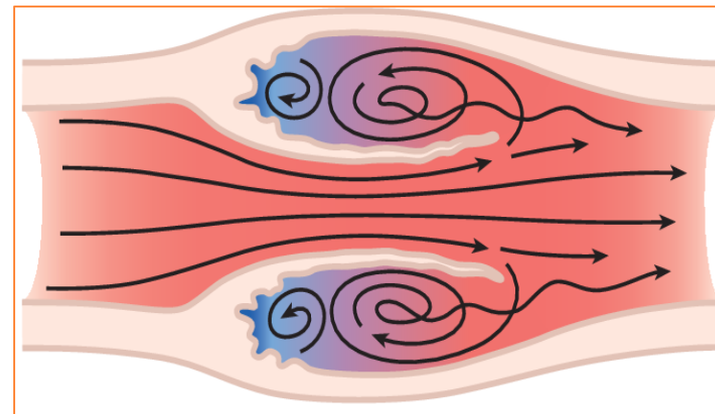
# LAMINAR BLOOD FLOW



# LAMINAR VS TURBULENT BLOOD FLOW



Turbulent Flow



# ○ Stasis

- *Stasis is a major factor in **venous** thrombi*
- Normal blood flow is *laminar* (platelets flow centrally in the vessel lumen, separated from the endothelium by a slower moving clear zone of plasma)
- Stasis and turbulence cause the followings:

Stasis and  
turbulence

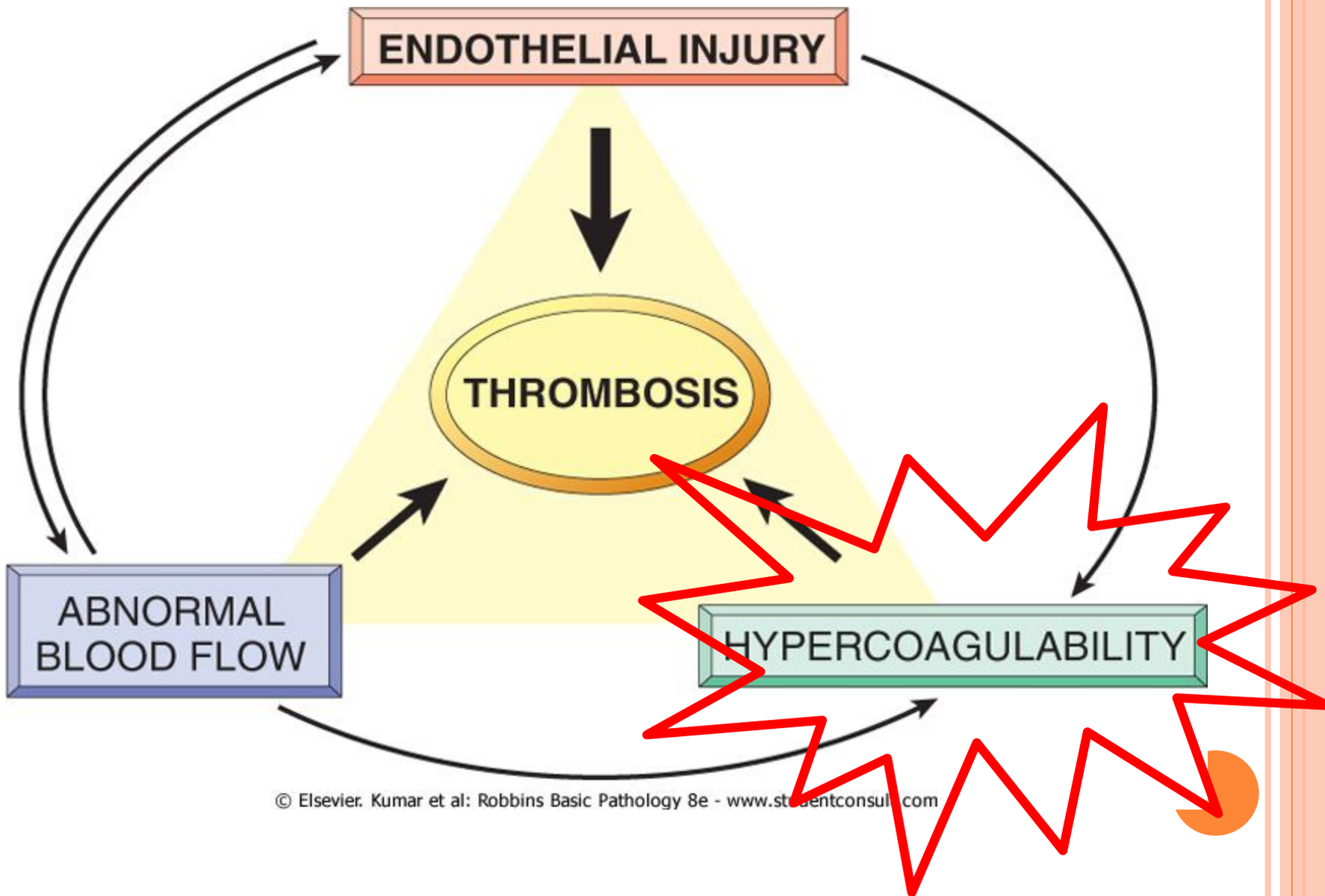
- Disrupt normal blood flow
- Prevent dilution of activated clotting factors by fresh flowing blood.
- Retard the inflow of clotting factor inhibitors
- Promote endothelial cell injury.



## ○ Causes of Stasis

1. *Atherosclerosis*
2. *Aneurysms*
3. *Myocardial Infarction (Non-contractile fibers)*
4. *Mitral valve stenosis (atrial dilation)*
5. *Hyper viscosity syndrome (PCV and Sickle Cell anemia)*
6. ....





# ○Hypercoagulability

## *A. Genetic (primary):*

- most common >> mutations in factor V gene and prothrombin gene

## *B. Acquired (secondary):*

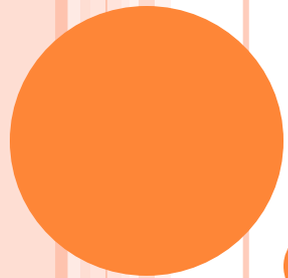
- multifactorial & more complicated
- causes include: Immobilization, MI, AF, surgery, fractures, burns, Cancer, Prosthetic cardiac valves  
...etc



# MORPHOLOGY OF THROMBI

- Can develop anywhere in the CVS (e.g., in cardiac chambers, valves, arteries, veins, or capillaries).
- Arterial or cardiac thrombi → begin at sites of endothelial injury or turbulence; and are usually superimposed on an atherosclerotic plaque
- Venous thrombi → occur at sites of stasis. Most commonly the veins of the lower extremities (90%)
- Thrombi are focally attached to the underlying vascular surface.
- The propagating portion of a thrombus is poorly attached → fragmentation and embolus formation



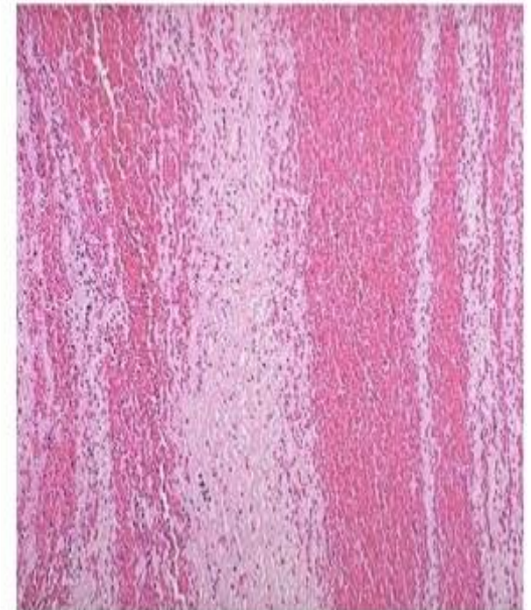
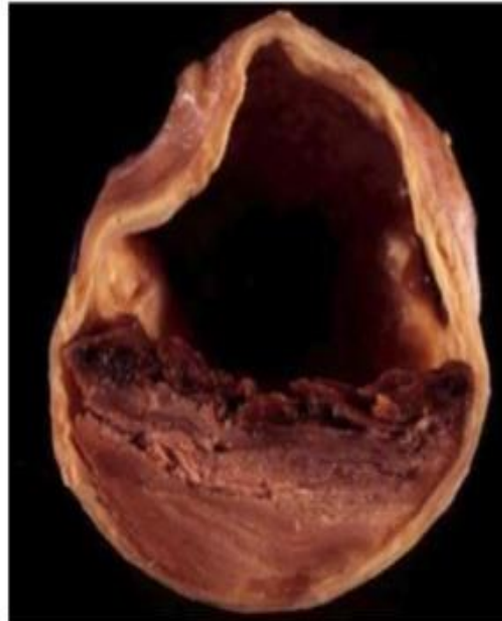


**TERMS TO REMEMBER ...**

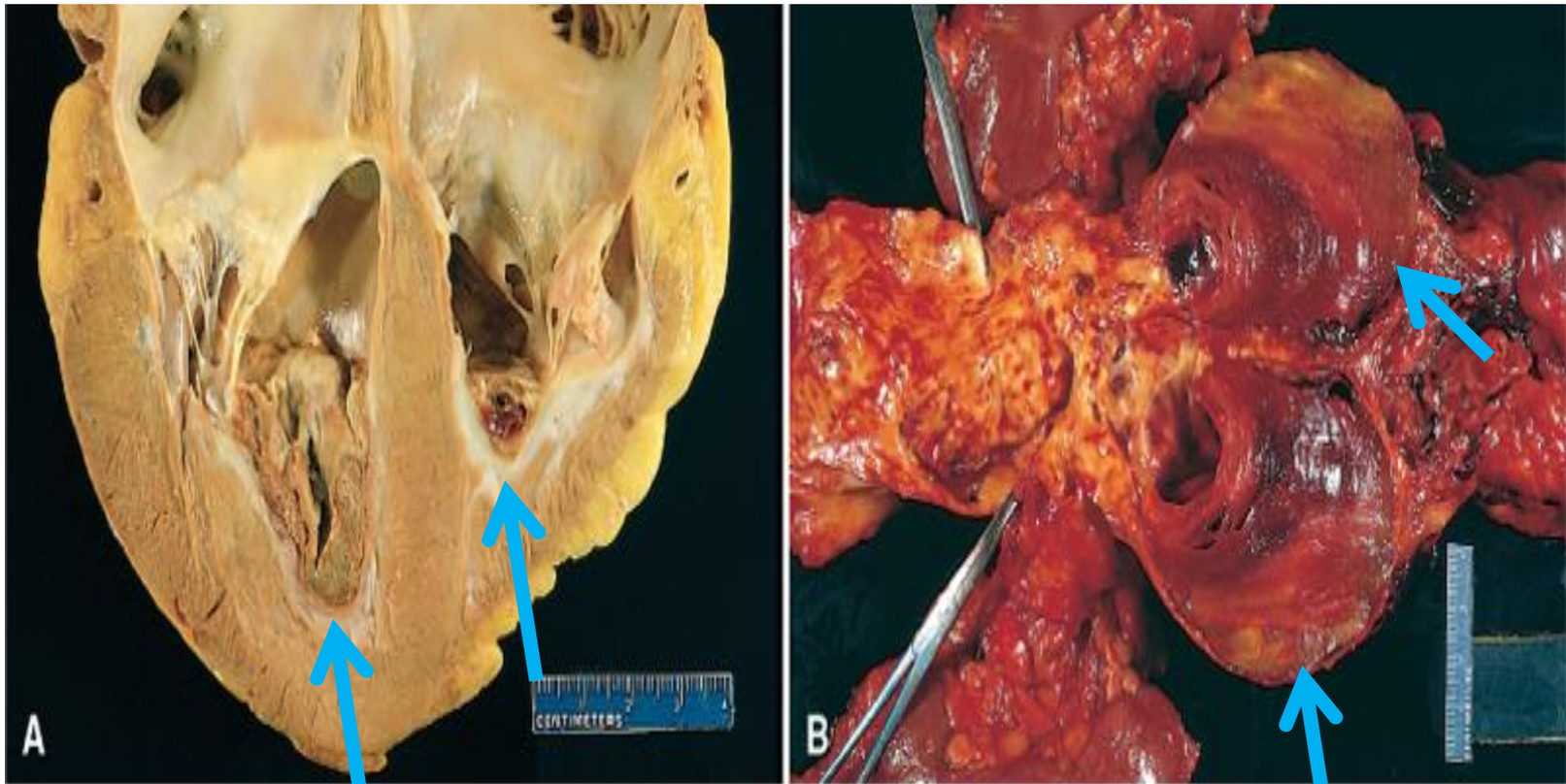


## LINES OF ZAHN

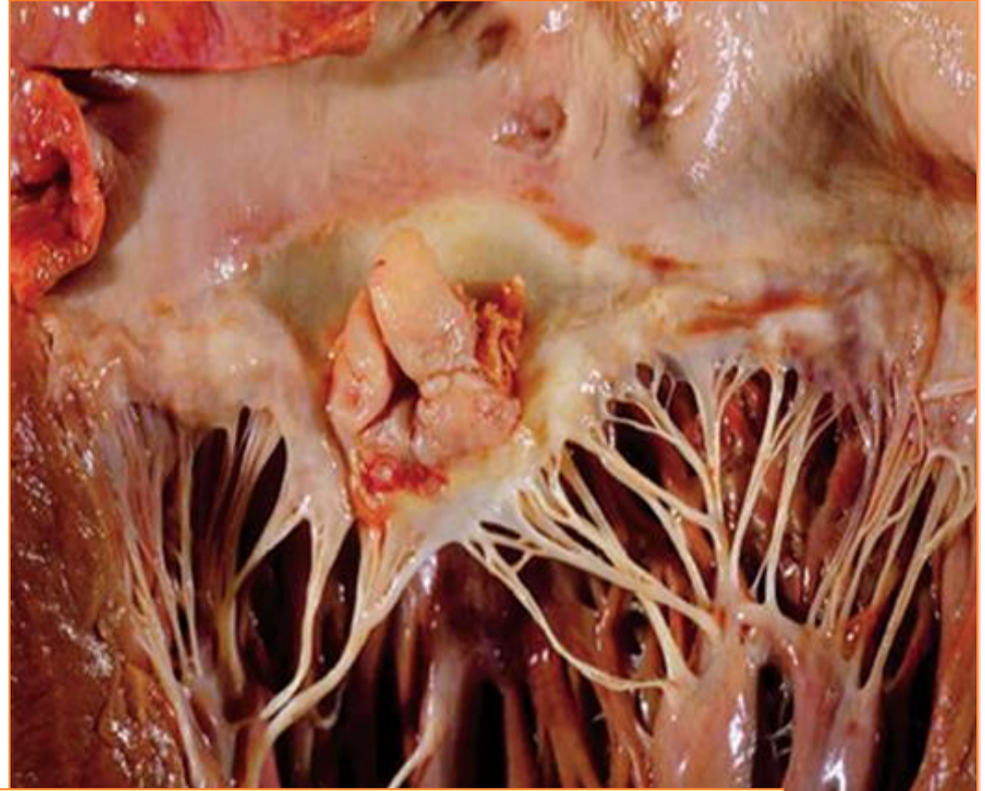
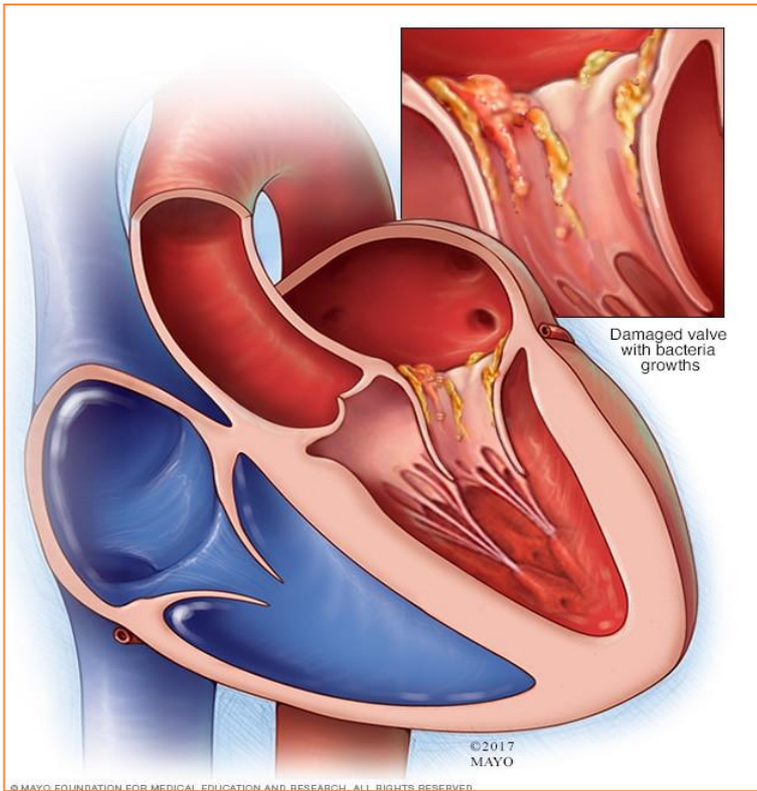
- gross and microscopically apparent laminations
- represent pale platelet and fibrin layers alternating with darker erythrocyte-rich layers
- Significance? distinguish **antemortem** thrombosis from postmortem clots
- postmortem blood clots are non-laminated clots (no lines of Zahn)



# MURAL THROMBI= - IN HEART CHAMBERS OR IN AORTIC LUMEN



# CARDIAC VEGETATIONS



= Thrombi on heart valves

## Types:

1- infectious (Bacterial or fungal blood-borne infections)

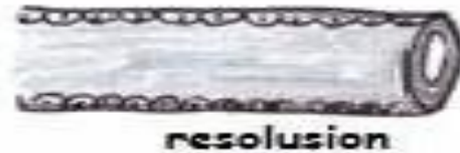
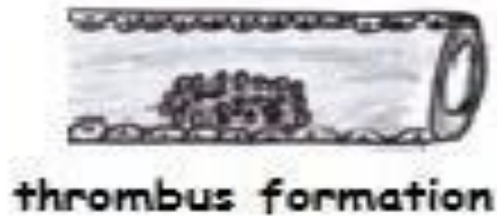
e.g. infective endocarditis

2-non- infectious:

e.g. non-bacterial thrombotic endocarditis

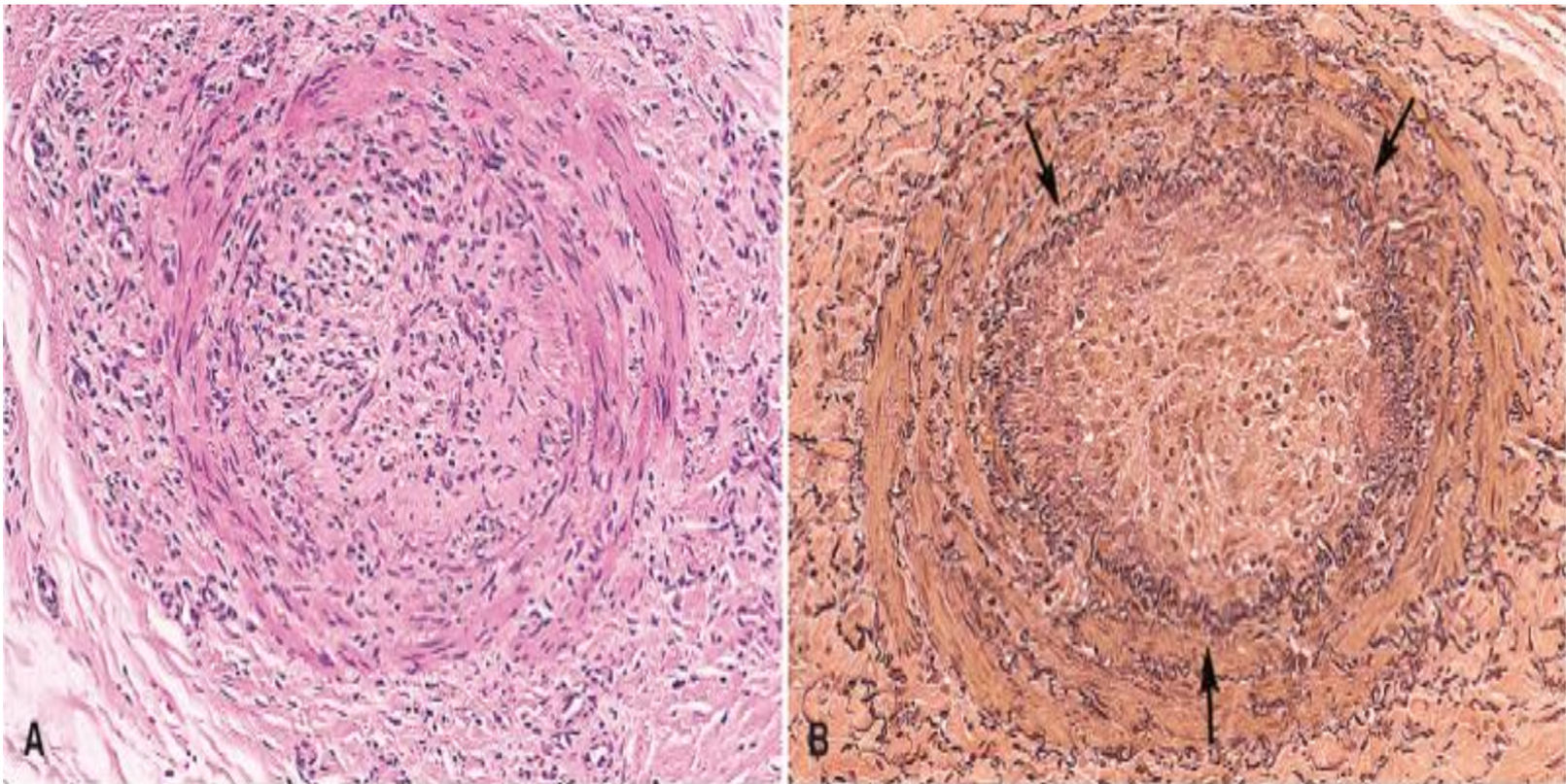


# FATES OF A THROMBUS



Mycotic aneurysm (discussed later)

# ORGANIZED ARTERIAL THROMBUS



## ○ Fate of thrombi

1. **Propagation** → accumulate additional platelets and fibrin, eventually causing **vessel obstruction**
2. **Embolization** → Thrombi dislodge or fragment and are transported elsewhere in the vasculature
3. **Dissolution** → Thrombi are removed by fibrinolytic activity (only in recent thrombi)
4. **Organization\* and recanalization** → Thrombi induce inflammation and fibrosis. These can *recanalize* (re-establishing some degree of flow), or they can be incorporated into a thickened vessel wall

*\*Organization refers to the ingrowth of endothelial cells, smooth cells and fibroblasts into the fibrin rich thrombus.*

5. **Superimposed infection (Mycotic aneurysm)**

