Heart Pump and Cardiac Cycle

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Objectives

- To understand the volume, mechanical, pressure and electrical changes during the cardiac cycle
- To understand the inter-relationship between all these changes
- To describe the factors that regulate Cardiac output and Stroke volume.
- Resources: Textbook of Medical Physiology By Guyton and Hall





Cardiac Cycle

- Cardiac cycle refers to all events associated with blood flow through the heart
 - Systole contraction of heart muscle
 - Diastole relaxation of heart muscle

Cardiac Cycle

- Atrial systole 0.1 second
- Atrial diastole 0.7 second
- Ventricular systole 0.3 second
 - Isovolumic contraction 0.01 seconds
 - Rapid ejection period
 - Slow ejection period
- Ventricular diastole 0.5 seconds
 - Isovolumic relaxation 0.02 seconds
 - Rapid filling
 - Slow filling (Diastasis)
 - Atrial contraction

Cardiac cycle ...cont

- End diastolic volume (EDV) End systolic volume (ESV) = Stroke volume (SV)
- SV X heart rate (HR) = cardiac output (CO)
- Ejection fraction = SV/EDV
- Inotropic vs. Chronotropic
- Autonomic control of cardiac cycle (pump)

- Ventricular filling mid-to-late diastole
 - Heart blood pressure is low as blood enters atria and flows into ventricles
 - AV valves are open, then atrial systole occurs

- Ventricular systole
 - Atria relax
 - Rising ventricular pressure results in closing of AV valves
 - Isovolumetric contraction phase
 - Ventricular ejection phase opens semilunar valves

- Isovolumetric relaxation early diastole
 - Ventricles relax
 - Backflow of blood in aorta and pulmonary trunk closes semilunar valves
- Dicrotic notch brief rise in aortic pressure caused by backflow of blood rebounding off semilunar valves





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Changes during Cardiac cycle

- Volume changes: End-diastolic volume, End-systolic volume, Stroke volume and Cardiac output.
- Aortic pressure: Diastolic pressure ~80 mmHg, Systolic pressure ~ 120 mmHg, most of systole ventricular pressure higher than aortic
- Ventricular pressure: Diastolic ~ 0, systolic Lt. ~120 Rt. ~ 25 mmHg.
- Atrial pressure: A wave =atrial systole, C wave= ventricular contraction (AV closure), V wave= ventricular diastole (Av opening)
- Heart sounds: S₁ = turbulence of blood around a closed AV valves, S₂ = turbulence of blood around a closed semilunar valves.

Heart Sounds



Heart Sounds

Heart sounds

 (lub-dup) are
 associated
 with closing
 of heart
 valves

Sounds of aortic semilunar valve are heard in 2nd intercostal space at right sternal margin Sounds of pulmonary semilunar valve are heard in 2nd intercostal space at left sternal margin Sounds of mitral valve are heard over heart apex, in 5th intercostal space in line with middle of clavicle Sounds of tricuspid valve are typically heard in

right sternal margin of

5th intercostal space; variations include over sternum or over left sternal margin in 5th

intercostal space

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Heart sounds

- Auscultation listening to heart sound via stethoscope
- Four heart sounds
 - S_1 -"lubb" caused by the closing of the AV values
 - S_2 "dupp" caused by the closing of the semilunar valves
 - S_3 a faint sound associated with blood flowing into the ventricles
 - S_4 another faint sound associated with atrial contraction

Cardiac Output (CO) and 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)
- 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

Cardiac Output: Example

CO (ml/min) = HR (75 beats/min) x SV (70 ml/beat) CO = 5250 ml/min (5.25 L/min)

Regulation of Stroke Volume

- SV = end diastolic volume (EDV) minus end systolic volume (ESV)
- EDV = amount of blood collected in a ventricle during diastole
- ESV = amount of blood remaining in a ventricle after contraction

Factors Affecting Stroke Volume

- Preload amount ventricles are stretched by contained blood
- Contractility cardiac cell 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 Heart

- Preload, or degree of stretch, of cardiac muscle cells before they contract is the critical factor controlling stroke volume
- Slow heartbeat and exercise increase venous return to the heart, increasing SV
- Blood loss and extremely rapid heartbeat decrease SV





(a) Preload

(b) Afterload

Cardiac Output





Extrinsic Factors Influencing Stroke Volume

- Contractility is the increase in contractile strength, independent of stretch and EDV
- Increase in contractility comes from:
 - Increased sympathetic stimuli
 - Certain hormones
 - Ca^{2+} and some drugs

Extrinsic Factors Influencing Stroke Volume

- Agents/factors that decrease contractility include:
 - Acidosis
 - Increased extracellular K⁺
 - Calcium channel blockers

Contractility and Norepinephrine

• Sympathetic stimulation releases norepinephrine and initiates a cyclic AMP secondmessenger system





LEFT VENTRICULAR PRESSURE/VOLUME P/V LOOP

LEFT VENTRICULAR PRESSURE (mmHg)



Valvular Function

- ≻ To prevent back-flow.
- Chordae tendineae are attached to A-V valves.
- Papillary muscle, attached to chordae tendineae, contract during systole and help prevent back-flow.
- Because of smaller opening, velocity through aortic and pulmonary valves exceed that through the A-V valves.

Valvular Function (cont'd)

Most work is external work or pressure-volume work.

A small amount of work is required to impart kinetic energy to the heart (1/2 mV²).
What is stroke-volume in previous figure?
External work is area of Pressure-Volume curve.
Work output is affected by "preload" (end-diastolic pressure) and "afterload" (aortic pressure).

Work Output of the Heart





Left Ventricular Volume



Left Ventricular Volume



Left Ventricular Volume

PRESSURE/VOLUME RELATIONSHIPS UNDER DIFFERENT CONDITIONS



Intrinsic Control of Stroke Volume (Frank-Starling Curve)



Regulation of Heart Rate

- Positive chronotropic factors increase heart rate
- Negative chronotropic factors decrease heart rate

Regulation of Heart Rate: Autonomic Nervous System

- Sympathetic nervous system (SNS) stimulation is activated by stress, anxiety, excitement, or exercise
- Parasympathetic nervous system (PNS) stimulation is mediated by acetylcholine and opposes the SNS
- PNS dominates the autonomic stimulation, slowing heart rate and causing vagal tone

Atrial (Bainbridge) Reflex

- Atrial (Bainbridge) reflex a sympathetic reflex initiated by increased blood in the atria
 - Causes stimulation of the SA node
 - Stimulates baroreceptors in the atria, causing increased SNS stimulation

Chemical Regulation of the Heart

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

Important Concepts About Cardiac Output (CO) Control

- Cardiac Output is the sum of all tissue flows and is affected by their regulation (CO = 5L/min, cardiac index = $3L/min/m^2$ (surface area in m²).
- CO is proportional to tissue O_{2.} use.
- CO is proportional to 1/TPR when AP is constant.
- CO = (MAP RAP) / TPR



Ventricular Pressure and Volume Curves (cont'd)

- During the latter part of the ejection phase how can blood still leave the ventricle if pressure is higher in the aorta? Momentum of blood flow
- Total energy of blood = $P + mV^2/2$

= pressure + kinetic energy

• Total energy of blood leaving ventricle is greater than in aorta.

Ejection Fraction

[©] End diastolic volume = 125 mlEnd systolic volume = 55 ml \mathbb{F} Ejection volume (stroke volume) = 70 ml $Figure E_{\text{jection fraction}} = 70 \text{ml}/125 \text{ml} = 56\%$ (normally 60%) FIF heart rate (HR) is 70 beats/minute, what is cardiac output? \bigcirc Cardiac output = HR * stroke volume = 70/min. * 70 ml= 4900 ml/min.

Ejection Fraction (cont'd)

- If HR =100, end diastolic volume = 180 ml, end systolic vol. = 20 ml, what is cardiac output?
- C.O. = 100/min. * 160 ml = 16,000 ml/min.
- Ejection fraction= 160/180%=~ 90%

Aortic Pressure Curve

- A ortic pressure starts <u>increasing</u> during systole after the <u>aortic valve opens.</u>
- Aortic pressure decreases toward the end of the ejection phase.
- After the aortic valve closes, an *incisura* occurs because of sudden cessation of back-flow toward left ventricle.
- Aortic pressure decreases slowly during diastole because of the elasticity of the aorta.

Frank-Starling Mechanism

→ Within physiological limits the heart pumps all the blood that comes to it without excessive damming in the veins.

Extra stretch on cardiac myocytes makes actin and myosin filaments interdigitate to a more optimal degree for force generation.

Ventricular Stroke Work Output



Autonomic Effects on Heart

- Sympathetic stimulation causes increased HR and increased contractility with HR = 180-200 and C.O. = 15-20 L/min.
- Parasympathetic stimulation decreases HR markedly and decreases cardiac contractility slightly. Vagal fibers go mainly to atria.
- Fast heart rate (tachycardia) can decrease C.O. because there is not enough time for heart to fill during diastole.

Effect of Sympathetic and Parasympathetic Stimulation on Cardiac Output



Right Atrial Pressure (mmHg)

Cardiac Contractility

- Best is to measure the C.O. curve, but this is nearly impossible in humans.
- dP/dt is not an accurate measure because this increases with increasing preload and afterload.
- (dP/dt)/P _{ventricle} is better. P _{ventricle} is instantaneous ventricular pressure.
- Excess K⁺ decreases contractility.
- Excess Ca⁺⁺ causes spastic contraction, and low Ca⁺⁺ causes cardiac dilation.

Thank You

