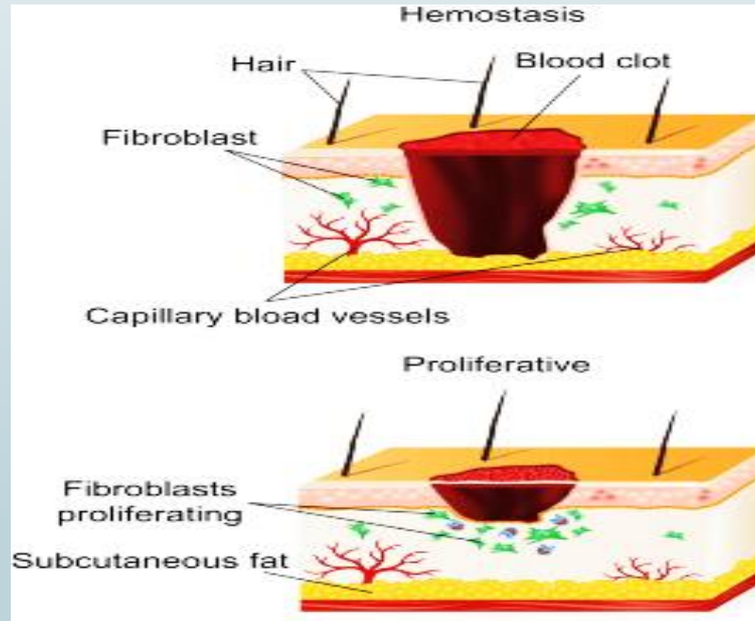




9. BLOOD LYSIS.



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*Causes of **fluidity of blood** inside the cardiovascular system [Factors against intravascular clotting]:

These factors prevent blood clotting in normal state and in cases of injury they limit the process of blood coagulation to the site of injury and help re-canalisation of thrombosed blood vessels:

1) Role of smoothness of endothelium in prevention of clotting:

a- Prevent contact activation of XII.

b- Protein that covers the endothelium has –ve charges which repels –ve charged platelet & clotting factors.

c- Protein (**thrombomodulin**) which cover endothelium binds with thrombin preventing its action and this complex activates protein- C that act as an anticoagulant.

d- Release of prostacyclin (major inhibitor of platelets aggregation) from the healthy endothelium.

2) Anticoagulant in the blood itself:

a- Blood flow: Removal of activated coagulation factors by the circulating blood and their inactivation in the liver, spleen and the bone marrow.

b- Antithrombin action of fibrin and antithrombin III:

Fibrin: adsorb about 90% of thrombin formed during this process preventing its spread into the blood causing more coagulation.

Antithrombin III (alpha globulin) Combine and inhibit the remaining thrombin and factor Xa.

c- **Protein C & protein S:** (Both are natural anticoagulants are formed in the liver in presence of vit.K)

Protein C: activated by thrombin and inhibits the clotting factors V and VIII and stimulate fibrinolysis.

Protein S: potentiate the effect of protein C.

d-Heparin :

- It is the most powerful anticoagulant
- It is negatively charged muco-polysaccharide.
- It is secreted by mast cells and basophile cells in minute amounts.

- Mechanism of its action:

1. It combines with anti-thrombin III aiding its inhibition of thrombin Also, it inhibits the activated factors IX, X and XI
 - 2.It inhibits platelet aggregation and stimulate fibrinolysis.
 - 3.Lipaemia clearing effect occur by activation of lipase enzyme to hydrolyse lipids and prevent its deposition in blood vessels so prevent the development of atherosclerosis.
- e- Alpha2-macroglobulin: bind and inhibit coagulation factors.
 - f- Alpha1-antitrypsin: Inactivate factor XIa and thrombin.
 - g-Plasmin (fibrinolysin):causes breakdown of fibrin, fibrinogen, prothrombin, factor V ,VIII and XII.

*Prevention of blood clotting **outside** the body [**Invitro anticoagulants**]

- (1) Blood is collected in silicon or paraffin coated test tube to prevent aggregation and activation of factor XII.
- (2) Cooling of the blood delay clotting.
- (3) Removal of Ca^{++} ions: by
 - Precipitation of ionized calcium by addition of Na oxalate \square Ca^{++} oxalate (toxic) or by EDTA.
 - Adding of Na citrate \rightarrow Chelation of Ca^{++} and formation of non-ionized Ca^{++} (Ca^{++} citrate). This compound is not toxic, and rapidly removed from the blood so citrate is used in blood transfusion.
 - EDTA (Ethylene diamine tetraacetic acid) \rightarrow Chelation of Ca^{++}
- (4) De-fibrination of blood by a glass rod.
- (5) Addition of heparin as in artificial kidney machine.

*Prevention of blood clotting **inside** the body [**Invivo anticoagulants**]

Drugs are used for prevention and treatment of thrombosis as in:

- 1- Deep venous thrombosis or pulmonary thrombosis.
- 2- Myocardial infarction.
- 3- After cardiac surgery.
- 4- Rheumatic valve disease complicated with embolism.
- 5- In hereditary deficiency of anti-thrombin III, protein C or S.

- There are two types of anticoagulants drugs:

	<i>Heparin</i>	<i>Coumarin</i>
<u>-Origin:</u>	- Animal origin from mast cells and basophils	- Plant origin as warfarin and Dicumarol
<u>-Mode of action:</u>	- Anti-thrombin - Inhibits platelet aggregation - Prevent activation of IX, X, XI - Lipaemia clearing effect	- Competitive inhibition with vit K in liver. So prevent formation of factors II, VII, IX & X and protein C & S.
<u>-Site of action</u>	-In vivo and in vitro	-In vivo only
<u>-Onset:</u>	-Rapid	-Delayed onset (1-3 days)
<u>-Duration:</u>	-Short duration (4-6 h.) then hydrolysed by Heparinase enzyme.	-Long duration (3 days)
<u>-Mode of administration:</u>	-Intravenous or intra-muscular (as it is digested by the stomach)	-Orally
<u>-Antidote:</u>	-Protamine sulphate 1% (It has strong positive charges to neutralize the negative charges of heparin)	-Vitamin K or blood transfusion

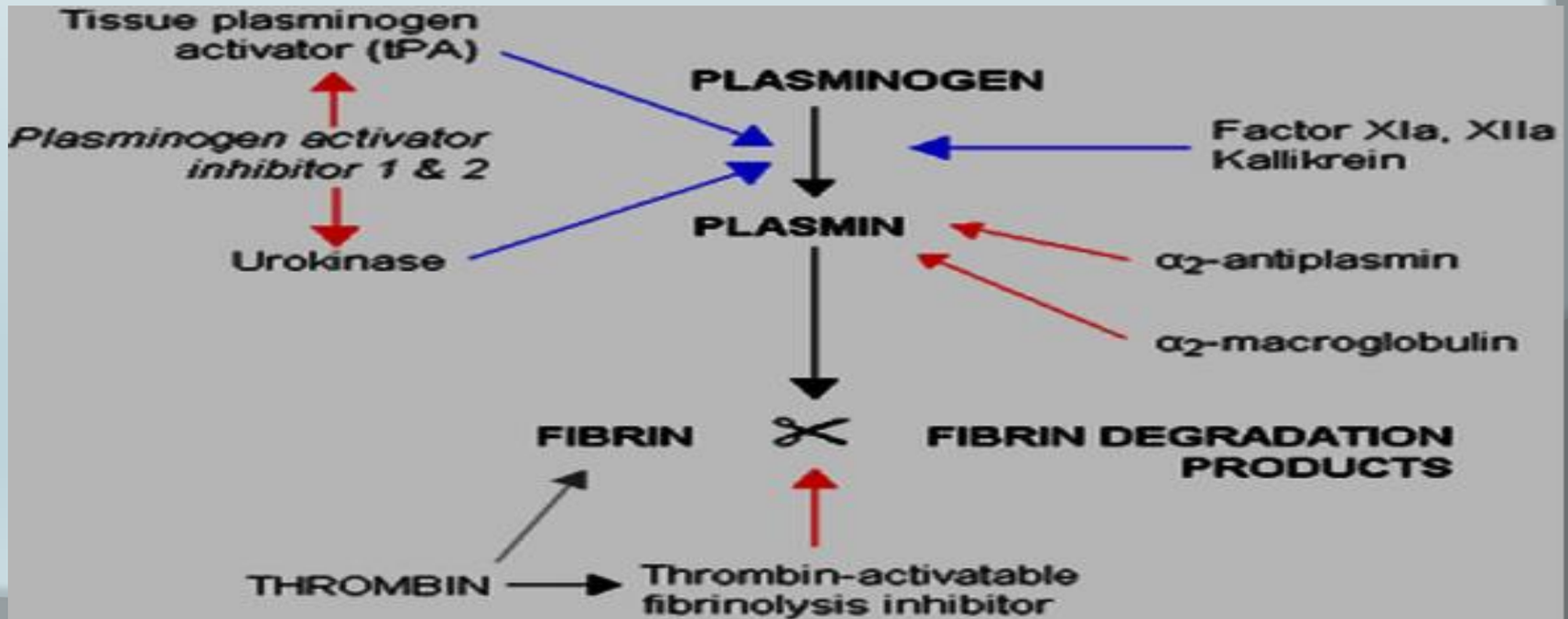
Fibrinolytic System

* Definition:

- Fibrinolysis means lysis and removal of blood clot after stoppage of bleeding and healing of the vascular wall.
- This is produced by enzyme called plasmin (fibrinolysin) which present in plasma as inactive plasminogen (pro-fibrinolysin).

* Mechanism:

- After the blood clotting stops the bleeding, tissue plasminogen activator (t-PA) converts plasminogen into plasmin which lyses the blood clot into fibrin degradation products (FDP).
- After lysis of the blood clot, plasmin, t-PA and FDP are removed by the phagocytic cells.
- Then the inhibitor to t-PA limit its effect to site of blood clot only.



***Activation of plasminogen & fibrinolysis:**

- (1) Tissue plasminogen activator: (t-PA): Released from injured tissue & endothelium but the plasma contains a physiological inhibitor to the t-PA to balance its effect.
- (2) Factor XII, Kallikrein & thrombin.
- (3) Other physiological activators as:
 - a- Urokinase enzyme in the urine to lyse blood clots in the urine.
 - b-Enzymes in pleural, peritoneal & uterine cavities to prevent blood clot in these sites and passage of uterine blood to outside.
- (4) Exogenous activators: as streptokinase enzyme from bacteria to treat acute myocardial infarction to dissolve clot.

***Inhibition of fibrinolysis:**

- (1) Inhibition of plasmin: by α 2 Anti-plasmin , α -2-macroglobulin & α -1 anti-trypsin.
- (2) Inhibition of tissue plasminogen activator.

*** Significance of fibrinolysis:**

- 1- Lysis of blood clots & reopening the blood vessels and prevent closure of capillaries by sluggish circulation.
- 2- Cleaning of the tissue from the blood clots formed outside the blood vessels
- 3- Removal & prevent bl. clots in the urinary tract (to prevent blocking of renal tubules), pleural, uterine & peritoneal cavities.
- 4-Treatment of early stages of myocardial infarction by:
 - Injection of tissue plasminogen activator.
 - Streptokinase & urokinase injection [direct on the clot].

Thank You