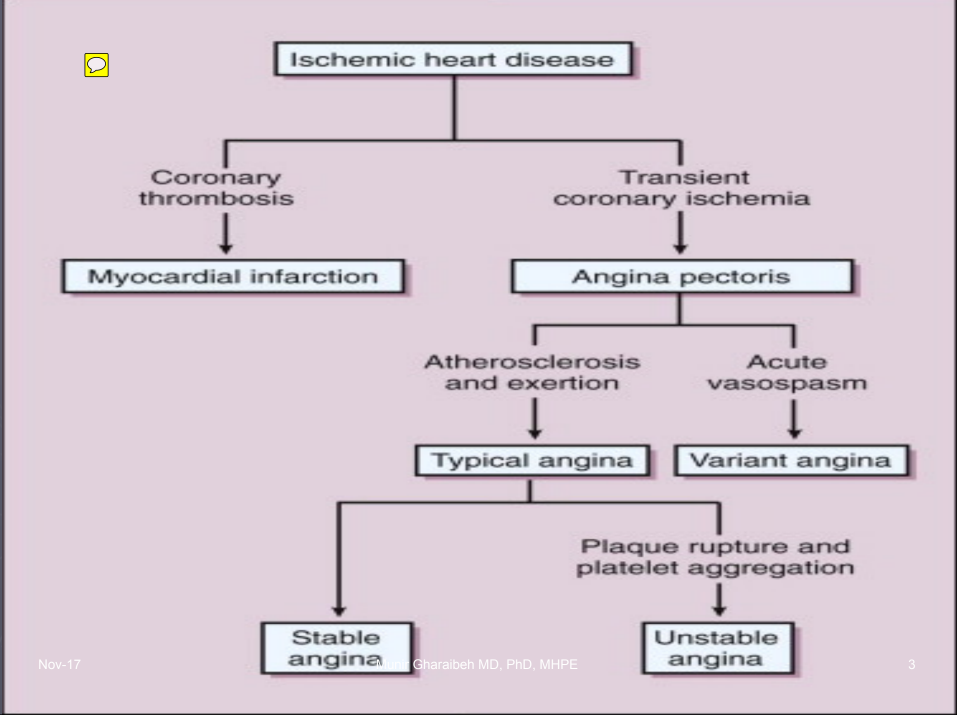
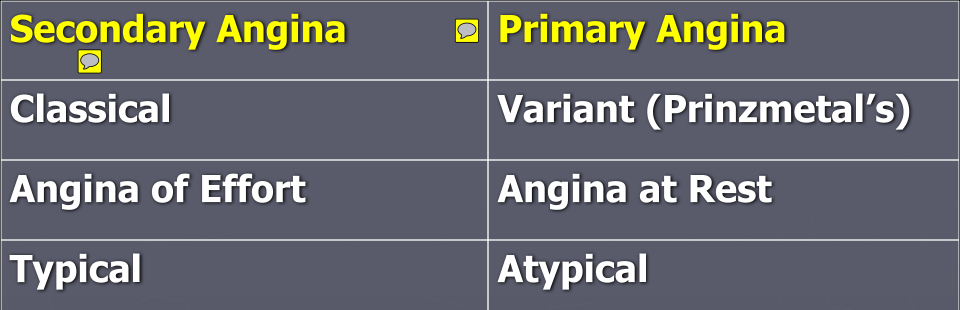
# **Drug Treatment of Ischemic Heart Disease**

Munir Gharaibeh, MD, PhD, MHPE
School of Medicine,
The University of Jordan
November, 2017

### **Categories of Ischemic Heart Disease**

Fixed "Stable", Effort Angina
Variant Angina "Primary Angina"
Unstable Angina
Myocardial Infarction





1768

**Small vessels** 

**Atherosclerosis** 

**ST**wdepression

Single or multiple p

1957

Single 🗩

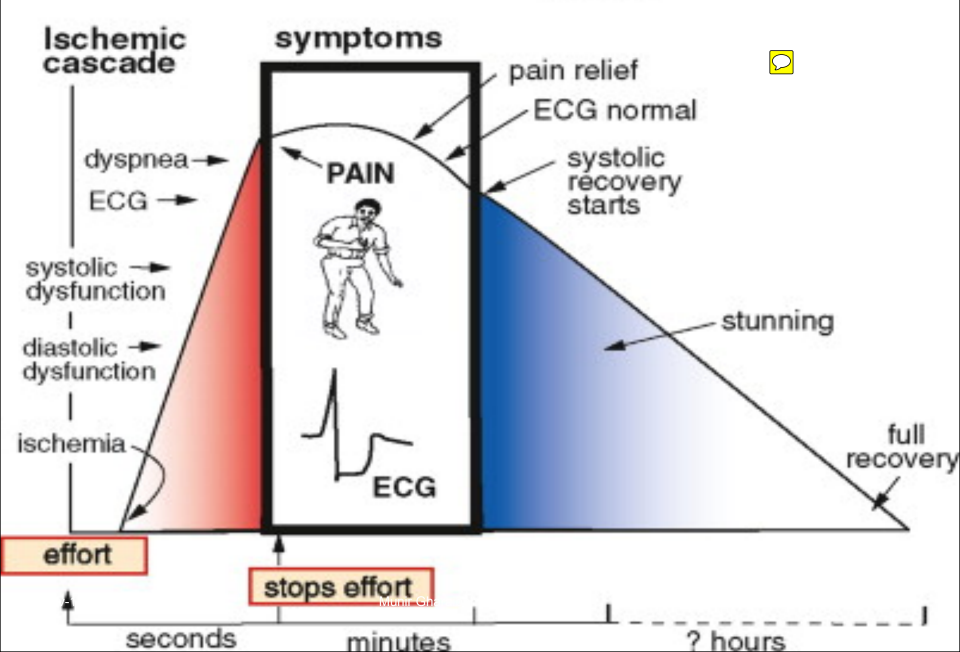
Munir Gharaibeh MDShT, Melevation

Vasospasm

Large vessels

### EFFORT ANGINA

Opie 2008

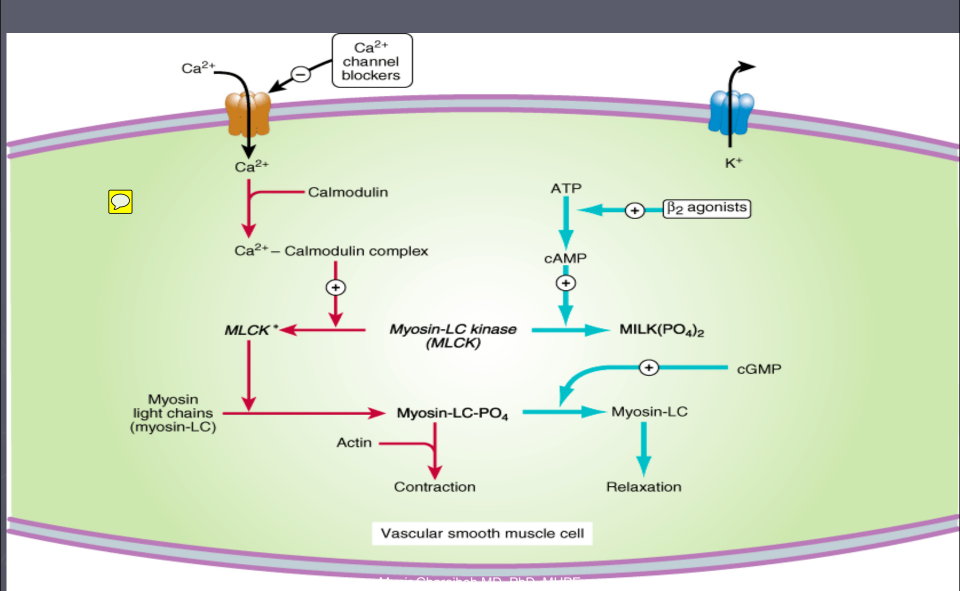


- Stunning?(مدوخ): □
  - Myocardial stunning is the reversible reduction of function of heart contraction after reperfusion not accounted for by tissue damage or reduced blood flow.

### **Control of smooth muscle contraction**

- Contraction is triggered by influx of calcium through L-type transmembrane calcium channels.
- Calcium combines with calmodulin to form a complex that converts the enzyme myosin light-chain kinase to its active form (MLCK\*).
- MLCK phosphorylates myosin light chains, thereby initiating the interaction of myosin with actin.
- ► Beta2 agonists (and other substances that increase cAMP) may cause relaxation in smooth muscle by accelerating the inactivation of MLCK and by facilitating the expulsion of calcium from the cell.

### Control of vascular smooth muscle contraction



Source: Katzung BG, Masters SB, Trevor AJ: Basic & Clinical Pharmacology, 11th Edition: http://www.accessmedicine.com
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# **Mechanism of IHD**

Imbalance of the ratio:

O<sub>2</sub> Supply (Coronary Blood Flow)

O<sub>2</sub> Demand (Work of the Heart)

# Major Determinants of Myocardial Oxygen Supply and Demand

#### Oxygen supply

Oxygen extraction (%)

Coronary blood flow

Aortic diastolic pressure

Coronary arteriolar

resistance

Metabolic autoregulation

Endocardial-epicardial

flow O

Coronary collateral

blood flow

Large coronary artery

diameter

#### Oxygen demand

Wall tension

Ventricular volume

Radius or heart size

Ventricular pressure

Systolic pressure

(afterload)

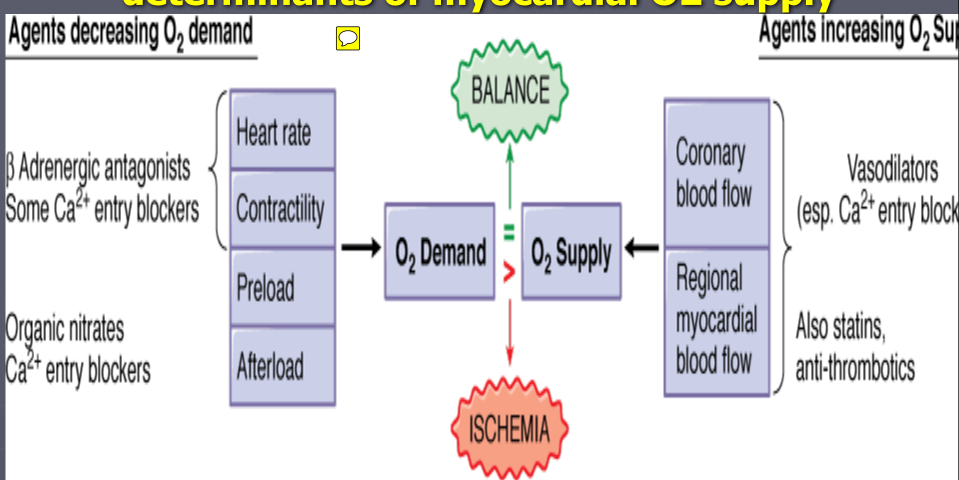
Diastolic pressure

(preload)

Heart rate

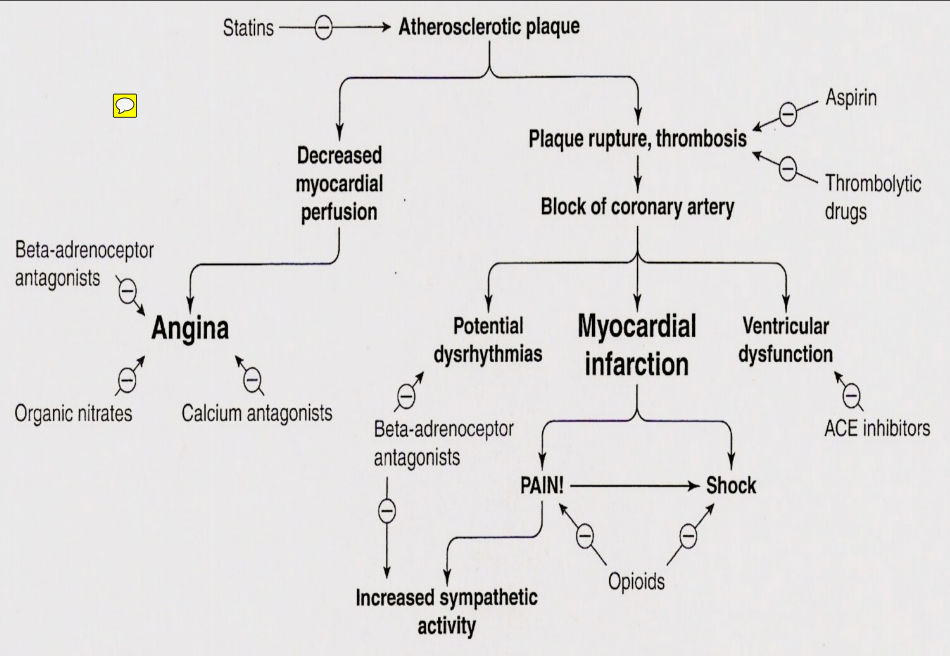
Contractility

# Pharmacological modification of the major determinants of myocardial O2 supply

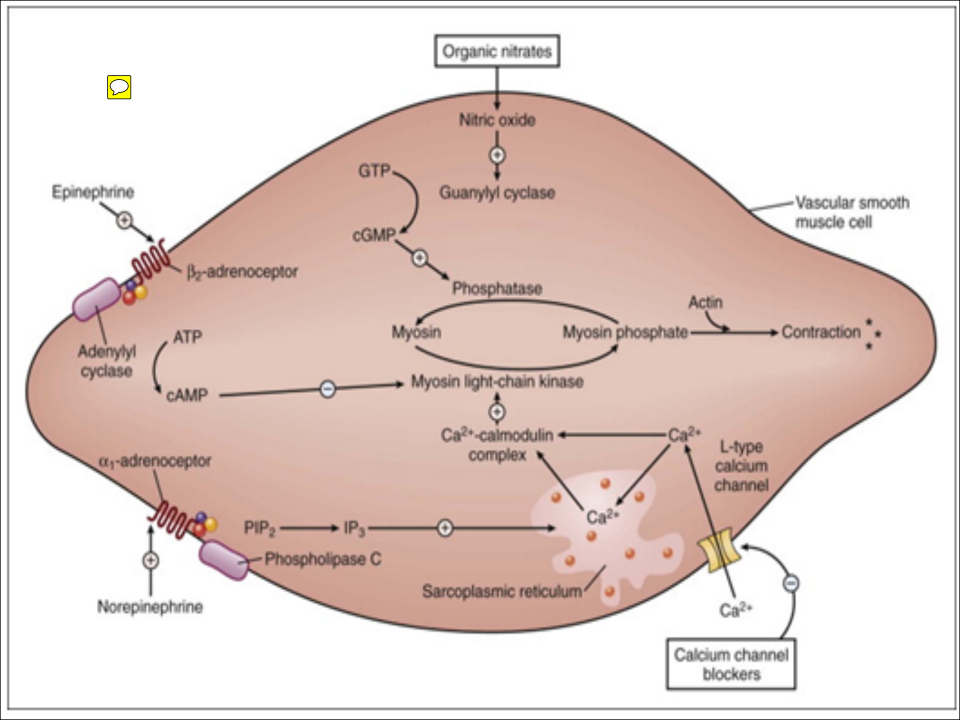


Source: Brunton LL, Chabner BA, Knollmann BC: Goodman & Gilman's The Pharmacological Basis of Therapeutics, 12th Edition: www.accessrnedicine.com

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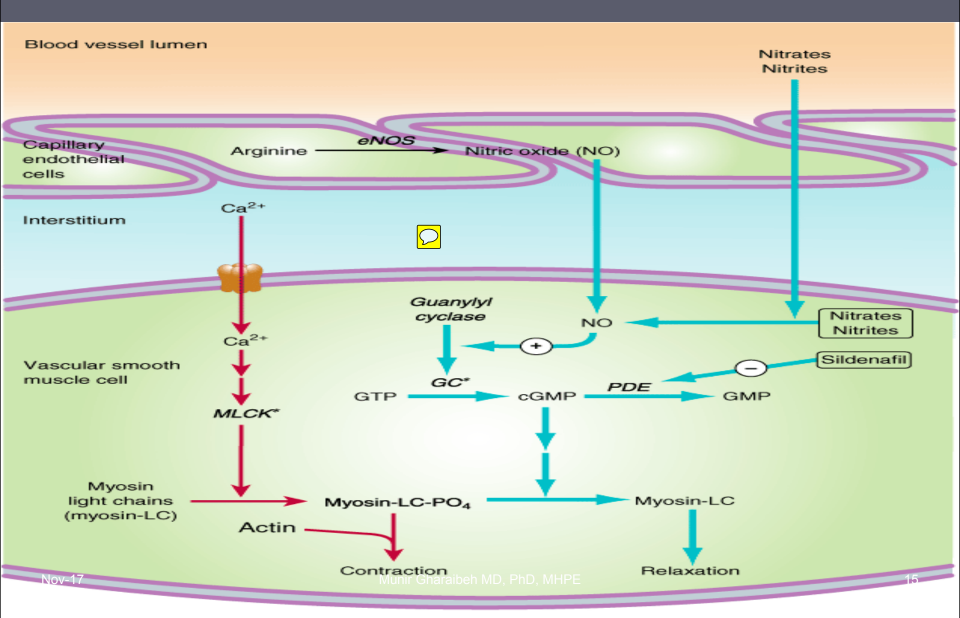
19.1 Simplified diagram of atherosclerosis, angina and myocardial infarction, and drugs used in treatment.



### **Organic Nitrates**

- ► <u>Nitroglycerine (GTN):</u> □
- ► Prototype, used for more than 150 years.
- ► Nonspecific smooth muscle relaxant.
- Action is due to release of NO, leading to activation of guanylyl cyclase.
- Action not antagonized by any known antagonist.

# Nitrates, nitrites, and other substances that increase the concentration of nitric oxide (NO) in vascular muscle



## Nitroglycerine (GTN)

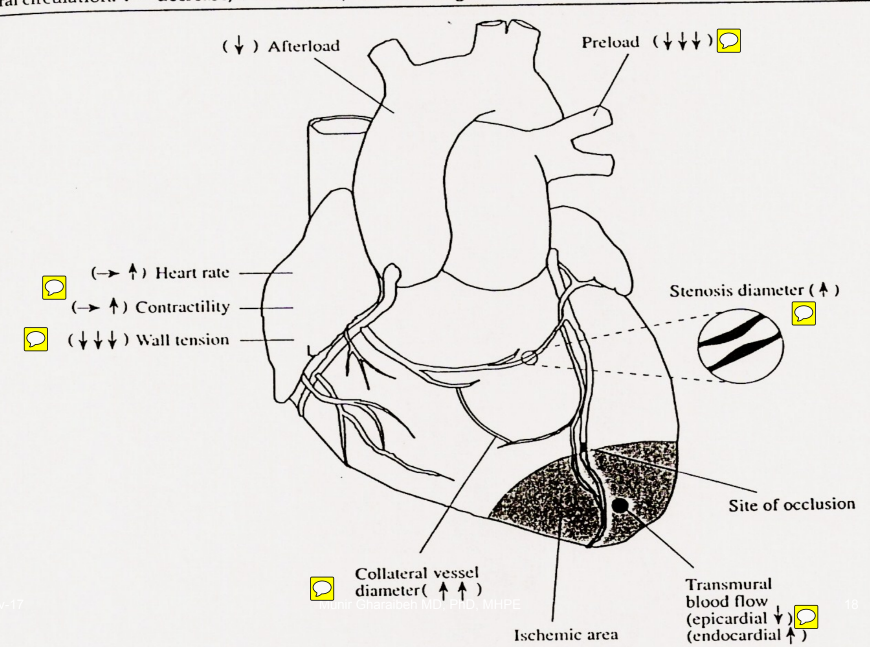
- ▶ Usually administered sublingually. ▶
- ► Can be administered by various routes.
- ► Fast onset of action(1-3minutes, Peaks at 10 minutes).
- ► Short duration (15-30minutes).
- Reductase enzyme, in liver, breaks down the drug.

## Nitroglycerine (GTN) **□**

- Causes general vasodilation:
- ► Arteriolar dilation: short lived (5-10 min)
  - Decreases systemic blood pressure (afterload), but causes reflex tachycardia and increased contractility might increase
     MVO2.
- **Venous dilation**: more intense, even with low doses, lasts for 30 minutes.
  - Decreases venous return (preload) and decreases MVO2.

Figure 19-2

A schematic drawing indicating the major actions of the nitrates on the ischemic heart and peripheral circulation.  $\downarrow = \text{decrease}$ ;  $\uparrow = \text{increase}$ ;  $\rightarrow = \text{unchanged}$ ;  $\updownarrow = \text{variable effect}$ .



## TABLE 12-2 Beneficial and deleterious effects of nitrates in the treatment of angina.

Effect	P	Result		
Potential beneficial effects				
Decreased ventricu Decreased arterial Decreased ejection	pressure	Decreased myocardial oxygen requirement		
Vasodilation of epi onary arteries	cardial cor-	Relief of coronary artery spasm		
Increased collatera	l flow	Improved perfusion to ischemic myocardium		
Decreased left ven diastolic pressure	tricular	Improved subendocardial perfusion		
Potential deleterious effects				
Reflex tachycardia		Increased myocardial oxygen requirement		
Reflex increase in o	contractility	Increased myocardial oxygen requirement		
Decreased diastolic	•	Decreased coronary perfusion on MD, PhD, MHPE 19		

### Nitroglycerine (GTN)

- **► Side Effects:**
- ► Headache.
- Hypotension and tachycardia.
- ► Increased intraocular and intracranial pressures.
- Methemoglobinemia.
- **▶** Tolerance: only for the arteriolar effects.
- ► Withdrawal: in workers in ammunition industry. □

### **Preparations of Nitrate**

Drug 🖸	Duration of Action
Short-acting:	
Nitroglycerin, sublingual	10-30 minutes
Isosorbide dinitrate, sublingual	10-60 minutes
Amyl nitrite, inhalant	3-5 minutes
Long-acting:	
Nitroglycerin, oral sustained- action	6–8 hours
Nitroglycerin, 2% ointment, transdermal	3–6 hours
Nitroglycerin, slow-release, buccal	3–6 hours
Nitroglycerin, slow-release patch, transdermal	8–10 hours
Isosorbide dinitrate, sublingual	1.5-2 hours
Isosorbide dinitrate, oral	4–6 hours
Isosorbide dinitrate, chewable oral	2-3 hours

## **Beta Adrenergic Blockers**

- ▶ Prevent actions of catecholamines, so more effective during exertion.
- Do not dilate coronary arteries, might constrict them.
- **▶** Do not increase collateral blood flow.
- ► Cause subjective and objective improvement: decreased number of anginal episodes, nitroglycerine consumption, enhanced exercise tolerance, and improved ECG.

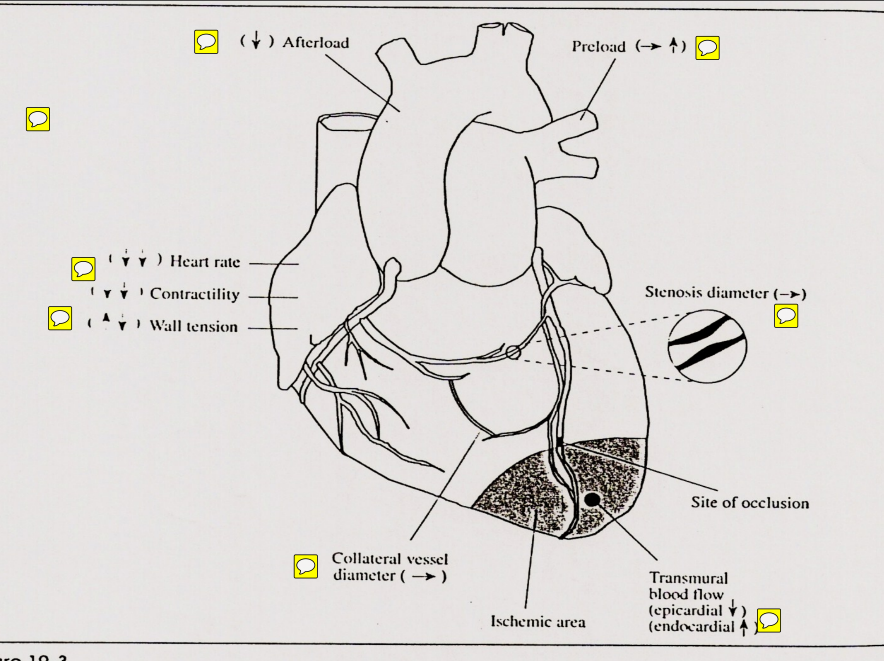


Figure 19-3

A schematic drawing indicating the major actions of the  $\beta$ -blockers on the ischemic heart and peripheral circulation. For key, see Fig. 19-2.

# Calcium Channel Blockers

Particularly beneficial in vasospasm.

Can also affect platelets aggregation.

May be dangerous in the presence of heart failure and in patients susceptible to hypotension.



Т

N

P/Q

Ca<sub>v</sub>1.1-

Ca<sub>v</sub>3.1-

Ca<sub>v</sub>3.3

Ca<sub>v</sub>2.2

Ca<sub>v</sub>2.1

Ca<sub>v</sub>2.3

Ca<sub>v</sub>1.3

#### Properties of Several Recognized Voltage-Activated Calcium Channels

Long, large,

Short, small,

Short, high

Long, high

**Pacemaking** 

threshold

threshold

low threshold

high threshold

Verapamil,

DHPs,

 $Cd^{2+}$ ,

sFTX,

Ni<sup>2+</sup>,

aga-IIIA

flunarizine,

mibefradil<sup>1</sup>

Ziconotide,3 ga

aga-IIIA, Cd<sup>2+</sup>

bapentin,4

CTX-GVIA.

-CTX-

MVIIC,

aga-IVA

SNX-482,

aga-IIIA

Properties of Several Recognized Voltage-Activated Calcium Chainleis.					
			<b>Properties of</b>	<b>Blocked By</b>	
	Name		the Calcium		
			Current p	$\bigcirc$	

Cardiac, skeletal, smooth

**Heart, neurons** 

Neurons, sperm<sup>2</sup>

Neurons

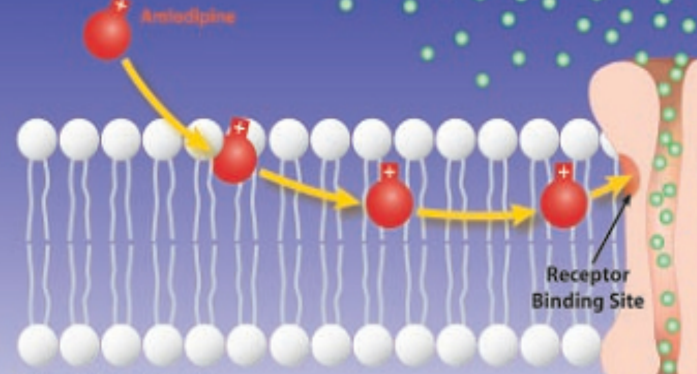
muscle, neurons (Ca<sub>v</sub>1.4 is found

in retina), endocrine cells, bone

Neurons, sperm<sup>2</sup>
Munir Gharaibeh MD, PhD, MHPE

### Cell Plasma Membrane

#### Calcium Ions

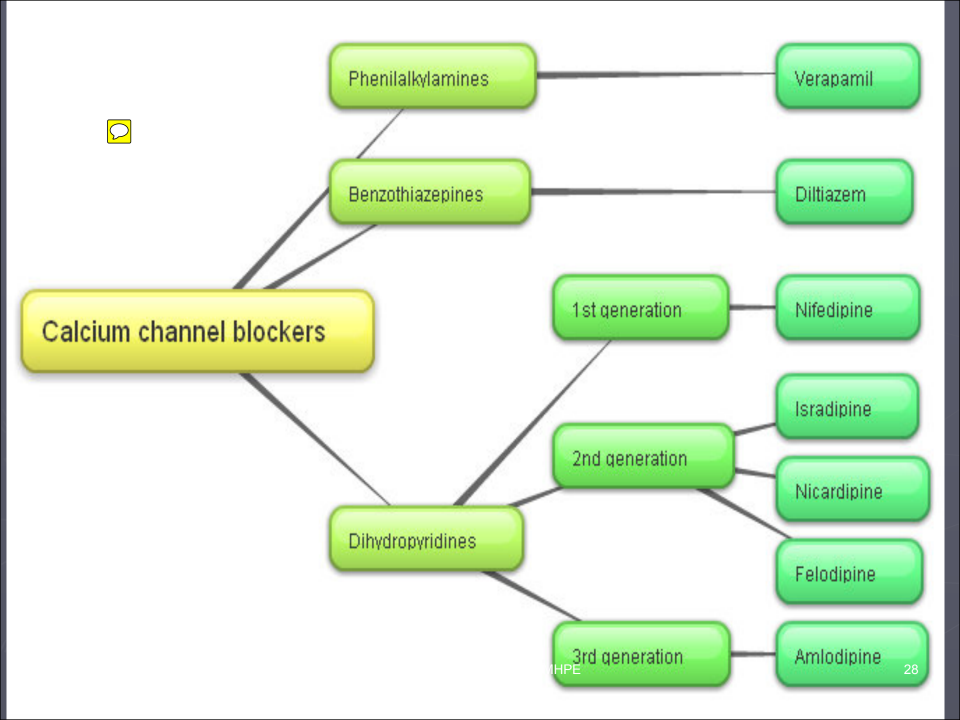


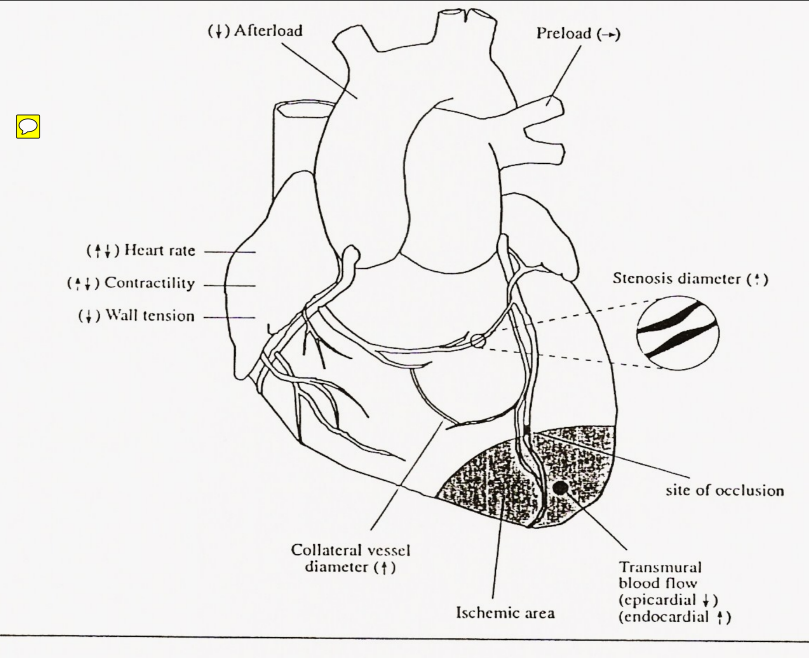
**Phospholipid Bilayer** 

L-type Calcium Channel

Source: Katzung BG, Masters SB, Trevor AJ: Basic & Clinical Pharmacology, 11th Edition: http://www.accessmedicine.com

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**Figure 19-4**A schematic drawing indicating the major actions of the calcium antagonists on the ischemic heart and coronary circulation. For key, see Fig. 19-2.

Drug	Oral Bioavailability (%)	Half-Life (hours)	Indication
Dihydropyridines			
Amlodipine	65-90	30-50	Angina, hypertension
Felodipine	15-20	11-16	Hypertension, Raynaud's phenomenon
Isradipine	15-25	8	Hypertension
Nicardipine	35	2-4	Angina, hypertension
Nifedipine	45-70	4	Angina, hypertension, Raynaud's phenomenon
Nimodipine	13	1-2	Subarachnoid hemorrhage
Nisoldipine	< 10	6-12	Hypertension
Nitrendipine	10-30	5-12	Investigational
Miscellaneous			
Diltiazem	40-65	3-4	Angina, hypertension, Raynaud's phenomenon
Verapamil	20-35 Munir Gharalbeh N	AD 6 D MHPE	Angina, hypertension, arrhythmias, migraine

### **Calcium Channel Blockers**

- **► Side Effects:**
- **►** Hypotension.
- ► Headache, dizziness.
- ► Flushing.
- ► Peripheral edema.

# Effects of Nitrates Alone and with Beta Blockers or Calcium Channel Blockers in Angina Pectoris. Nitrates Alone Beta Blockers or Combined

Nitrates Alone

Beta Blockers or Calcium Channel Blockers or Calcium Channel Blockers

Calcium Channel Blockers or Calcium Channel Blockers

Decrease

Decrease

**Increase** 

Decrease

MDradree

Decrease

Decrease

Non

Non

Non or decrease

Reflex<sup>1</sup> increase

Reflex<sup>1</sup> increase

Decrease

Decrease

Decrease

**Heart rate** 

**Arterial pressure** 

**End-diastolic** 

Contractility

**Ejection time** 

volume

# Dipyridamole

- ► Thought to be a good coronary dilator.
- Increases the blood flow to the normal area i.e. "Coronary Steal Phenomenon". 

  ■
- Still used as an antiplatelet drug (in Nov-17 ☐ TIAs), but not better than aspirin.

### **Others**

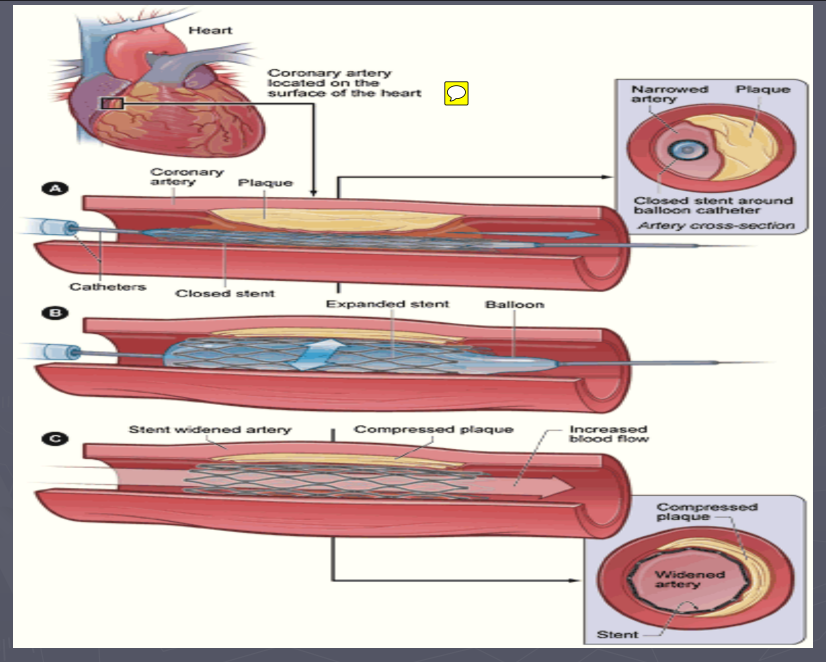
► ACEI. □

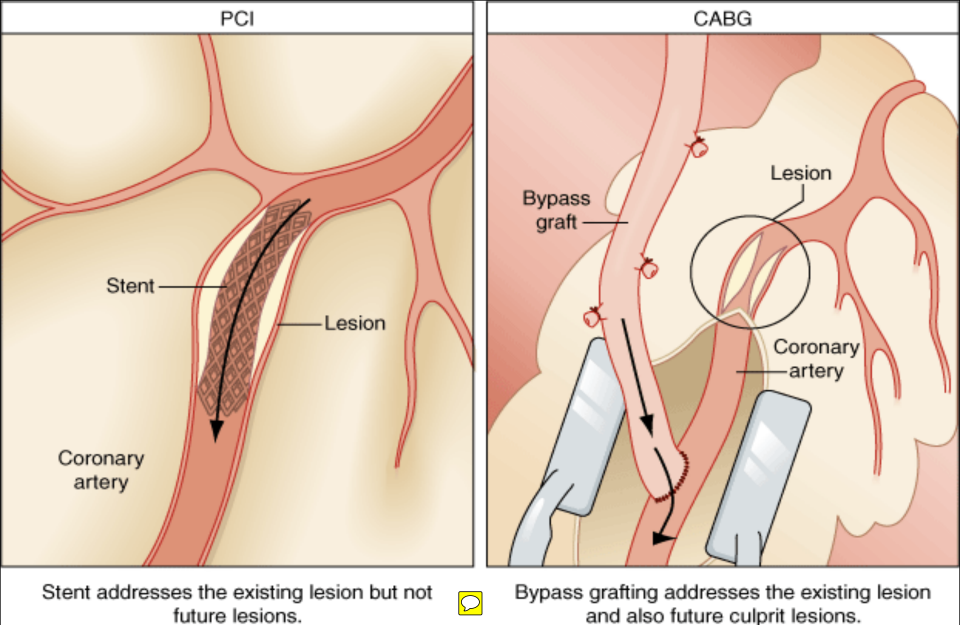
Anticoagulants and/or Thrombolytic Therapy.

Cholesterol Lowering Agents.

Angioplasty







Source: Fauci AS, Kasper DL, Braunwald E, Hauser St., Longo DL, Jameson JL, Loscalzo J: *Harrison's Principles of Internal Medicine*, 17th Edition: http://www.accessmedicine.com

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## Newer Antianginal Drugs

- ► Metabolic modulators: Ranolazine.
- **▶** Direct bradycardic agents: Ivabradine.
- ► Potassium channel activators: Nicorandil.
- ► Rho-kinase inhibitors: Fasudil.
- > Sulfonylureas: Glibenclamide.
- **►** Thiazolidinediones.
- **►** Vasopeptidase inhibitors.
- ► Nitric oxide donors: L- arginine.
- ► Capsaicin.
- **A**miloride.