Antiarrhythmic Drugs

Munir Gharaibeh MD, PhD, MHPE School of Medicine, The University of Jordan November 2017 **Types of Cardiac Arrhythmias Abnormalities of Impulse Formation:** Rate disturbances. **Triggered automaticity. Abnormalities of Impulse Conduction: Blocks**. **Reentry.**



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Causes of Cardiac Arrhythmias Cardiac Causes: Ischemic heart disease. Inflammation. Trauma e.g. heart surgery. Congestive heart failure. Hypotension.

11/27/17

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Causes of Cardiac Arrhythmias

Non Cardiac Causes: Electrolyte imbalance. Acid-Base imbalance. Hypoxia. Drugs: Digitalis, Anesthetics, Tricyclic, **Diuretics**, **Bronchodilators**. G.I. reflexes. Neural reflexes.



Source: Katzung BG, Masters SB, Trevor AJ: *Basic & Clinical Pharmacology,* 11th Edition: http://www.accessmedicine.com Munir Gharaibeh MD, PhD, MHPE

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SA node automaticity

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Normal Circuitry

Purkinje twig



A. Normal conduction

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Re-entry Rhythm



B. Unidirectional block

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Pre-requisites for Reentry (Circus Movement)

Anatomic or physiologic obstacle.

Unidirectional block.

Conduction time around the circuit must be longer than the effective refractory period.



Туре	Chromosome Involved	Defective Gene	lon Channel or Proteins Affected	Result
LQT-1	11	KCNQ1	I _{Ks}	LF
LQT-2	7	KCNH2 (HERG)	Ι _{κr}	LF
LQT-3	3	S CN5 A	I _{Na}	GF
LQT-4	4	Ankyrin-B ¹		LF
LQT-5	21	KCNE1 (minK)	I _{Ks}	LF
LQT-6	21	KCNE2 (MIRP1)	I _{Kr}	LF
LQT-7 ²	17	KCN J2	I _{KIr}	LF
LQT-8 ³	12	CACNA1c	l _{ca}	GF
SQT-1	7	KCNH2	I _{Kr}	GF
SQT-2	11	KCN Q 1	l _{ks}	GF
SQT-3	17	KCN J2	I _{KIr}	GF
CPVT-1 ⁴	1	h Ry R2	Ryanodine receptor	GF
CPVT-2	1	CAS Q2	Calsequestrin	LF
Sick sinus syndrome	15 or 3	HCN4 or SCN5A ⁵		LF
Brugada syndrome	3	S CN5 A	I _{Na}	LF
PCCD	3	S CN5 A	I _{Na}	LF
Familial a trial fibrillation	11	KCN Q1	l _{Ks}	GF

TABLE 14-1 Molecular and genetic basis of some cardiac arrhythmias.

ECG of some Arrhythmias

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Source/ Katzung BG, Masters///#BGMMARMD/RDD MARic & Clinical Pharmacollogy, 11th Edition: http://www.accessmedicine.com

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Torsade de Pointes Polymorphic Ventricular Tachycardia LQT, syncope, and sudden death. Causes: Familial long QT interval Drug - Induced (drugs which prolong APD) <u>Mechanisms:</u> Increased inward current (GF), or **Decreased outward (LF) current during** the plateau. **Genetic Studies:** 300 different mutations in at least 8 ion channel genes. 11/27/17 Munir Gharaibeh MD, PhD, MHPE

Figure 14-8



Source: Katzung BG, Masters SB, Trevor AJ: Basic & Clinical Pharmacology, 12th edition: www.accessmedicine.com

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Electrocardiogram from a patient with the long QT syndrome during two episodes of torsades de pointes. The polymorphic ventricular tachycardia is seen at the start of this tracing and spontaneously halts at the middle of the panel. A single normal sinus beat (NSB) with an extremely prolonged QT interval follows, succeeded immediately by another episode of ventricular tachycardia of the torsades type. The usual symptoms include dizziness or transient loss of consciousness. (Reproduced, with permission, from Basic and Clinical Phartyperdiagy, 10th edition, McGraw-Hill, 2007.) Munir Charalbeh MD, PhD, MHPE 13

Torsade de Pointes

Risk Factors:

- Bradycardia.
- Hypokalemia.
- Triggered upstrokes.
- Drugs which [↑] APD.
- **Treatment:**
 - K+
- ✓ Triggered upstrokes (β Blockers or Mg++)
 ↓ APD (Pacemaker <u>or</u> isoproterenol).

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Other Congenital Arrhythmias Short QT Syndrome:

- GF mutations in three potassium channel genes(KCNH2, KCNQ1, and KCNJ2).

Chatecholaminergic Polymorphic Ventricular Tachycardia (CPVT):

- Stress or emotion-induced syncope.
- Caused by mutations in sarcoplasmic proteins that control calcium.

Other Congenital Arrhythmias Sick Sinus Syndrome: Mutations in HCN4 and SCN5A Brugada Syndrome: - Ventricular fibrillation, persistent ST elevation, and BBB. Linked to LF mutations in SCN5A Familial Atrial Fibrillation: Linked to GF mutation in the potassium channel gene, KCNQ1.

Nonpharmacologic TherapySurgery.

Radiofrequency Catheter Ablation.

Implantable Cardioverter- Defibrillator (ICD)

Gene therapy!!!!.

Mechanism of Action of Antiarrhythmic Drugs

- Readily bind to activated channels or inactivated channels, but bind poorly to rested channels.
 i.e.: Use –Dependent or State-Dependent.
- Channels in normal cells will rapidly lose the drug from the receptors during the resting portion of the cycle.
- This selectivity is lost with increasing doses, leading to drug-induced arrhythmias.

 Also, these drugs may become" *Proarrhythmic or Arrhythmogenic*" during fast heart rates, acidosis, hyperkalemia, or ischemia.



Source: Brunton LL, Lazo JS, Parker KL: Coodman & Gilman's The Pharmacological Basis of Therapeutics, 11th Editory GREED, N.C. Stressmedicine.com

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Possible Effects of Drugs on Action Potential



	\bigcirc	Example	Mechanism of action	Electrophysiological actions	Clinical use	
Vaughan Williams classification	Class la	Disopyramide	Na ⁺ channel block	Reduced rate of depolarisation of action potential, increased ERP, decreased AV conduction	Ventricular fibrillation, especially associated with myocardial	
	Class II	Propranolol, atenolol	β-Adrenoceptor antagonism	Slowed pacemaker activity, increased AV refractory period	Dysrhythmia prevention in myocardial infarction; paroxysmal atrial fibrillation due to sympathetic activity	
	Class III	Amiodarone, sotalol	K⁺ channel block	Increased action potential duration and increased ERP	Atrial fibrillation; ventricular fibrillation	
	Class IV	Verapamil	Ca2+ channel block	Decreased APD, slowed AV conduction	Supraventricular tachycardias; atrial fibrillation	
Not classified by system		Adenosine	K ⁺ channel activation	Slowed pacemaker activity, slowed AV conduction	Given i.v. for supraventricular tachycardias	
	\bigcirc	Digoxin	K ⁺ channel activation (vagal action)	Slowed AV conduction (block)	Atrial fibrillation	
		Magnesium chloride	? Ca ²⁺ channel block		Ventricular fibrillation; digoxin toxicity	

Table 17.1 The mechanism of action, the electrophysiological actions and clinical uses of selected antidysrhythmic drugs

APD, action potential duration; AV, atrioventricular; EMP, effective leftactury period.MEIPE



de, an action potential is triggered when the pacemaker potential reaches a critical threshold (approx. -60 mV).

		Effect on AV	$\overline{\mathbf{D}}$		Usefulness in Arrhythmias			
Drug	Effect on SA Nodal Rate	Nodal Refractory Period	PR Interval	QRS Duration	QT Interval	Supra- ventricular	Ventricular	Half-Life
Adenosine	↓↑	↑↑↑	↑↑↑	0	0	+++++	?	< 10 s
Amiodarone	μĻ,	↑ ↑	Variable	↑	$\uparrow\uparrow\uparrow\uparrow$	+++	++++	(weeks)
Diltiazem	¢↓	↑ ↑	↑	0	0	+++	-	4–8 h
Disopyramide	↑↓ ^{1,2}	1,↓2	†↓²	$\uparrow\uparrow$	$\uparrow\uparrow$	+	+++	7–8 h
Dofetilide	↓(?)	0	0	0	† †	++	None	7 h
Dronedarone					1	+++	-	24 h
Esmolol	$\downarrow\downarrow$	$\uparrow\uparrow$	† †	0	0	+	+	10 min
Flecainide	None,↓	↑ (↑	^† †	0	+3	++++	20 h
lbutilide	↓ (?)	0	0	0	$\uparrow\uparrow$	++	?	6 h
Lidocaine	None ¹	None	0	0	0	None ⁴	+++	1–2 h
Mexiletine	None ¹	None	0	0	0	None	+++	12 h
Procainamide	\downarrow^1	↑↓²	†↓²	↑ ↑	† †	+	+++	3–4 h
Propafenone	0,↓	î	1	$\uparrow\uparrow\uparrow$	0	+	+++	5–7 h
Propranolol	$\downarrow\downarrow$	↑↑	† †	0	0	+	+	5 h
Quinidine	↑↓ ^{1,2}	1¢↓2	1¢↓2	$\uparrow\uparrow$	↑ ↑	+	++++	6 h
Sotalol	$\downarrow\downarrow$	↑ ↑	† †	0	$\uparrow\uparrow\uparrow$	++++	++++	7 h
Verapamil	↓↓	$\uparrow\uparrow$	^	0	0	++++	-	7 h
Vernakalant		1	Munir Gharaib	eh MD, PhD,	MHPE	+++	-	2 h ²⁹

TABLE 14-3 Clinical pharmacologic properties of antiarrhythmic drugs.

Class 1A Drugs



Cinchona tree \rightarrow Antipyretic.

- Antimalarial.
 - Prototype,

Quinidine:

Inhibits α and muscarinic receptors.

Slows upstroke, conduction, and prolongs APD and QRS duration.

Quinidine

Use restricted to patients with normal hearts(no failure, no ischemia), but have atrial or ventricular arrhythmias.
 Occasionally used in acute severe malaria.

Quinidine

Side Effects: Toxic

Nausea (18%), Diarrhea (33%).

- Headache, Dizziness, and tinnitus= Cinchonism
- Hypersensitivity, fever, rash, angioedema.
- Thrombocytopenia.
- Excessive prolongation of QT interval, slowed conduction and sudden death (TdP)
- Hypotension.
- Serum Digoxin levels.
- 1 Warfarin effects.
- Sudden death.

Class 1A Drugs

Procainamide:



 Oral, but has short t¹/₂.
 L.E. (30% of patients Tx over 6 moths)

- Acetylated \rightarrow NAPA (Class III) action
 Disopyramide
- More anticholinergic effects but less diarrhea than quinidine

Class 1B Drugs

Lidocaine: 🛛

- High affinity to bind with activated and inactivated Na+ channels with rapid kinetics.
 - Acts selectively in ischemic tissue to promote conduction & block reentry. More effective with ↑ K+. Not effective in atrial arrhythmias. ☑

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Lidocaine

Class 1B Drugs

<u>Lidocaine:</u> Kinetics:

- Well absorbed, but ineffective orally, due to first pass effect, so given IV.
- Well distributed, including the brain.

Side Effects:

- Least cardiotoxic of the class, except for hypotension with high doses due to depression of the myocardium.
- CNS: parasthesia, tremor, nausea, slurred speech, and convulsions.
- Was routinely given to all MI patients to prevent ventricular arrhythmias.

Class 1B Drugs

<u>Tocainide:</u>

- Oral analog of lidocaine.
- CNS, GI and blood dyscrasia.

<u>Mexiletine:</u>

- Oral analog of lidocaine.
- Neurologic side effects.

Phenytoin: 🗩

- Digitalis induced arrhythmias.
 - Epilepsy.

- Arrhythmias after congenital heart surgery.
- Congenital prolonged QT interval.

Class 1C Drugs

Flecainide:

Potent blocker of Na + and K+ channels. **Negative inotropic effect. Proarrhythmic** \rightarrow **ventricular**. **Effective in supra ventricular** tachycardia with normal hearts. Side Effects: Ventricular arrhythmias, CNS, and sudden death.

Class 1C Drugs

Propafenone:

Blocks Na+ channels but also has beta blocking and Ca++ blocking activity.
 No effect on QT interval.
 Used for supraventricular arrhythmias.
 Side effects: metallic taste, constipation, and arrhythmias.

Class II Drugs

<u>Propranolol:</u>

- Besides beta blocking, membrane stabilization, and intrinsic sympathmimetic activities, has effective antiarrhythmic activity
- Very effective, well tolerated, and documented to reduce mortality after acute myocardial infarction by reducing arrhythmias.

<u>Esmolol:</u>

Class II Drugs

- Short acting, used in intraoperative and acute arrhythmias
- β1 selective
- No membrane stabilization effect.

<u>Acebutolol:</u>

- Short acting, used in intraoperative and acute arrhythmias.
- β1-selective.
- Also has direct membrane stabilizing. Munir Gharaiben MD, PhD, MHPE

Class III Drugs

<u>Amiodarone:</u>



- Blocks K+ channels and markedly prolongs APD.
- Class I actions.
- **Blocks** α and β Receptors.
 - Ca++ blocking actions.
- So, effect is due to alteration of lipid membrane.
 - Reserved for life-threatening atrial and ventricular arrhythmias.
- Slows heart rate and AV conduction.
- Low incidence of TdP despite significant QT prolongation.
- **11**^{27/17} Peripheral vasodifator (orby with IV).

Class III Drugs

<u>Amiodarone:</u>

Given IV (Loading dose 10gm) and orally.
 Slow kinetics (t¹/₂ 25-110 days), metabolized by CYP3A4 enzymes.

<u>Toxicity:</u> mainly extracardiac and dose related.

- Lung fibrosis (1%).
- CNS.
- Thyroid(hypo and hyper).
- Gl and liver.
- Corneal deposits,
- Skin: photodermatitis and discoloration.
- Digoxin & Anticoagulants.

Interactions: affected by CYP3A4 activity Munir Gharaibeh MD, PhD, MHPE

Class III Drugs

Sotalol:

- Beta blocker and Class III actions.
- For atrial and ventricular arrhythmias.
- Bradycardia, HF, Prolongation of QT.

Bretylium Tosylate:

- Originally an antihypertensive, but tolerance develops.
- Releases NE, then \$\frac{1}{2}\$ Release / Reuptake
- Rarely used except in the prevention of ventricular fibrillation after failure of cardiversion and lidocaine.
- Hypotension, Parotid swelling.
- Ibutilide.
- Dofetilide.

Class IV Drugs (Ca++ Channel Blockers)

<u>Verapamil</u>

- <u>Diltiazem</u>
- Block activated and inactivated L-type Ca++ channels.
- Effects more marked in tissues that fire frequently, less completely polarized at rest, and those dependant on Ca++ (SA node and AV node).
- Paroxysmal Supraventricular Tachycardia.
- Vasodilators and have negative inotropic effects.
- Can cause severe AV block in diseased hearts.
- Safe: Constipation, gastric discomfort, vertigo, headache, nervousness, pruritis.
- Digoxin levels.

Miscellaneous Drugs

Digoxin: 🗩

- Old fashioned agent for atrial arrhythmias.
 - Direct Actions.
 - Vagotonic Effects.
 - ↑ AV refractoriness.

Miscellaneous Drugs

<u>Magnesium:</u>

- Works on Na+/K+ ATPase, Na+ channels, certain K+ channels and Ca++ channels.
- Effective IV in refractory digitalis- induced ventricular arrhythmias only in hypomagnesemic patients.
- Also, in TdP patients even if serum Mg++ is normal.

Potassium salts:

- For digitalis- induced arrhythmias with hypokalemia.
- Depress ectopic pacemakers and slow Munir Gharaibeh MD, PhD, MHPE conduction.

Miscellaneous Drugs

Adenosine:

- Naturally occurring nucleoside.
 - Stimulates purinergic(P1) receptors.
- Activates inward rectifier K+ current and inhibits Ca++ current.
 - Very short acting (t 1/2 10 seconds).
 - \downarrow Phase 4 depolarization in SA node.
 - \downarrow AV conduction.
 - No effect on ventricles.

Miscellaneous Drugs Adenosine:

- 90-95% effective in supraventricular tachycardia, replaced verapamil.
- Less effective in the presence of adenosine receptor blockers, e.g. theophylline and caffeine.

Can cause transient flushing (20%), chest tightness, AV block, headache, hypotension, nausea, and paresthesia.