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Stages of H	lemostasis
1. Vessel Spasm	Response to inflammation. Initiated by endothelial injury. Reflex vessel restriction by the smooth muscle layer reducing blood flow. Only last 1 minute. Thromboxane A2 released from platelets contribute to vasoconstriction. This happens locally at the site of the injury
2. Formation of Platelet Plug	Platelets are attracted to damaged vessel wall by the release of von willebrand factor. Once they encounter vWF they activate and change from disk shaped to star shaped then flat sphere like shaped. Then they adhere to collagen and aggregation occurs. Aggregation is mediated by the release of granules- ADP and TXA2 (more of these = more aggregation). Glycoprotein IIb and IIIa receptors bind fibrinogen and link platelets together. This leads to the platelet plus formation.
3. Blood Coagul- ation	Results in conversion of inactive soluble fibrinogen to insoluble fibrin. <i>Vitamin K</i> is necessary for synthesis of factors VII , IX , X , prothrombin , protein C . <i>Calcium</i> is required by activated factor X to convert prothrombin to thrombin. Involves intrinsic, extrinsic and common pathway. Regulated by natural anticoagulants (Antithrombin II, Protein C-inactivates factor V and VIII, plasmin-breaks down fibrin).
4. Clot Retraction	Serum is squeezed out of the clot and the edges of the vessels are joined. Failure of clot retraction is indicative of low platelet count.
5. Clot Dissol- ution	Needed for permanent tissue repair. Process known as fibrinolysis (getting rid of clot). Plasmin digests fibrin, factors V, VIII, XII, prothrombin. Plasminogen is activated to plasmin by enzymes (one is factor XII or Hageman factor).

These 5 stages are the holy grail of wound healing.

vWF is produced by endothelial cells, platelets and connective tissue. disorder?

Hemostasis		Blood Testing for Coagulability		Increased Clotting Activity (cont)		Bleeding Disorders (cont)	
Definition:	The process which causes the bleeding to stop. Maintains blood fluidity and prevents blood from leaving the vascular compartments	(cont) PTT- Partial thromb opl- astic time	Tests <i>intrinsic</i> pathway. Used to monitor <i>heparin</i> . Normal range is 30-50 seconds.	Secondary:	Acquired. Stasis due to bed rest (slows normal blood flow and allows accumu- lation of clotting factors)cancer, birth control,	Thromb- ocytop- enia:	Low circulating platelets. Due to decreased production by bone marrow (aplastic anemia, leukemia, HIV) or increased pooling
Main Factors:	1. Cell membrane 2. Platelets 3. Coagulation	Hypercoa platelet f	agulability (increased unction)	Antiphosp-	smoking and obesity, MI. AKA Hughes		of platelets in the spleen, or decreased platelet
	cascade unction of : thrombosis (inapp-	platelet a	agulability results in Idhesion and formation which leads to disruption	holipid Syndrome:	syndrome. Autoimmune hypercoagulable state caused by		survival or nutrit- ional deficiencies (B12, iron, folic acid),
	tting) or bleeding/- ng *insufficient n Cascade		d Clotting Activity		antiphospholipid antibodies. Provokes blood clots in arteries in veins. Can be		Types: idiopathic, thrombotic or hemolytic uremia syndromes or heparin induced.
					primary or secondary (due to lupus).	Decreased platelet function:	Caused by asprin, uremia (increased urea in blood coats the platelets
				Bleeding Dis	orders		causing glycop- roteins not to function) or genetic disorders



Genetics. Mutations in factor V and prothrombin genes. Results in inability of factor V a to be deactivated by protein C. Examples: Factor V Leiden disorder where clotting persists and predisposes to DVT. Other disorders are inherited deficiencies of antithrombin III, protein C/S.

Plateletnormal range:Disorders:150,000- 400,00-0/ml. Signs of0/ml.disorders include:Petechia, purpura,ecchmyosis,bleeding frommucous membrane	
ecchmyosis, bleeding from	 150,000- 400,00- 0/ml. Signs of
	ecchmyosis, bleeding from

Blood Testing for Coagulability

PT-	Tests extrinsic and
Prothr	common pathway.
ombin	Looking at time to clot.
time	Used to monitor
	warfarin. Normal is 11-
	13 seconds. PT is
	increased with warfarin

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Bleeding Di	sorders (cont)	Bleeding Diso	orders (cont)	Anticoagula	ants (cont)	Novel Oral A	nticoagulants (cont)
	Genetic disorders: Bernard Soulier- GpIIb disorder so vWf has nowhere to bind, Von Willebrand Disease-no vWF to bind platelets.	ency: X-linke disorder, affe Soft tissue bl knee, elbow a Can lead to jo contractures. replacement	cts mostly males. eeding of GI, hip, and ankle joints. oint fibrosis and Tx is factor VIII therapy. Only	Hepari- n(IV)/ LMW Heparin (lovenox):	Induces a confor- mational change in antithrombin III making it more accessible to proteases -> increase inacti- vation of thrombin		<i>Uses:</i> A- reduces stroke and systemic embolism, prophy- laxis of DVT/PE after hip or knee surgery. R- same but prophylaxis of
	Leads to decreased platelet adhesion *Vasopressin can stimulate release of	Anticoagulant			Uses: Prophylaxis and tx of thromb- oembolic diseases, unfractionated (IV		venous thromb- oembolic events for hip/knee surgery pts.
	vWF for tx. Glanzmann thrombocytopenia- GpIIb-IIIa so	(Coumadin):	antagonist. Blocks epoxidase reductase, leads to depletion of		heparin) used with antiplatelet agents for tx of acute coronary		<i>AE:</i> easy bruising, bleeding, back or muscle pain, hypotension.
Coagul- ation Cascade	platelets cant bindtogetherDeficiencies orimpairments of oneor more coagul-		reduced vit K (which is essential for synthese of factors II,		syndromes. Lovenox is an efficient cataly- zation of factor Xa inactivation.	Dabigatran (Pradaxa):	Direct thrombin inhibitor which prevents conversion of fibrinogen to fibrin.
Disorders:	ation factors due to defective synthesis, inherited disease or increased consum- ption. Prevents		VII,IX,X, protein c/s) Uses: Prevention of thrombosis in	Neural Orral	AE: bleeding and heparin induced thrombocytopenia		<i>Uses</i> : Prevents thromboembolism in pts with AF, DVT, PE
	fibrinogen from converting to fibrin. Will see bleeding in deep tissues like		predisposed patients. AE- bleeding	Apixaban (Eliquis), Rivaro-	Anticoagulants Direct inhibitor of free and clot-bound factor Xa which	Betrixaban (Bevyxxa):	Cofactor-indepe- ndent direct inhibitor of factor Xa.
	hematomas. Elevated PTT and PT.			xaban (Xarelto):	prevents the conversion of prothrombin to thrombin. Prevents clot formation.		<i>Uses</i> : prophylaxis of VTE in moderate to severe restricted mobility patients.

Rivaroxaban interacts with Aspirin. All drugs will have bleeding as a side effect!

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Inhibition of	Anticoagulation	Antiplatelet	Agents	Antiplatel	et Agents (cont)	Antiplatelet	Agents (cont)
Protamine:	Antagonist of heparin. <i>Uses</i> : IV administr- ation if there is life threatening hemorrhage/h- eparin excess	Aspirin (ASA)	Non selective COX inhibitor. Irreve- rsible inhibition of COX-1= inhibits platelet aggregation for 10 days. Stops conversion of		<i>Uses</i> : Intermittent claudication symptoms (by widening the vessels in teh legs which helps with blood flow).		<i>Uses</i> : reduces risk of MI/stroke, bette than aspirin in decreasing CV outcomes <i>AE, DI</i> : upper RTI joint, chest pain,
Thrombolytic Agents Strept- Forms a stable			arachidonic acid to thromboxane A2 (potent platelet		<i>AE, DI</i> : heart failure, tachycardia, interacts with		depression, bleeding. DI- Ibuprofen
okinase:	complex with		aggregation inducer). <i>Uses</i> : Pain/infl- ammation/fever, reduces risk of MI/unstable angina, prevents strokes due to blood clots		NSAIDs and aspirin.	GPIIb-IIIa A	ntagonist
	plasminogen which then cleaves other plasminogen molecules into plasmin			Pentox- ifylline (Trental)	Inhibits erythrocyte phosphodiesterase - > increases cAMP activity, decreases blood viscosity by reducing plasma fibrinogen concentra- tions and increasing fibrinolytic activity	Abicixmab (Reopro)	Binds to intact platelet GPIIb/IIIa receptor and blocks access of large molecules to receptor through steric hinderance or conformational change. Prevents cell adhesion <i>Uses</i> : prevents cardiac ischemic complications in vascular surgeries or pts w/ unstable angina, intended
	<i>Uses</i> : PE, STEMI, arterial thrombosis, DVT. <i>AE</i> :systemic fibrinolysis,						
			<i>AE</i> : hemorrhagic stroke, GI bleeding				
	hemorrhage	PDE Inhibi	ors		Uses: Intermittent		
Recomb- inant Tissue	formed thrombi and (Pl makes it a potent activator of plasmi- nogen. Cleaves	Cilostazol (Pletal)	Antiplatelet and vasodilator. Inhibitors phosph- odiesterase II -> suppresses cAMP degradation -> increases cAMP in platelets and blood vessels -> inhibition of platelet aggreg- ation and vasodi- lation		claudication, chronic occlusive arterial disease		
Plasmi- nogen Activator:					<i>AE</i> : muscle aches, headaches, GI discomfort		
				ADP Receptor Pathway Inhibitor			for use with aspirin and heparin
	fibrin degradation products			Clopid- ogrel (Plavix)	Irreversibly binds to P2Y12 which prevents the binding		AE: N&V, hypote- nsion, vision
	<i>Uses</i> : PE,STEMI, Acute ischemia			、 /	of ADP receptors on platelets which		changes, back pain
	stroke. AE: bleeding				prevents GPIIb-IIIa activation -> inhibits platelets aggregation		

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