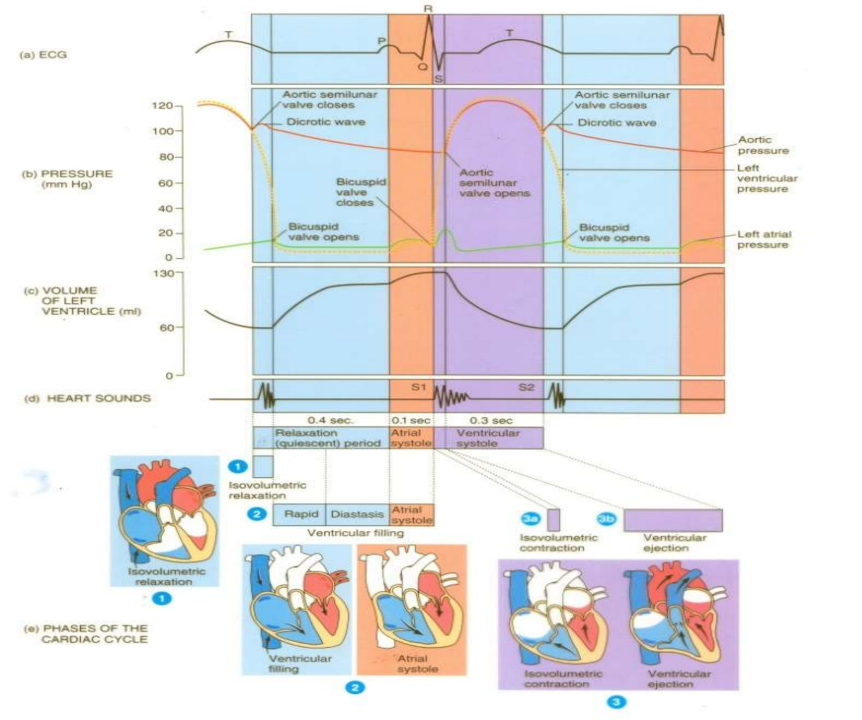
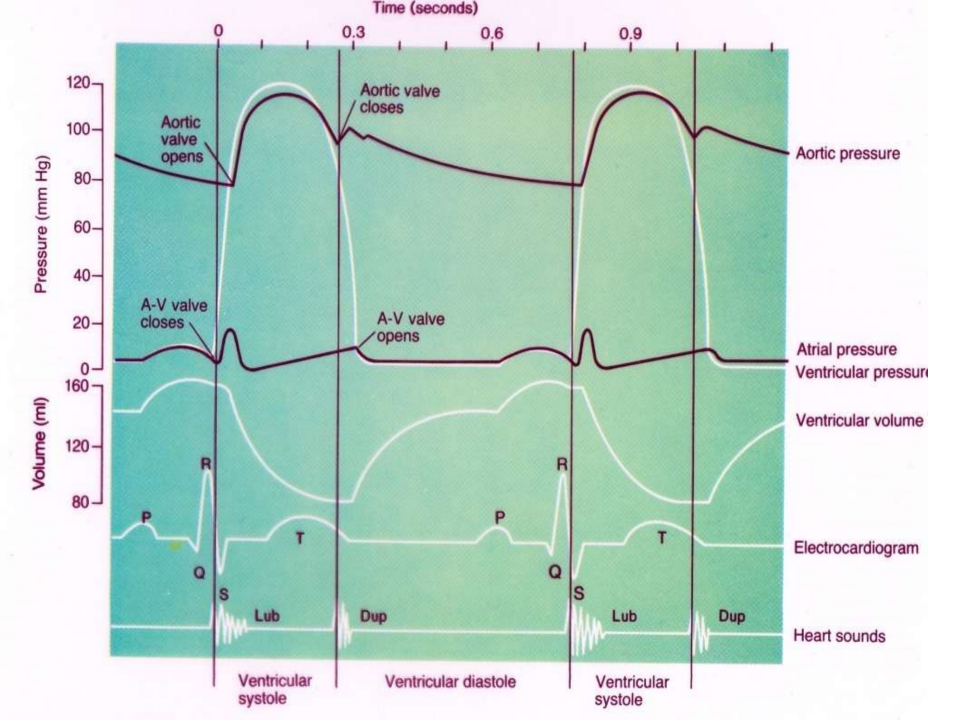
#### **Heart Pump and Cardiac Cycle**

Faisal I. Mohammed, MD, PhD

## **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

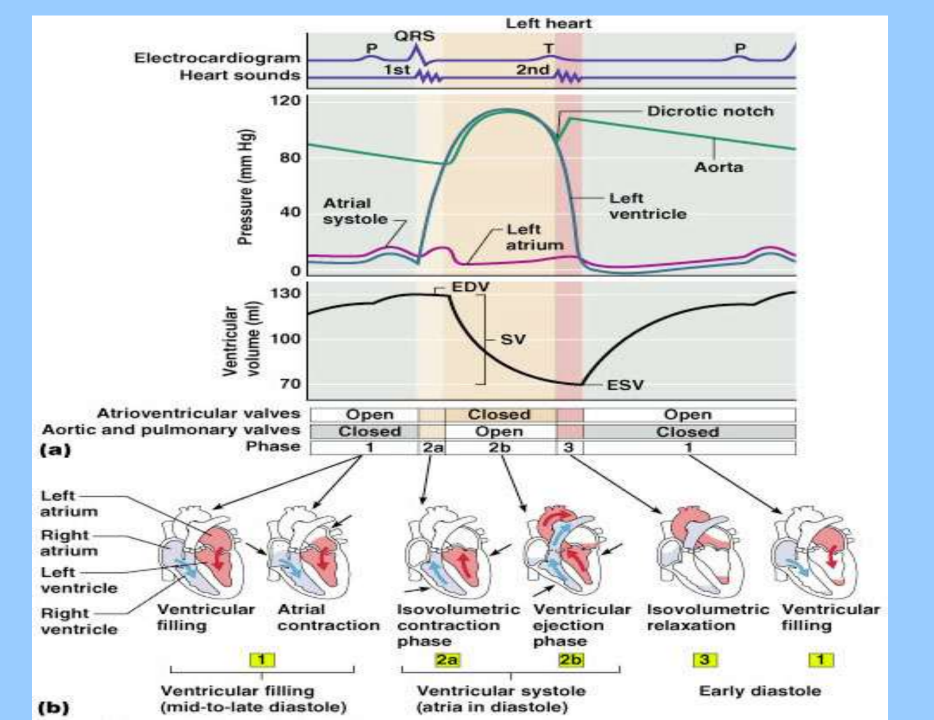
#### Cardiac cycle ...cont

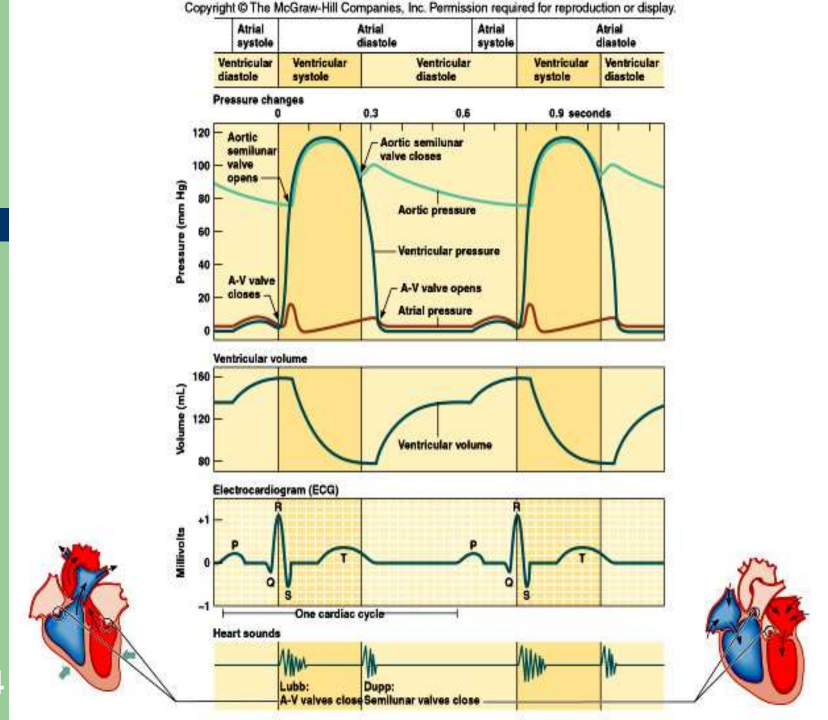
- End diastolic volume (EDV) End systolic volume (ESV) = Stroke volume (SV)
- SV X heart rate (HR) = cardiac output (CO)
- Ejection fraction = SV/EDV
- Inotropic vs. Chronotropic
- Autonomic control of cardiac cycle (pump)

- 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

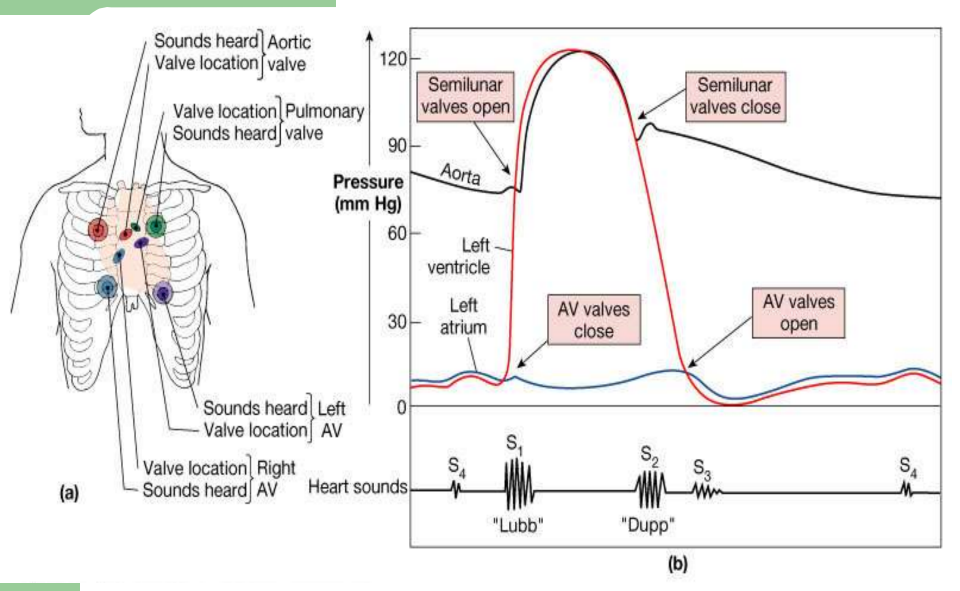




#### **Changes during Cardiac cycle**

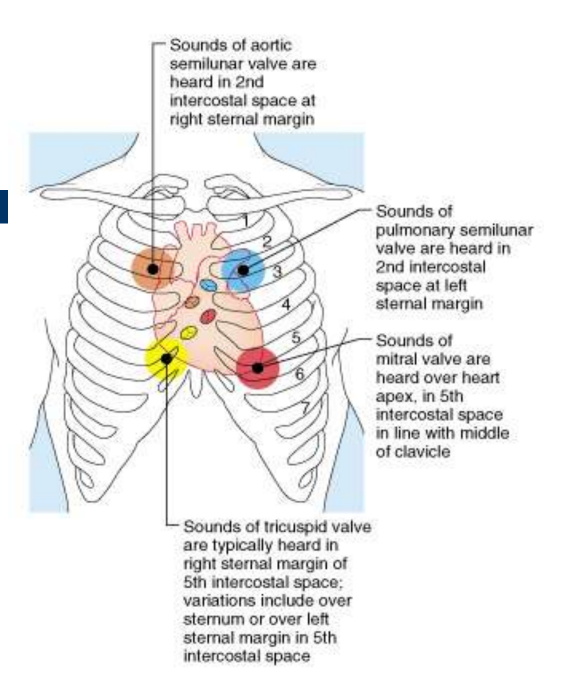
- Volume changes: End-diastolic volume, End-systolic volume,
   Stroke volume and Cardiac output.
- Aortic pressure: Diastolic pressure ~80 mmHg, Systolic pressure
   ~120 mmHg, most of systole ventricular pressure higher than aortic
- Ventricular pressure: Diastolic ~ 0, systolic Lt. ~120 Rt. ~ 25 mmHg.
- Atrial pressure: A wave =atrial systole, C wave= ventricular contraction (AV closure), V wave= ventricular diastole (Av opening)
- Heart sounds:  $S_1$  = turbulence of blood around a closed AV valves,  $S_2$  = turbulence of blood around a closed semilunar valves.

#### **Heart Sounds**



#### **Heart Sounds**

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



#### **Heart sounds**

- Auscultation listening to heart sound via stethoscope
- Four heart sounds
  - $-S_1$  "lubb" caused by the closing of the AV valves
  - S<sub>2</sub> "dupp" caused by the closing of the semilunar valves
  - $-S_3$  a faint sound associated with blood flowing into the ventricles
  - S<sub>4</sub> 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

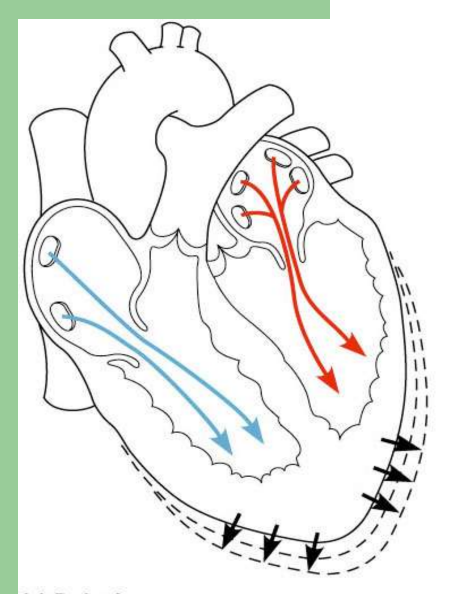
- SV = end diastolic volume (EDV)
   minus end systolic volume (ESV)
- EDV = amount of blood collected in a ventricle during diastole
- ESV = amount of blood remaining in a ventricle after contraction

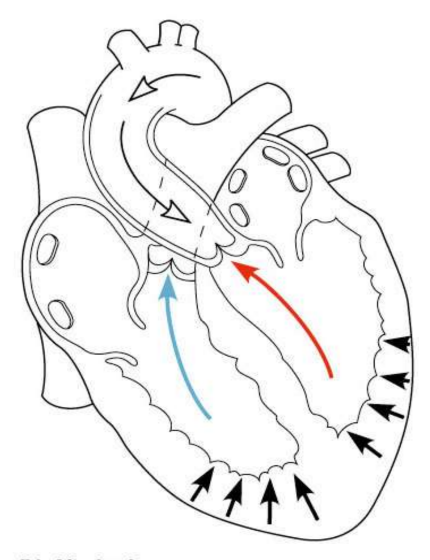
## **Factors Affecting Stroke Volume**

- 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

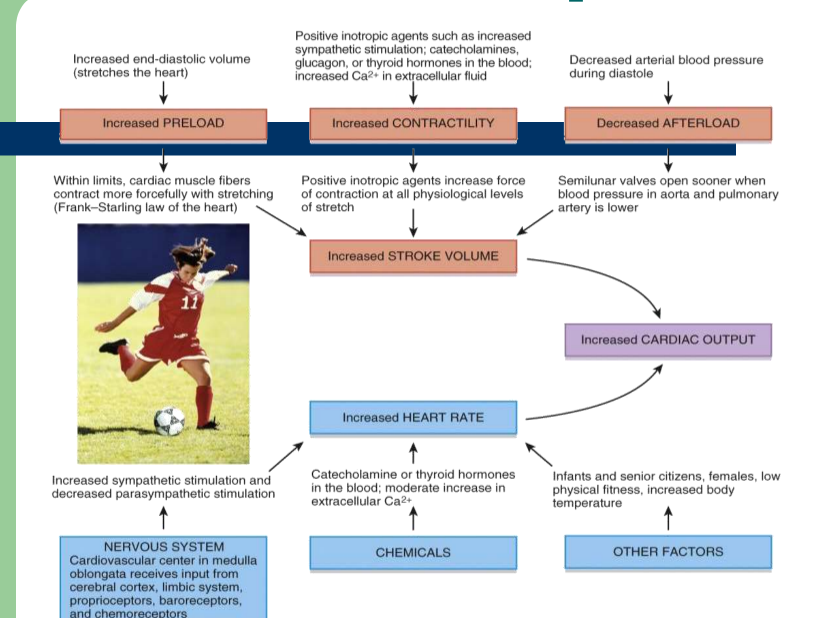


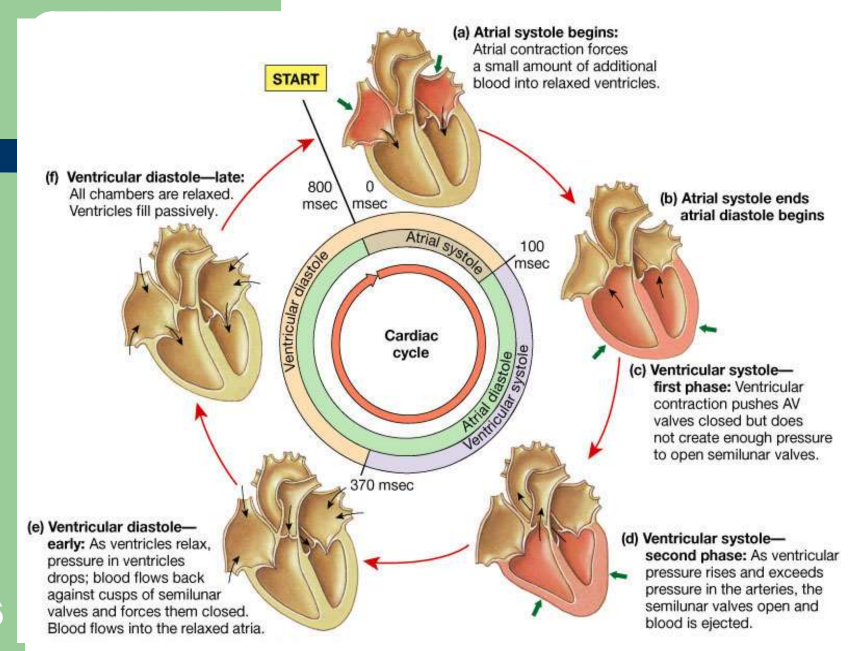


(a) Preload

(b) Afterload

### **Cardiac Output**





# **Extrinsic Factors Influencing Stroke Volume**

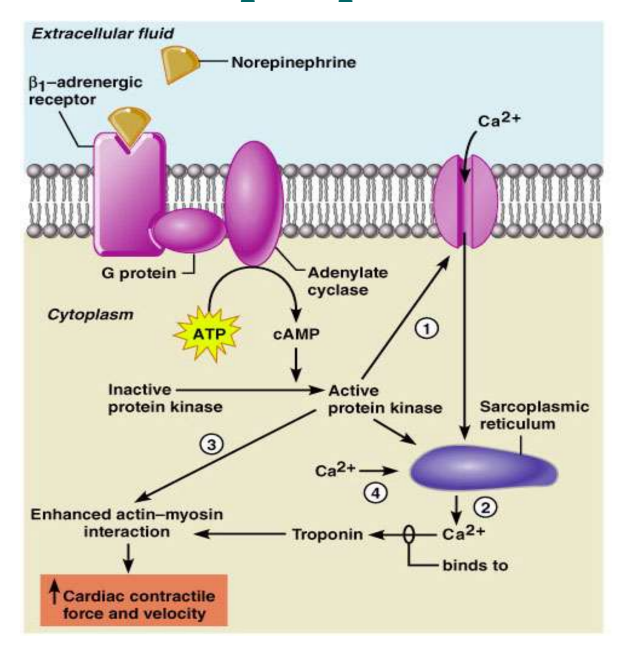
- Contractility is the increase in contractile strength, independent of stretch and EDV
- Increase in contractility comes from:
  - Increased sympathetic stimuli
  - Certain hormones
  - Ca<sup>2+</sup> and some drugs

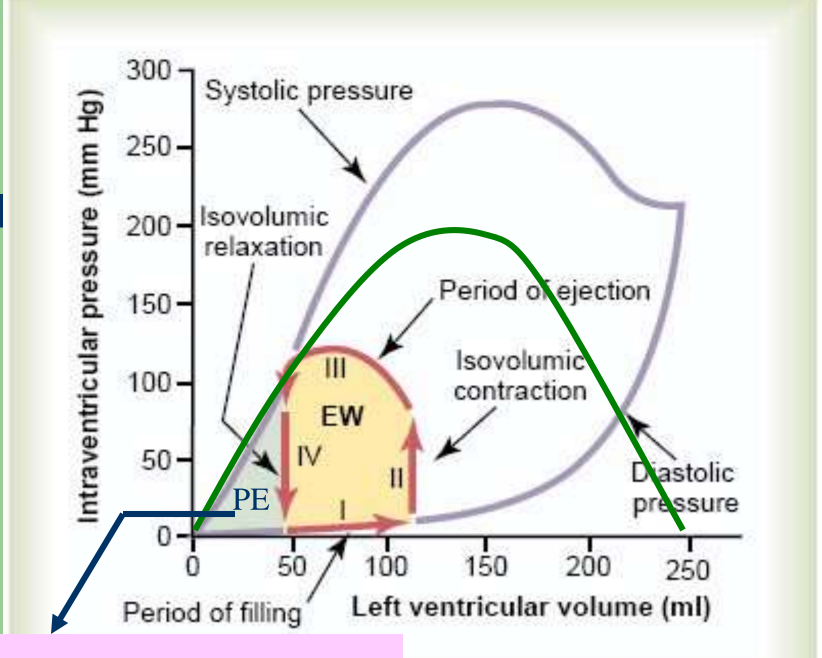
# **Extrinsic Factors Influencing Stroke Volume**

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

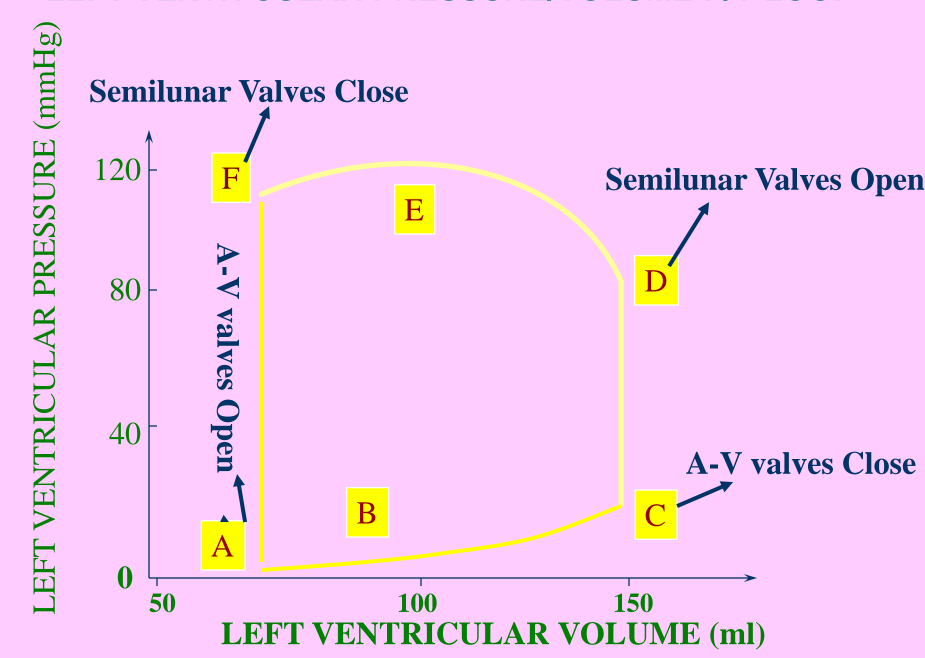
## Contractility and Norepinephrine

Sympathetic stimulation releases norepinephrine and initiates a cyclic AMP secondmessenger system





#### LEFT VENTRICULAR PRESSURE/VOLUME P/V LOOP



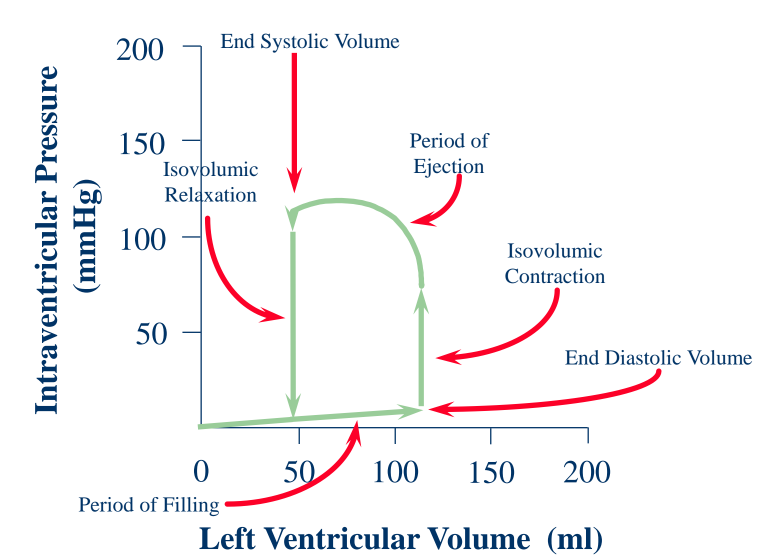
#### **Valvular Function**

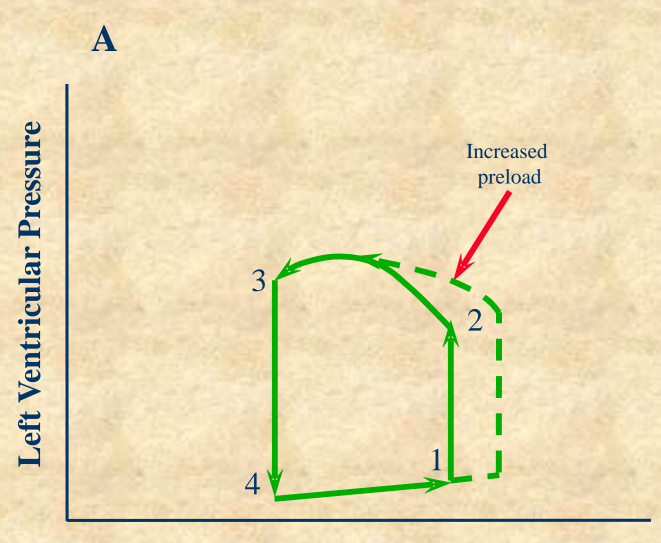
- To prevent back-flow.
- > Chordae tendineae are attached to A-V valves.
- Papillary muscle, attached to chordae tendineae, contract during systole and help prevent back-flow.
- ➤ Because of smaller opening, velocity through aortic and pulmonary valves exceed that through the A-V valves.

#### Valvular Function (cont'd)

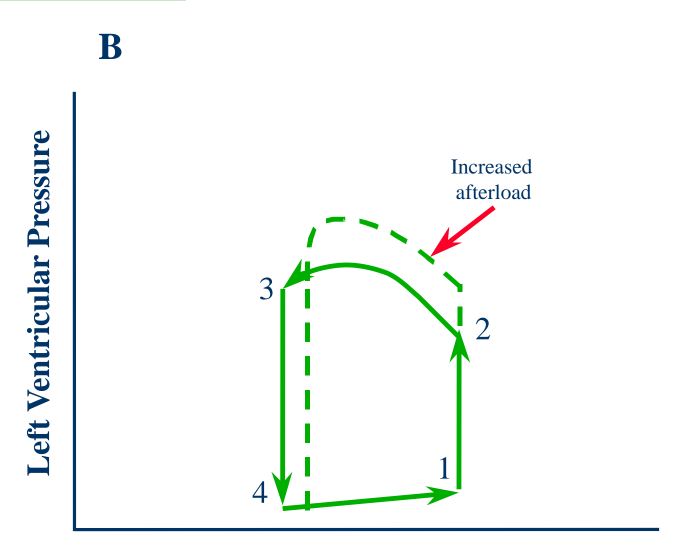
- Most work is external work or pressure-volume work.
- A small amount of work is required to impart kinetic energy to the heart (1/2 mV²).
- ☼ What is stroke-volume in previous figure?
- External work is area of Pressure-Volume curve.
- ₩ Work output is affected by "preload" (end-diastolic pressure)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

#### **Work Output of the Heart**

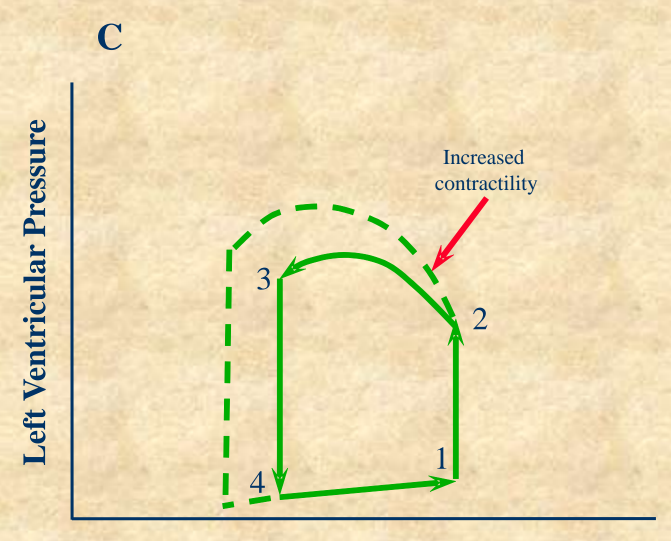




**Left Ventricular Volume** 

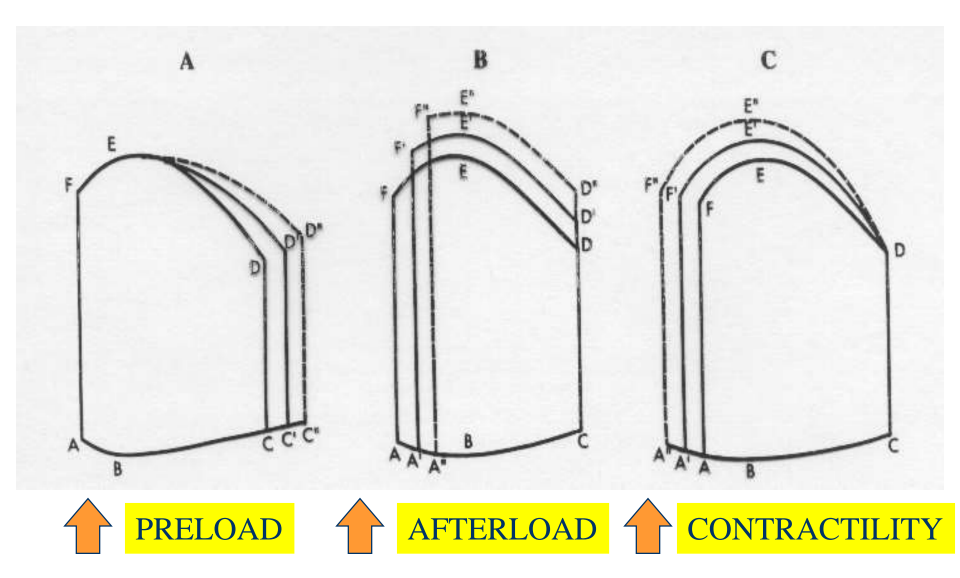


**Left Ventricular Volume** 

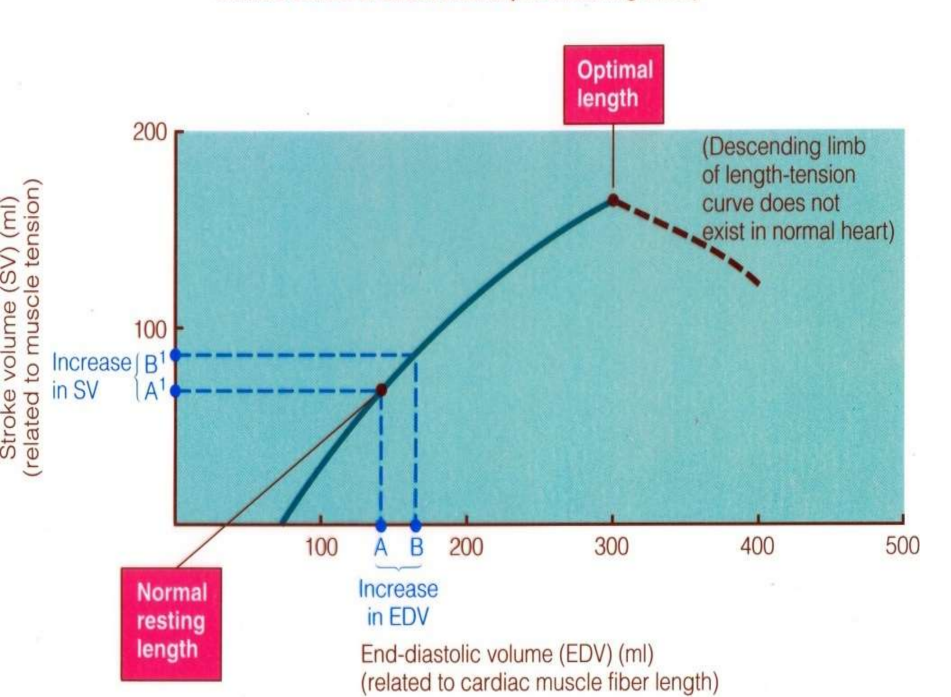


**Left Ventricular Volume** 

# 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

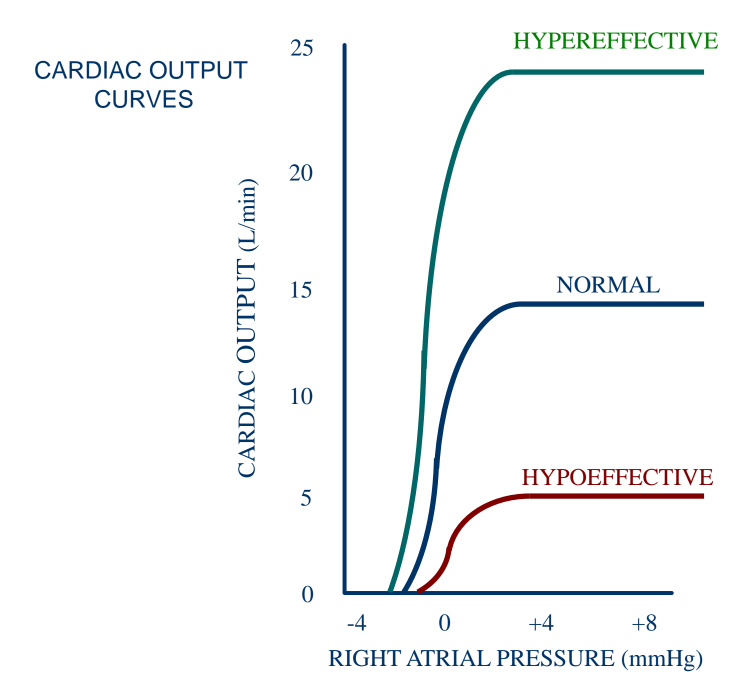
- Atrial (Bainbridge) reflex a sympathetic reflex initiated by increased blood in the atria
  - Causes stimulation of the SA node
  - Stimulates baroreceptors in the atria,
     causing increased SNS stimulation

### **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  $m^2$ ).
- CO is proportional to tissue O<sub>2</sub> 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 \text{ ml}
End systolic volume
                                    = 55 ml
Ejection volume (stroke volume) = 70 ml
Ejection fraction = 70\text{ml}/125\text{ml} = 56\%
      (normally 60%)
If heart rate (HR) is 70 beats/minute, what is
      cardiac output?
Cardiac output = HR * stroke volume
                   = 70/\text{min.} * 70 \text{ ml}
                          = 4900 \text{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%

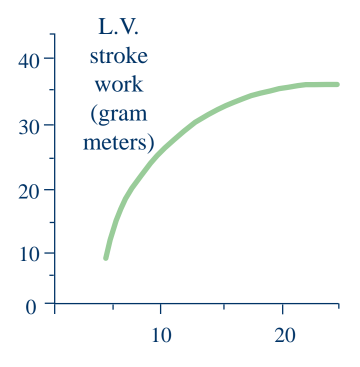
#### **Aortic Pressure Curve**

- ❖ Aortic 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.

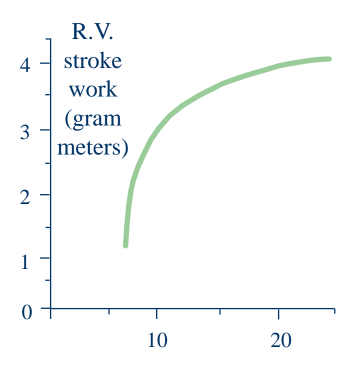
### Frank-Starling Mechanism

- → 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



Left Atrial Mean Pressure (mm Hg)

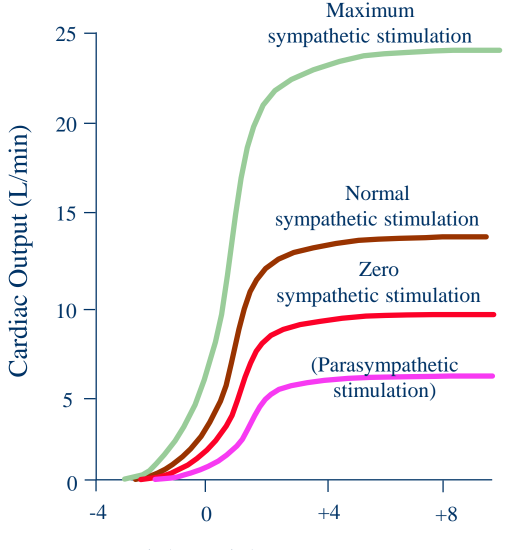


Right Atrial Mean Pressure (mm Hg)

#### **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 <sub>ventricle</sub> is better. P <sub>ventricle</sub> is instantaneous ventricular pressure.
- Excess K<sup>+</sup> decreases contractility.
- Excess Ca<sup>++</sup> causes spastic contraction, and low Ca<sup>++</sup> causes cardiac dilation.

# Thank You

