

Anti-arrhythmic drugs

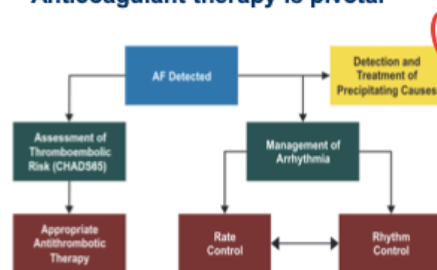
Class	Drugs/Dose	MOA	Indication	ADRs/CI
Class I Anti-arrhythmia: sodium channel blockers	1A: Quinidine , procainamide, ajmaline, prajmalium	<ul style="list-style-type: none"> Moderate sodium channel block, inhibition of potassium channels (prolonged repolarization) 	<ul style="list-style-type: none"> Atrial flutter & fibrillation Ventricular tachycardia Extrasystoles Quinidine = BRUGADA'S syndrome (congenital defects) 	<p>S/E: hypotension, negative inotropic effect (HF), blood count, arrhythmia (prolongation of AP and QT-interval prolongation), overdosing = AV block</p> <p>C/I: bradycardia, DHF, conduction defects, 1st trimester pregnancy</p> <p>Quinidine S/E: in addition to above = GI (NVD), overdosing = cinchonism (headache, dizziness, tinnitus)</p>
	1B: Lidocaine, mexiletine	<ul style="list-style-type: none"> Mild sodium channel block, shortened repolarization LIDOCAINE: Mainly ventricular effects; decreased depolarization velocity especially in case of decreased resting potential; increase recovery time of sodium channels with high activation frequencies (USE DEPENDENCY); High first pass effect (IV) MEXILETINE: PO admin, 2-3x/day (600-1200mg/d) (neuropathy dose: 450-750mg/d) 	<p>LIDOCAINE</p> <ul style="list-style-type: none"> Acute therapy of ventricular tachycardia Prevention of ventricular fibrillation after heart attack Least cardiotoxic Na blocker BUT NO PROPHYLACTIC USE <p>MEXILETINE</p> <ul style="list-style-type: none"> Ventricular arrhythmia Neuropathic pain 	<p>LIDOCAINE S/E: hypotension, negative inotropic effects (HF), overdosing = seizures</p> <p>LIDOCAINE C/I: AV block grade II and III</p> <p>Mexiletine S/E: dose related, neurologic effects include tremor, blurred vision, lethargy and nausea</p>
	1C: Flecainide, Propafenone	<ul style="list-style-type: none"> Marked sodium channel block, no effect on repolarization (slow dissociation from sodium channels → inhibits fast sodium influx during phase 0 strongly → QRS prolongation but no QT interval or APD prolongation) 	<ul style="list-style-type: none"> Limited, supraventricular arrhythmia Life threatening ventricular arrhythmias Some congenital arrhythmia syndromes (type 3 LQTS and RYR2 mutations) 	<p>PROPAFENONE: CYP2D6 SUBSTRATE (SLOW METABOLIZER)</p> <p>S/E distinct pro-arrhythmic potential (even at normal doses), blurred vision (F), exacerbation of congestive heart failure</p> <p>Flecainide S/E: increased mortality for patients after myocardial infarction</p>
Class II anti arrhythmia: sympatholytic drugs (beta blockers)	Propranolol	<ul style="list-style-type: none"> 1st line therapy (safety and efficacy has been proven) = reduces mortality after MI Negative inotropic, chronotropic and dromotropic effects, membrane stabilizing effects (decreases heart rate, decreases intracellular calcium overload, inhibition of after-depolarization automaticity) 	<ul style="list-style-type: none"> Sinus tachycardia Supraventricular arrhythmia Ventricular extrasystoles Atrial fibrillation 	<p>Negative inotropic, AV block, bronchospasm, fatigue, hypotension, aggravation of HF</p>
	Esmolol			
	Sotalol			
Class III anti arrhythmias: prolongation of refractory period	Inhibition of potassium channels → prolongation of repolarization and APD → reverse use-dependency (= high affinity to channels in normal physiological state so sotalol is able to prevent the development of arrhythmias whereas drugs like lidocaine are only effective when there is already arrhythmias i.e., has to be high fire rate)			Torsades de pointes arrhythmia (strong substance specificity)
	Sotalol	<ul style="list-style-type: none"> Beta and potassium channel blocker 	<ul style="list-style-type: none"> Life threatening ventricular arrhythmias Maintenance of sinus rhythm in patients with atrial fibrillation Also used in pediatrics 	
	Amiodarone	<ul style="list-style-type: none"> Multi-channel inhibitor (dirty drug) = blocks inactivated sodium channels, decreases calcium current and transient outward delayed rectifier and inward rectifier potassium currents, prolongs action potential and refractoriness, delays repolarization 	<ul style="list-style-type: none"> Serious supra and ventricular arrhythmia Effective in otherwise difficult to treat arrhythmia Broadly used (1st line) No increase in mortality in patients with HF or CAD 	<p>S/E: Long t_{1/2} (20 – 100 d), accumulation in several tissues including skin, cornea, photodermatitis (changing eye colour, hyper or hypothyroidism, LUNG FIBROSIS (doses ≥ 400mg/d), bradycardia, AV block</p> <p>Lots of interactions: substrate of cyp3a4, inhibitor of cyp1a2, cyp2d6, cyp3a4, inhibition of p-GP, increased anticoagulation of coumarins</p>

	Dronedaron	<ul style="list-style-type: none"> Structural analog of amiodarone No iodine atoms → reduced lipophilicity → reduced t/12 (25-35h) 	<ul style="list-style-type: none"> Atrial fibrillation Atrial flutter 	S/E: diarrhea, impaired sense of tasting, bradycardia, GI problems, liver toxicity, black box warning against ADHF or NYHA IV HF Less risk of accumulation and S/E (but also less effective) Less interactions with thyroid hormones, cyp3a4 substrate, cyp2d6 inhibitor
	Ibutilide	<ul style="list-style-type: none"> IV admin only 	<ul style="list-style-type: none"> Acute conversion of atrial flutter fibrillation (within 20 minutes) 	Pronounced QT interval prolongation and torsade-de-pointes (→ ECG monitoring for 4 hours after use)
Class IV: calcium channel blockers	Verapamil, diltiazem, and bepridil	<ul style="list-style-type: none"> Block calcium channels in cardiac tissue and thus have anti-arrhythmic effects Verapamil: inhibition of activated and inactivated L-type calcium channels → active in very frequently firing tissue, less completely polarized tissue at rest and in SA and AV node (activation depends exclusively on calcium), prolongation of AV conduction Diltiazem similar to verapamil 	<ul style="list-style-type: none"> Verapamil: extracardiac effects (vasodilatation) Supraventricular arrhythmia (2nd choice after adenosine) Rate control in ventricular flutter or fibrillation 	
Miscellaneous anti-arrhythmia	Adenosine	<ul style="list-style-type: none"> Increased potassium influx, decreased calcium influx Marked hyperpolarization, suppression of calcium dependent AP, shortening of AP Bolus: inhibits AV nodal conduction (less effects on SA) Very short duration of action (10s) Less effective in presence of adenosine receptor blockers (e.g. theophylline, caffeine) 	<ul style="list-style-type: none"> 1st choice for paroxysmal supraventricular tachycardia to sinus rhythm (high efficacy) 	S/E: transient asystole, flushing, shortness of breath, chest burning (≥10% of patients), atrial fibrillation C/I: Grade II and III AV block, atrial fibrillation or flutter, obstructive lung disease
	Cardiac glycosides: Digoxin	<ul style="list-style-type: none"> Reduced intracellular potassium levels → hyperpolarization → decreased conduction velocity Vagotonic actions → inhibition of calcium current in AV node Increased calcium exceeds storage capacity of endoplasmic reticulum → extrasystoles Positive inotropic Digitoxin = enterohepatic circulation (t1/2: 168-192h); digoxin = renal elimination (t1/2: 40h) 	<ul style="list-style-type: none"> Supraventricular tachycardia Atrial fibrillation and flutter Standby medication 	S/E: low therapeutic index, arrhythmia, disturbances of cognitive functions, nausea, blurred vision, hyperkalemia C/I: ventricular fibrillation (NEEDS TO BE CLEAR DIAGNOSIS FOR USE)
	Magnesium	<ul style="list-style-type: none"> 1-2g MgSO4 IV MOA unknown effective in patients with normal magnesium serum levels and low magnesium levels 	<ul style="list-style-type: none"> Prevention of recurrent episodes of torsade-de-pointes arrhythmia Digitalis associated arrhythmia 	S/E: flush, bradycardia, hypotension, impaired AV conduction
	Ivabradine	<ul style="list-style-type: none"> Selective inhibitor of If channels in SA node → decreases heart rate → effect in inappropriate sinus tachycardia but is NOT YET approved 		
	Vernakalant	<ul style="list-style-type: none"> Multic-ion channel blocker, frequency and voltage dependent block of sodium current → mild QT interval prolongation Rapid termination of atrial fibrillation but NOT YET approved 		

Condition	Exclude/use with caution
Heart failure	Flecainide, disopyramide
Sinus or AV node dysfunction	Digoxin, β -blockers, amiodarone, verapamil, diltiazem
History of myocardial infarction	flecainide
Prolonged QT interval	Quinidine, procainamide, sotalol, ibutilide, amiodarone
Cardiac transplant	adenosine

Guidelines for Treatment of Atrial Fibrillation

- Most common sustained arrhythmia = atrial fibrillation
- Prevalence: 0.5% \leq 65 years; 10 % \geq 80 years
- **Treatment goal:** relieve of symptoms, prevention of complications of thromboembolism an tachycardia-induced heart failure
- **Anticoagulant therapy is pivotal**



Major Goals of AF/AFL Arrhythmia Management

- Identify and treat underlying structural heart disease and other predisposing conditions
- Relieve symptoms
- Improve functional capacity/quality of life
- Reduce morbidity/mortality associated with AF/AFL
 - ✓ Prevent tachycardia-induced cardiomyopathy
 - ✓ Reduce/prevent emergency room visits or hospitalizations secondary to AF/AFL
- Prevent stroke or systemic thromboembolism

Excerpt from Guideline from Canadian Society of Cardiology

- Treatment objectives:**
1. ventricular rate control (60-80 bpm; Ca²⁺ blocker and/or β -blocker; 2nd line digoxin in case of heart failure)

2. Sinus rhythm control

Rate control has better benefit-to-risk ratio compared to rhythm control