

Class	Drugs/Dose	MOA	ADME	ADRs/CI
Cyclooxygenase (COX) Inhibitor [NSAID]	<p>ASA (e.g. Aspirin)</p> <p>Low dose ASA 81mg/d: used for prevention of heart attacks</p> <p>Dose: 75-150mg/day</p>	<ul style="list-style-type: none"> Low dose = more effective in inhibiting platelet COX (irreversible acetylation) and does not affect COX1 in BV (in EC promote prostacyclin = VD, prevent clumping/activation, reversible in EC b/c has a nucleus) Irreversible acetylation – once blocked stays for the lifetime of platelet (7-10 days); platelets have no nucleus Inhibits platelet COX activity by blocking TXA2 formation = abolishes ability of platelets to release TXA2 = ↓ overall platelet aggregation In vascular endothelial cells – also contain COX activated by PGI2 synthesis – but forms prostacyclin → causes VD 	<ul style="list-style-type: none"> Onset of antiplatelet effect <60mins (rapid) Good efficacy Inexpensive Complete inactivation of platelet COX w/ 160mg/daily Duration: 7-10 days b/c irreversible 	<ul style="list-style-type: none"> Prolongs bleeding time GI bleed [in digestive tract COX1 makes PGs that help protect cells lining stomach against acids/digestive enz] Hemorrhage stroke bleeding within brain (rare) <p>Weakest antiplatelet agent (only blocks one pathway in aggregation)</p>
ADP-inhibitors	<p>Clopidogrel (P) 1st gen</p> <p>Dose: 300mg LD (need for immed. anti-aggreg. effect) → 75mg OD for LT without food</p>	<ul style="list-style-type: none"> selectively (and irreversibly) – binding of ADP to its platelet receptor (P₂Y₁₂) – activation of the GP IIb/IIIa R – TXA2 generation = – platelet aggregation <ul style="list-style-type: none"> prevent ↑ Ca+ AND GPIIb/IIIa R from binding = prevent sustained aggregation 	<ul style="list-style-type: none"> Prodrug (P) Esterase covert 85% into inactivate metabolites – only 15% active 	<ul style="list-style-type: none"> Drug interactions – CYP3A4, 3A5, 2C9 Interpatient variability – resistance (due to genetic polymorphs of CYP450 [↓ 30% in active clopidogrel – cant convert prodrug to active metabolite]) Rash (back, chest, abd) ~4% Major bleeding
	<p>Prasugrel (P) 1st gen</p> <p>Dose: 60mg LD → 10mg OD</p>		<ul style="list-style-type: none"> Prodrug (P) – intestinal (esterase) and hepatic (CYP450) – require gut and liver 10x more potent > clopidogrel Rapid action Can be taken together with ASA ✓ 	<ul style="list-style-type: none"> Increased risk of bleeding (higher risk vs Clopidogrel – b/c more efficacious) Less sensitive to polymorphisms and drug interactions (vs Clopidogrel) – CYP 3A4, CYP2B6, CYP2C9
	<p>Ticagrelor 2nd gen</p> <p>Dose: 180mg LD → 90mg BID</p>		<ul style="list-style-type: none"> Direct and reversible P₂Y₁₂ antagonists – its changes the conformation of the receptor Does not require metabolic activation for its antiplatelet effects (not a prodrug) 	<ul style="list-style-type: none"> More rapid onset Metabolized by CYP 3A4/5 Can be taken together with ASA ✓
GPIIb/GPIIIa	<p>Eptifibatid</p> <p>Dose: 180ug/kg IV STAT then continuous infusion 2ug/kg/min until hospital d/c or initiation of CABG up to 72 hrs</p>	<ul style="list-style-type: none"> Reversible, small molec, synthetic <u>peptide</u>, resembles RGD sequence of fibrinogen 	<ul style="list-style-type: none"> MW: 832 Da RoA: IV Short T½: 2-4h Restricted use b/c IV 	<ul style="list-style-type: none"> Severe bleeding – can stop drug and wait for platelet fn to return to normal as drug is cleared
	<p>Tirofiban</p> <p>Dose: infusion rate of 0.4 ug/kg/min for 30 min followed by maintenance infusion rate of 0.1 ug/kg/min for at least 48 hrs</p>	<ul style="list-style-type: none"> Reversible, small molec, synthetic <u>non-peptide</u> antagonist (inhibitor) of fibrinogen binding to the GPIIb/IIIa receptor (RGD mimetic) <p>Both: MoA: reversibly blocks GP IIb/IIIa receptors → prevent fibrinogen connecting platelets → doesn't make fibrin = block platelets from aggregating</p>	<ul style="list-style-type: none"> Smaller than Eptifibatide MW: 495Da RoA: IV Rapid onset Short duration (4 hrs) 	<ul style="list-style-type: none"> Severe bleeding
Organic Nitrate/Nitrates	<ul style="list-style-type: none"> Nitroglycerin (P) Isosorbide dinitrate Isosorbide 5-mononitrate 	<ul style="list-style-type: none"> Used for both tx of acute attacks of angina and for prophylaxis Reduces amount of blood entering into the heart – main effect is VD in veins ALL need to convert into NO via mtALDH2 Main effect: relaxation of smooth muscles of veins – affects arteries too but main is in the veins = ↓ preload = ↓ EDV = ↓ SV Coronary perfusion happens when heart relaxes (at diastole: backflow & closes aorta valve – fluid can only go in 1 direction into coronary BV) 3 Effects: <ol style="list-style-type: none"> Venous VD*main effect = ↓ preload = ↓ ventricular size, ↓ vent. wall stress (↓ pressure of branches of CBVs and allow flow) Direct coronary VD – myocardial perfusion Arterial VD – ↓ afterload = ↓ BP [b/c VD in arteries = ↓ TPR] 	<ul style="list-style-type: none"> Prodrugs (P) – require enz processing to form NO Poor oral bioavailability (10-20%) due to first pass liver metab Dosage forms: SL, PO, patch, spray 	<ul style="list-style-type: none"> Flushing of face/neck (due to dilation of arterioles in face/neck) Reflex tachycardia [b/c VD of arteriole = ↓ BP = baroreceptor starts firing = ↑ HR] Repeated doses can lead to tolerance (b/c mtALDH2 gets all used up – need to let it recover – about 8hr drug free intervals) Throbbing headache Interactions w/ PDES inhibitors (e.g. sildenafil, tadalafil) – b/c ↑↑ cGMP
Direct	<ul style="list-style-type: none"> Sodium Nitroprusside 	<ul style="list-style-type: none"> Release NO in circulation Relax both arteries and veins – b/c both have smooth muscles Tx of HTN crisis [v high BP] DIRECT – no need to be converted 	<ul style="list-style-type: none"> *sensitive to light 	<ul style="list-style-type: none"> Cyanide poisoning: SNP → cyanide → thiocyanate) *add Na thiosulfate into infusion to change into inactive thiocyanate
HCN Channel Blocker	Ivabradine	<ul style="list-style-type: none"> Pure heart rate reduction Blocks HCN channel in SA node: prevents Na+ from entering into slow Na+ channel (aka HCN channel) by: ↓ slow Na influx into HCN channel → delayed depolarization → ↓HR → more time for heart to be relaxed (b/c main amount of flow is during diastole) 		
Thrombolytic/fibrinolytic	Alteplase (Activase)	<ul style="list-style-type: none"> Clot specific – active only at site of clot tPA only active in presence of fibrin (fibrin specificity) – converts plasminogen into plasmin → fibrin mesh digested ↑ plasmin = greater degradation 	<ul style="list-style-type: none"> Short T ½ = 3-4 mins 	
	tPA = tissue plasminogen activator		<ul style="list-style-type: none"> Short T ½ = 14-15 mins Both: only given in hospital, IV 	