Semey State Medical University



Physiology of the Heart

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Plan:

- *Functions of the Heart

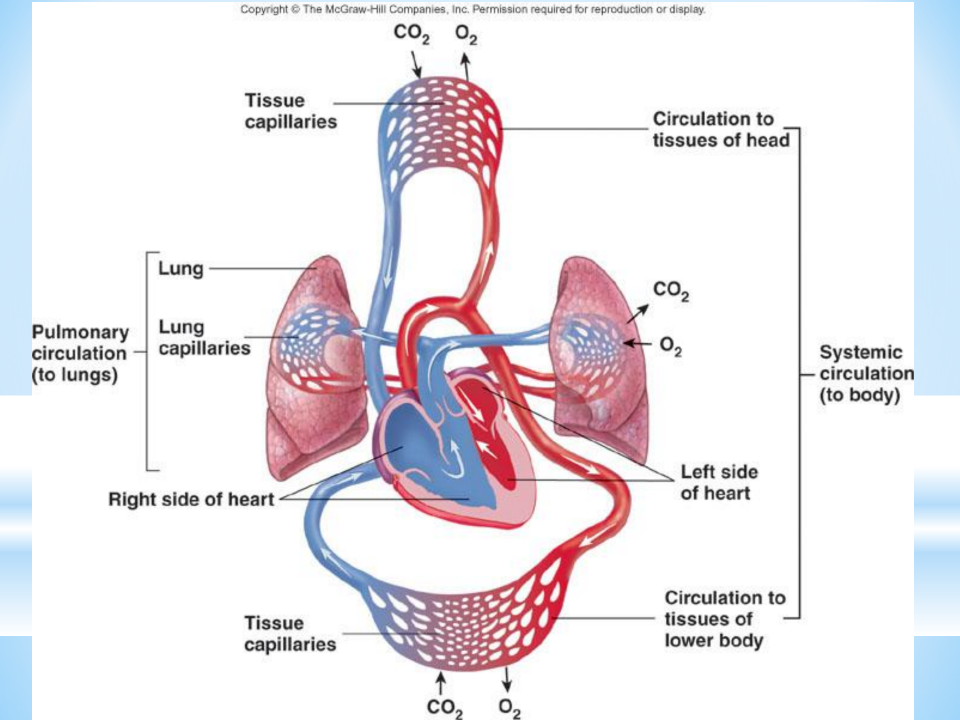
 * Conducting System of Heart
- *An Electrocardiogram
- *The Cardiac Cycle
 *Regulation of the Heart

*Functions of the Heart

- *Generating blood pressure
- *Routing blood: separates pulmonary and systemic circulations
- *Ensuring one-way blood flow: valves
- *Regulating blood supply
 - *Changes in contraction rate and force match blood delivery to changing metabolic needs

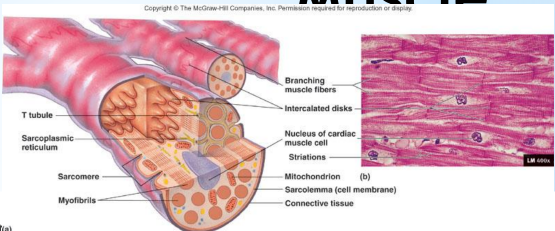
*The cardiovascular system is divided into two circuits

- *Pulmonary circuit
 - *blood to and from the lungs
- *Systemic circuit
 - *blood to and from the rest of the body
- *Vessels carry the blood through the circuits
 - *Arteries carry blood away from the heart
 - *Veins carry blood to the heart
 - *Capillaries permit exchange



- *Elongated, branching cells containing 1-2 centrally located nuclei
- *Contains actin and myosin myofilaments
- *Intercalated disks: specialized cell-cell contacts.
 - *Cell membranes interdigitate
 - *Desmosomes hold cells together
 - *Gap junctions allow action potentials to move from one cell to the next.
- *Electrically, cardiac muscle of the atria and of the ventricles behaves as single unit
- Mitochondria comprise 30% of volume of the cell vs. 2% in skeletal





- *Structural Differences in heart chambers
 - *The left side of the heart is more muscular than the right side
- *Functions of valves
 - *AV valves prevent backflow of blood from the ventricles to the atria
 - *Semilunar valves prevent backflow into the ventricles from the pulmonary trunk and aorta

*Heart chambers and valves

*Heart muscle:

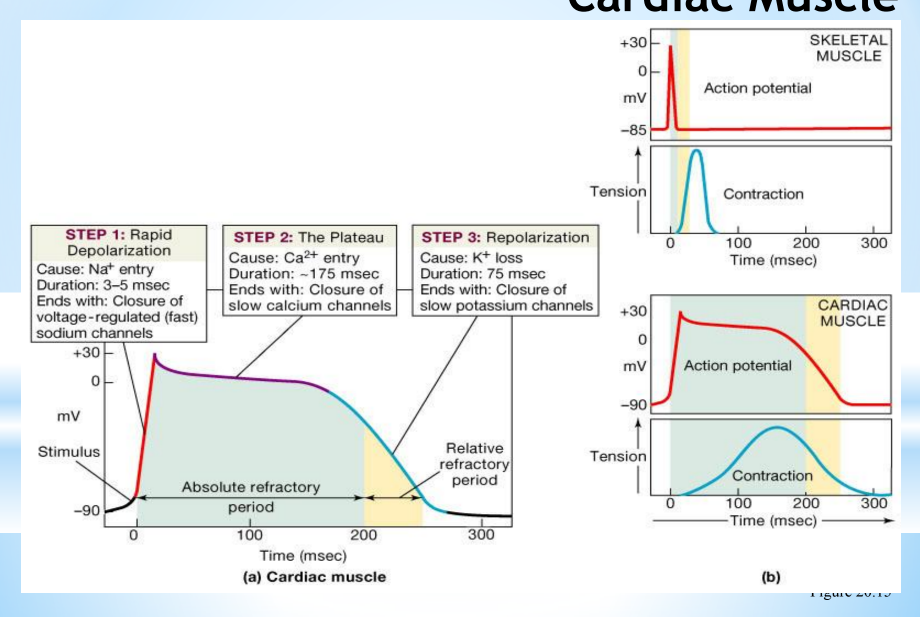
- *Is stimulated by nerves and is self-excitable (automaticity)
- *Contracts as a unit; no motor units
- *Has a long (250 ms) absolute refractory period
- *Cardiac muscle contraction is similar to skeletal muscle contraction, i.e., sliding-filaments

*Cardiac Muscle Contraction

*Differences Between Skeletal and Cardiac Muscle Physiology

- * Action Potential
 - * Cardiac: Action potentials conducted from cell to cell.
 - * Skeletal, action potential conducted along length of single fiber
- * Rate of Action Potential Propagation
 - Slow in cardiac muscle because of gap junctions and small diameter of fibers.
 - * Faster in skeletal muscle due to larger diameter fibers.
- * Calcium release
 - * Calcium-induced calcium release (CICR) in cardiac
 - * Movement of extracellular Ca²⁺ through plasma membrane and T tubules into sarcoplasm stimulates release of Ca²⁺ from sarcoplasmic reticulum
 - * Action potential in T-tubule stimulates Ca⁺⁺ release from

*The Action Potential in Skeletal and Cardiac Muscle



* Electrical Properties of Myocardial

- 1. Rising phase of action potential
 - Due to opening of fast Na⁺ channels

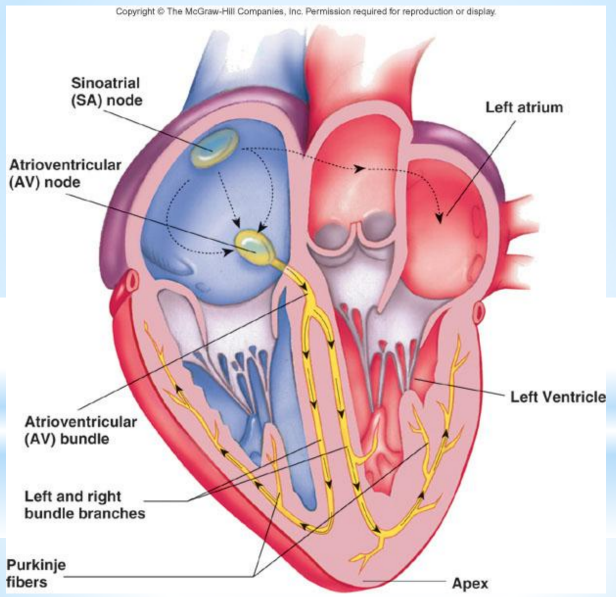
2. Plateau phase

- Closure of sodium channels
- Opening of calcium channels
- Slight increase in K⁺ permeability
- Prevents summation and thus tetanus of cardiac muscle

3. Repolarization phase

- Calcium channels closed
- Increased K⁺ permeability

*Conducting System of Heart



*SA node: sinoatrial node. The pacemaker. *Sa rode: Heart

- * Specialized cardiac muscle cells.
- *Generate spontaneous action potentials (autorhythmic tissue).
- * Action potentials pass to atrial muscle cells and to the AV node
- *AV node: atrioventricular node.
 - * Action potentials conducted more slowly here than in any other part of system.
 - *Ensures ventricles receive signal to contract after atria have contracted
- *AV bundle: passes through hole in cardiac skeleton to reach interventricular septum
- *Right and left bundle branches: extend beneath endocardium to apices of right and left ventricles
- *Purkinje fibers:
 - *Large diameter cardiac muscle cells with few myofibrils.
 - * Many gap junctions.

*Autorhythmic cells:

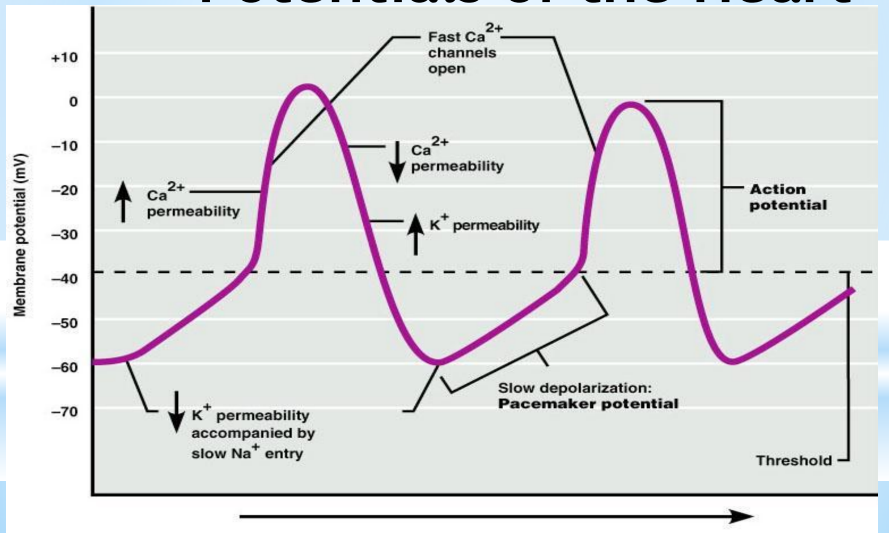
- *Initiate action potentials
- *Have unstable resting potentials called pacemaker potentials
- *Use calcium influx (rather than sodium) for rising phase of the action potential

*Heart Physiology: Intrinsic Conduction System

*Depolarization of SA Node

- *SA node no stable resting membrane potential
- *Pacemaker potential
 - *gradual depolarization from -60 mV, slow influx of Na⁺
- *Action potential
 - *occurs at threshold of -40 mV
 - *depolarizing phase to 0 mV
 - * fast Ca²⁺ channels open, (Ca²⁺ in)
 - *repolarizing phase
 - *K⁺ channels open, (K⁺ out)
 - *at -60 mV K⁺ channels close, pacemaker potential starts over
- *Each depolarization creates one heartbeat

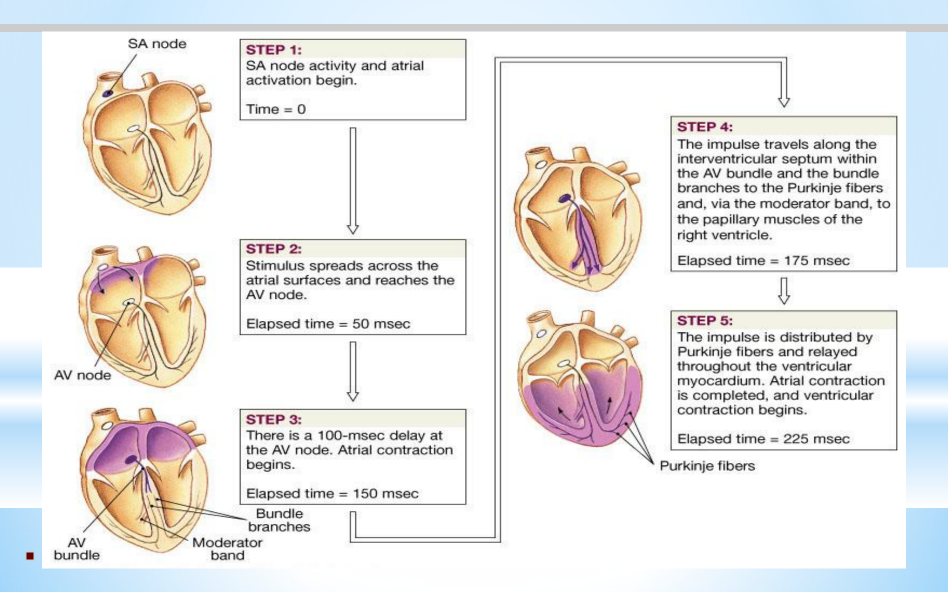
*Pacemaker and Action Potentials of the Heart



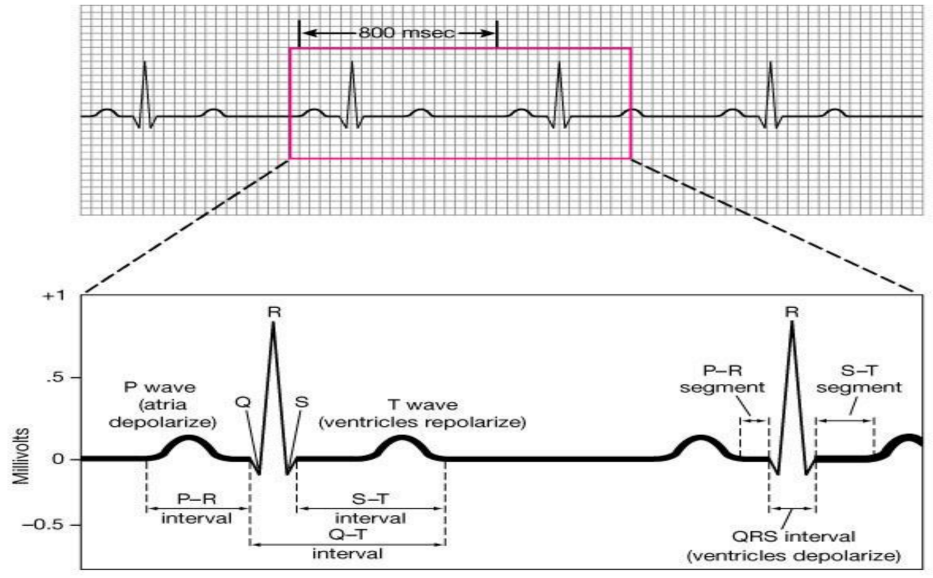
- *Sinoatrial (SA) node generates impulses about 75 times/minute
- *Atrioventricular (AV) node delays the impulse approximately 0.1 second
- *Impulse passes from atria to ventricles via the atrioventricular bundle (bundle of His) to the Purkinje fibers and finally to the myocardial fibers

*Heart Physiology: Sequence of Excitation

*Impulse Conduction through the Heart

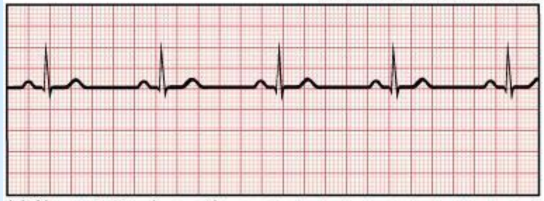


*An Electrocardiogram

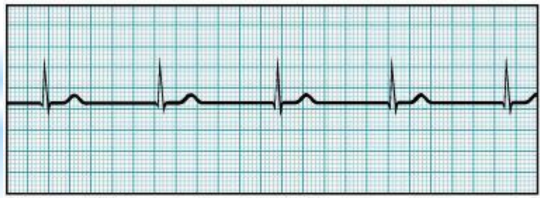


*Electrocardiogram

- *Record of electrical events in the myocardium that can be correlated with mechanical events
- *P wave: depolarization of atrial myocardium.
 - *Signals onset of atrial contraction
- *QRS complex: ventricular depolarization
 - *Signals onset of ventricular contraction..
- *T wave: repolarization of ventricles
- *PR interval or PQ interval: 0.16 sec
 - *Extends from start of atrial depolarization to start of ventricular depolarization (QRS complex) contract and begin to relax
 - *Can indicate damage to conducting pathway or AV node if greater than 0.20 sec (200 msec)
- *Q-T interval: time required for ventricles to undergo a single cycle of depolarization and repolarization
 - * Can be lengthened by electrolyte disturbances, conduction problems, coronary ischemia, myocardial damage

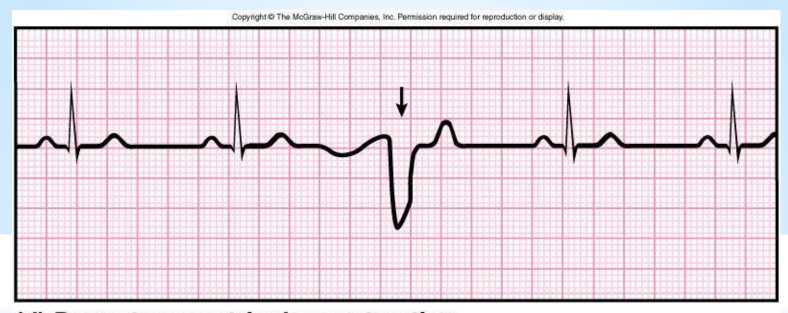


(a) Sinus rhythm (normal)



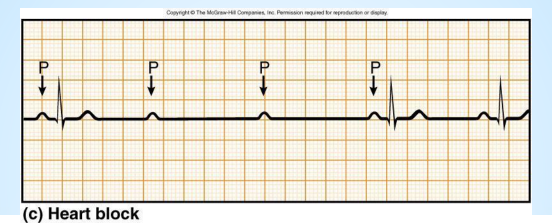
(b) Nodal rhythm - no SA node activity

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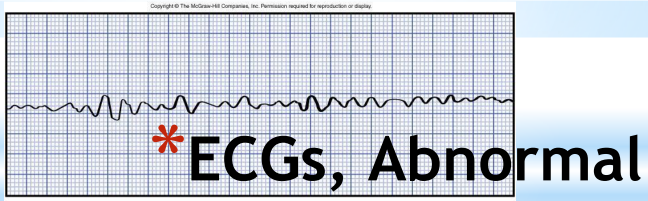


(d) Premature ventricular contraction **ECGs, Abnormal

Extrasystole: note inverted QRS complex, misshapen QRS and T and absence of a P wave preceding this contraction.



Arrhythmia: conduction failure at AV node



(e) Ventricular fibrillation

No pumping action occurs

- *Cardiac cycle refers to all events associated with blood flow through the heart from the start of one heartbeat to the beginning of the next
- *During a cardiac cycle
 - *Each heart chamber goes through systole and diastole
 - *Correct pressure relationships are dependent on careful timing of contractions

*The Cardiac Cycle

*Phases of the Cardiac Cycle

- *Atrial diastole and systole -
 - *Blood flows into and passively out of atria (80% of total)

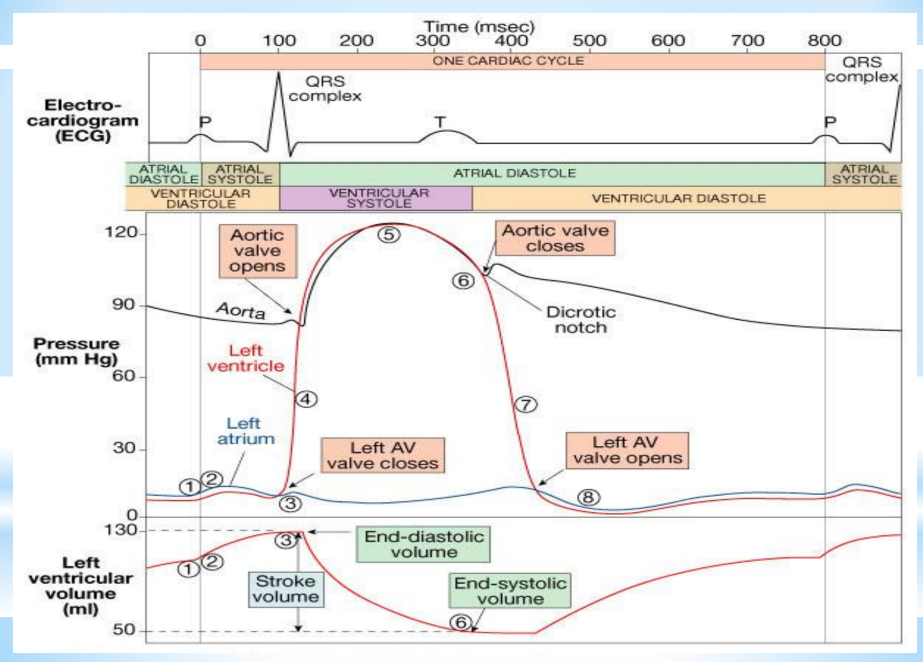
 *AV valves open
 - *Atrial systole pumps only about 20% of blood into ventricles
- *Ventricular filling: mid-to-late diastole
 - *Heart blood pressure is low as blood enters atria and flows into ventricles
 - *80% of blood enters ventricles passively
 - *AV valves are open, then atrial systole occurs
 - *Atrial systole pumps remaining 20% of blood into

*Phases of the Cardiac *Ventricular systole Cycle

- *Atria relax
- *Rising ventricular pressure results in closing of AV valves (1st heart sound 'lubb')
- *Isovolumetric contraction phase
 - *Ventricles are contracting but no blood is leaving
 - *Ventricular pressure not great enough to open semilunar valves
- *Ventricular ejection phase opens semilunar valves
 - *Ventricular pressure now greater than pressure in arteries (aorta and pulmonary trunk)

*Phases of the Cardiac Cycle

- *Ventricular diastole
 - *Ventricles relax
 - *Backflow of blood in aorta and pulmonary trunk closes semilunar valves (2nd hear sound "dubb"
 - *Dicrotic notch brief rise in aortic pressure caused by backflow of blood rebounding off semilunar valves
 - *Blood once again flowing into relaxed atria and passively into ventricles



Pressure and Volume Relationships in the Cardiac Cycle

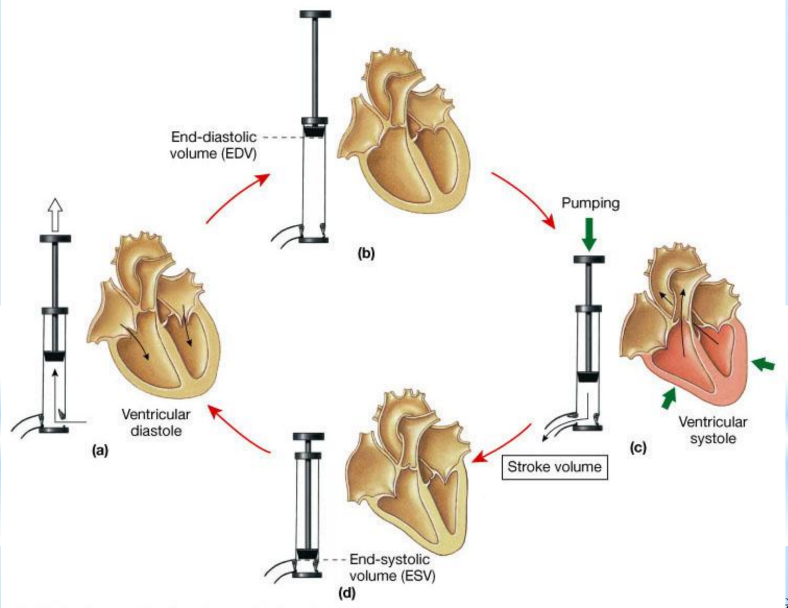
*Cardiac Output (CO) and Cardiac Reserve

- *CO is the amount of blood pumped by each ventricle in one minute
- *CO is the product of heart rate (HR) and stroke volume (SV)

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CO = HR x SV
(ml/min) = (beats/min) x ml/beat
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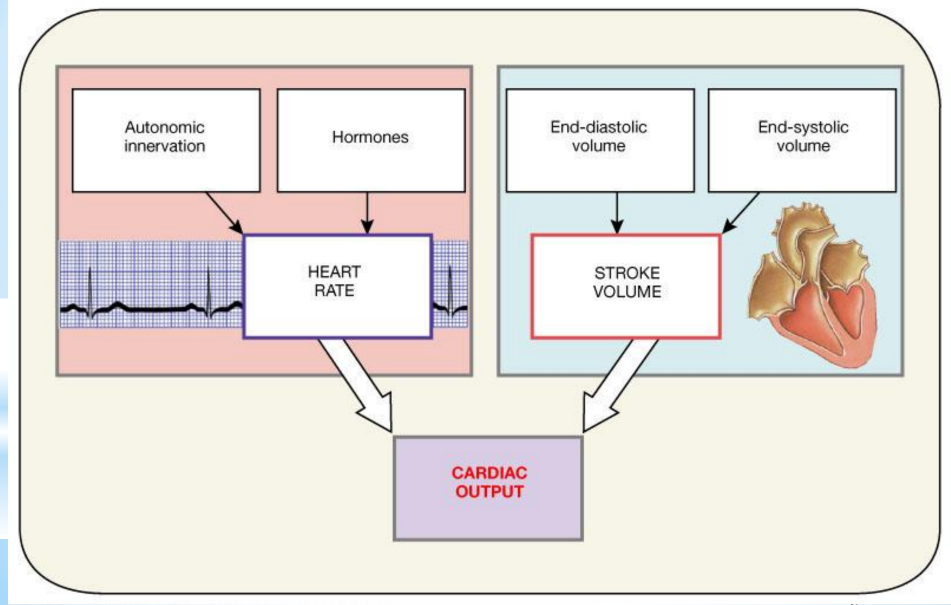
- *HR is the number of heart beats per minute
- *SV is the amount of blood pumped out by a ventricle with each beat
- *Cardiac reserve is the difference between resting and maximal CO

*A Simple Model of Stroke Volume



- *CO (ml/min) = HR (75 beats/min) x SV (70 ml/beat)
- *CO = 5250 ml/min (5.25 L/min)
- *If HR increases to 150 b/min and SV increases to 120 ml/beat, then
 - *CO = 150 b/min x 120 ml/beat
 - *co = 18,000 mi/minardiacooutput: An Example

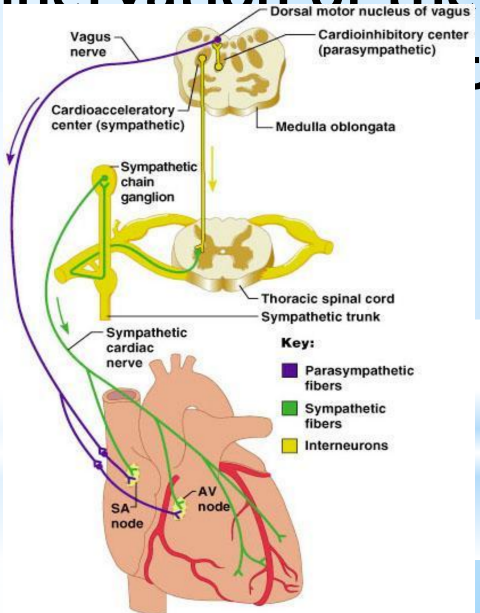
*Factors Affecting Cardiac Output



*Extrinsic Innervation of the

*Vital centers of medulla

- 1. Cardiac Center
- *Cardioaccelerator center
 - *Activates sympathetic neurons that increase HR
- *Cardioinhibitory center
 - *Activates parasympathetic neurons that decrease HR
- *Cardiac center receives input from higher centers (hypotha-lamus), monitoring blood pressure and dissolved gas concentrations



*Regulation of the Heart

*Neural regulation

- * Parasympathetic stimulation a negative chronotropic factor
 - *Supplied by vagus nerve, decreases heart rate, acetylcholine is secreted and hyperpolarizes the heart
- *Sympathetic stimulation a positive chronotropic factor
 - *Supplied by cardiac nerves.
 - *Innervate the SA and AV nodes, and the atrial and ventricular myocardium.
 - *Increases heart rate and force of contraction.
 - *Epinephrine and norepinephrine released.
 - *Increased heart beat causes increased cardiac output.
 Increased force of contraction causes a lower end-systolic volume; heart empties to a greater extent. Limitations: heart has to have time to fill.

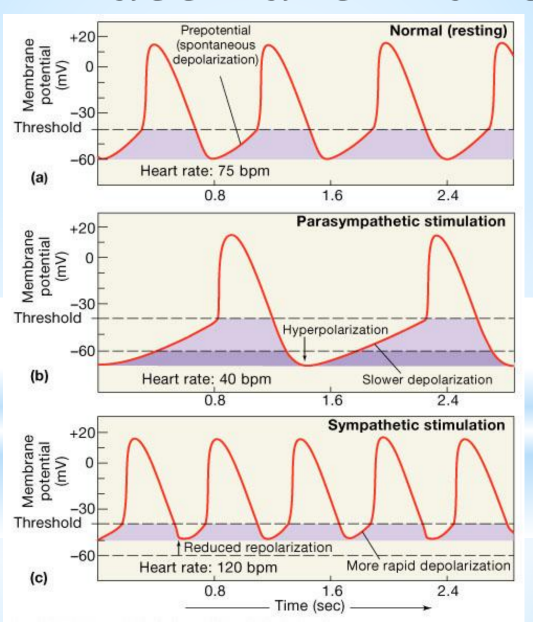
*Hormonal regulation

- * Epinephrine and norepinephrine from the adrenal medulla.
 - *Occurs in response to increased physical activity, emotional excitement, stress

- *SA node establishes baseline (sinus rhythmn)
- *Modified by ANS
- *If all ANS nerves to heart are cut, heart rate jumps to about 100 b/min
 - *What does this tell you about which part of the ANS is most dominant during normal period?

*Basic heart rate established by pacemaker cells

*Pacemaker Function



- *The hormones epinephrine and thyroxine increase heart rate
- *Intra- and extracellular ion concentrations must be maintained for normal heart function

*Chemical Regulation of the Heart

*Regulation of Stroke Volume

*SV: volume of blood pumped by a ventricle per beat

SV= end diastolic volume (EDV) minus end systolic volume (ESV); SV = EDV - ESV

- *EDV = end diastolic volume
 - *amount of blood in a ventricle at end of diastole
- *ESV = end systolic volume
 - *amount of blood remaining in a ventricle after contraction
- *Ejection Fraction % of EDV that is pumped by the ventricle; important clinical parameter
 - *Ejection fraction should be about 55-60% or higher

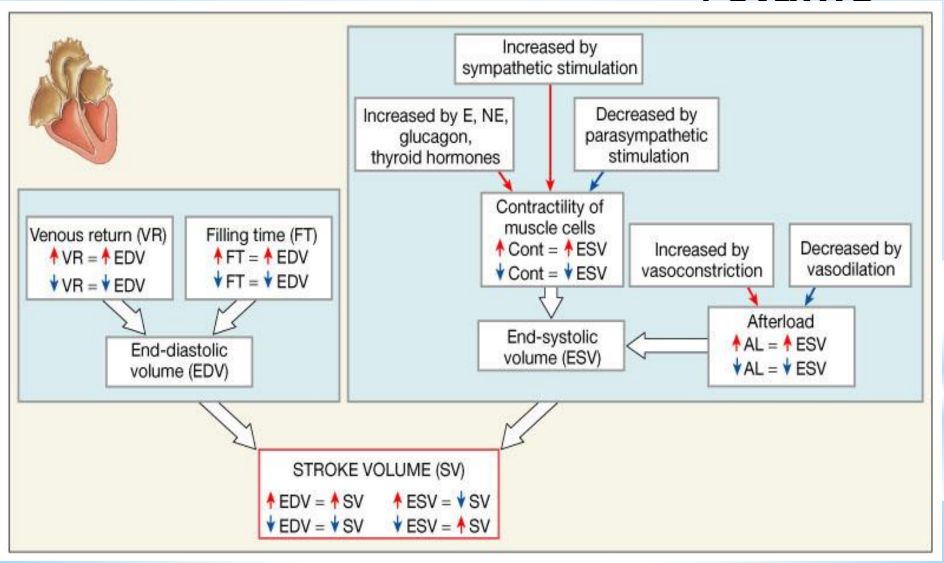
*Factors Affecting Stroke Cted by Volume

- *EDV affected by
 - *Venous return vol. of blood returning to heart
 - *Preload amount ventricles are stretched by blood (=EDV)
- *ESV affected by
 - *Contractility myocardial contractile force due to factors other than EDV
 - *Afterload back pressure exerted by blood in the large arteries leaving the heart

*Frank-Starling Law of the

- *Preload, or degree of stretch, of cardiac muscle elite before they contract is the critical factor controlling stroke volume; ↑EDV leads to ↑stretch of myocard.
 - *\force of contraction $\rightarrow \uparrow$ SV
 - *Unlike skeletal fibers, cardiac fibers contract MORE FORCEFULLY when stretched thus ejecting MORE BLOOD (↑SV)
 - *If SV is increased, then ESV is decreased!!
- *Slow heartbeat and exercise increase venous return (VR) to the heart, increasing SV
 - *VR changes in response to blood volume, skeletal muscle activity, alterations in cardiac output
 - $^*\uparrow VR \rightarrow \uparrow EDV$ and $\downarrow in VR \rightarrow \downarrow in EDV$
 - *Any \downarrow in EDV $\rightarrow \downarrow$ in SV
- *Blood loss and extremely rapid heartheat decrease SV

*Factors Affecting Stroke Volume



- *Contractility is the increase in contractile strength, independent of stretch and EDV
- *Referred to as extrinsic since the influencing factor is from some external source
- * Increase in contractility comes from:
 - *Increased sympathetic stimuli
 - *Certain hormones
 - *Ca²⁺ and some drugs
- *Agents/factors that decrease contractility include:
 - *Acidosis
 - *Increased extracellular K⁺
 - *Calcium channel blockers

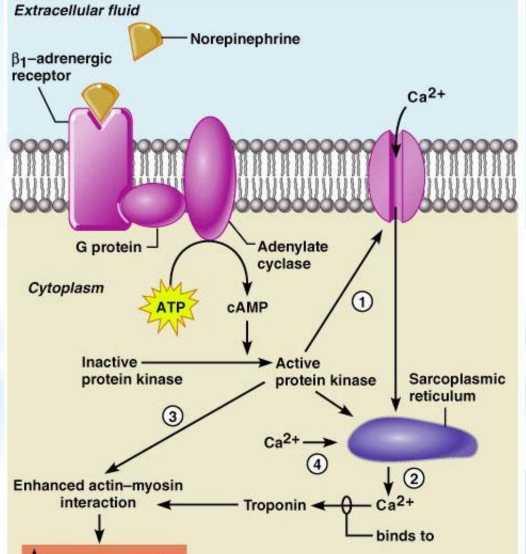
*Extrinsic Factors Influencing Stroke Volume

*Effects of Autonomic Activity on Contractility

- *Sympathetic stimulation
 - *Release norepinephrine from symp. postganglionic fiber
 - *Also, EP and NE from adrenal medulla
 - *Have positive ionotropic effect
 - *Ventricles contract more forcefully, increasing SV, increasing ejection fraction and decreasing ESV
- *Parasympathetic stimulation via Vagus Nerve -CNX
 - *Releases ACh
 - *Has a negative inotropic effect
 - *Hyperpolarization and inhibition
 - *Force of contractions is reduced, ejection fraction

*Sympathetic stimulation releases norepinephrine and initiates a cyclic AMP 2nd-messenger system

*Contractility and Norepinephrine



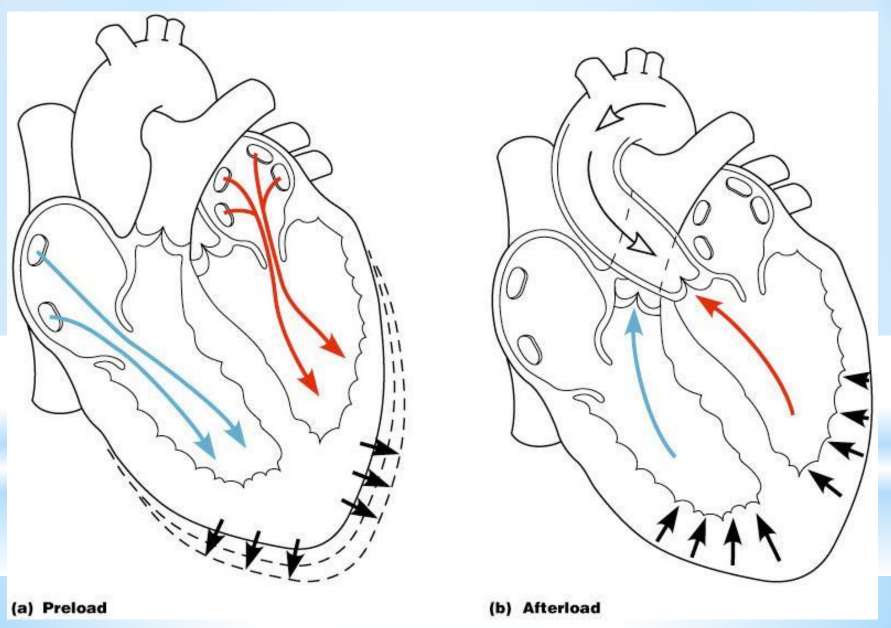


Figure 18.21

*Effects of Hormones on Contractility

- *Epi, NE, and Thyroxine all have positive ionotropic effects and thus †contractility
- *Digitalis elevates intracellular Ca⁺⁺ concentrations by interfering with its removal from sarcoplasm of cardiac cells
- *Beta-blockers (*propanolol*, *timolol*) block beta-receptors and prevent sympathetic stimulation of heart (neg. chronotropic effect)

- *Internet resources
- *Textbook of Marya Human phisiology

References