

CVS Block Stroke Volume Cardiac Output & its Regulation (Preload, Contractility & Afterload) (Physiology L No.6) Dr. Hayam Gad MBBS, MSc, PhD A. Professor of Physiology College of Medicine, KSU



جــامـعـة الملك سعود King Saud University

### **Learning Objectives**

- Define stroke volume, end-systolic volume, and end-diastolic volume
- Define cardiac output, venous return, cardiac index & cardiac reserve
- Understand the concept of preload and afterload
- Understand the determinants of CO and how CO is regulated
- Understand the factors affecting the EDV (the venous return)
- Understand the factors affecting the ESV
- Know how cardiac contractility & heart rate changes affect CO
- Identify the factors that affect heart rate
- Know the method for measurement of CO (The direct Fick's method)
- Describe the factors affecting the SV & CO

**Remember What Are P End-diastolic volume (EDV)**:

Volume of blood in ventricles at the end of diastole  $\approx 110-130 \text{mL}$ 

### End-systolic volume (ESV):

Amount of blood left in ventricles at the end of systole  $\approx 40-60mL$ 

### Stroke volume (SV):

Amount of blood ejected from ventricles during systole  $\approx$  70-80 mL/beat.

### **Dejection fraction (EF):**

Fraction of EDV ejected during a heart beat = $SV/EDV \approx 60-65\%$ .











If the amount of blood pumped by the left ventricle/<u>beat</u> = stroke volume (~70ml/beat, in adult at rest).

The Cardiac Output

• What is the amount of blood pumped by the left ventricle / minute? = (The Cardiac Output)

> Cardiac output (CO) = Stroke volume (SV) X heart rate (HR) ≈ 5L/min. in adult at rest

The Cardiac Output (CO)

- It is the volume of blood pumped by each ventricle per minute ≈ 5L/min.
- Since the normal adult blood volume is about <u>5 L</u>, the entire blood supply passes through the body once each minute.



## Normal Values of CO at Rest

## • Resting CO:

- The average CO for the resting adult is 5L/min.
- For men, CO **≈ 5.6L/min.**
- For women, CO ≈
   4.9L/min.

### • The factors affecting CO:

- 1. Body metabolism
- 2. Pregnancy
- 3. Body temperature.
- 4. Sympathetic activity
- 5. Exercise
- 6. Hyperthyroidism

## Physiological Changes in CO

During the first 3 hours after meals,  $\uparrow$  CO by  $\approx$  30% to enhance blood flow in the intestinal circulation.

- Later months of pregnancy ↑ CO by ≈ 30% due to increased uterine blood flow.
- > At environmental temperature above 30°C , $\uparrow$  CO due to  $\uparrow$  skin blood flow.
- During anxiety and excitement, sympathetic activity \u0357 CO up to 50% 100%.
- Sitting or standing from the lying position \$\\$ CO by 20-30%.
  Exercise \$\\$

## Effects of Exercise on Heart Rate, SV and CO

### Moderate Exercise

- ↑ HR ≈ 200% of resting, (140 beats/min.) ↑ SV ≈ 120% (85 ml)
- $\uparrow$  CO  $\approx$  240% (12 L)

#### Severe Exercise

- ↑ HR ≈ 300% of resting (200 beats/min.)
  ↑ SV ≈ 175% (125 ml)
- ↑ CO ≈ 500% 700% (25 35 L)



In athletes, maximum CO may be 35L or more - can't increase maximum HR beyond 200 beats/min. Hence, the SV  $\uparrow$  to 175 ml.

### What is the Cardiac Index (CI) 🦓

- Since : CO vary with size of individual, age & gender (e.g: women have smaller CO than men, children have smaller CO than adults).
- Thus it is important to relate heart performance to the size of the individual.

Cardiac index is CO per square meter of body surface area  $CI = CO/m^2$ Normal CI = (3.2 Liters/min/m<sup>2</sup> body surface area) What is the Cardiac Reserve?



Cardiac reserve = The maximum volume of blood that the heart is capable of pumping/min — The resting CO at rest

### **Preload**

- It is the **amount of blood** that returns to the heart from veins (VR).
- It is the load on the muscle in the relaxed state.
- ↑VR→ ↑EDV and stretches or lengthens the ventricular muscle fibers.

### **Preload** ↑ in:

- Hypervolemia
- Heart failure



What Are Preload and Afterload

### Afterload

- It is the **resistance** against which the ventricles contract.
- Afterload on the right ventricle \u03c6, when pulmonary artery pressure \u03c6

### Afterload ↑ in:

- Aortic/pulmonary stenosis
- Hypertension
- Vasoconstriction



- CO is crucial since it is the amount of blood that flows into the circulation and is responsible for transporting substances to and from the tissues.
- Thus, the body has strict control mechanisms that maintain adequate CO.
- There are 2 major factors which determine CO.
- These are the SV and HR.



To increase cardiac output

Increase stroke volume or

Increase heart rate or increase both







- *EDV* = is the volume of blood in the ventricles prior to ventricular ejection.
- How does the EDV affect the SV and hence the CO?

↑↑ The stretch in the myocardium(↑↑ initial fiber length)

**↑**↑ EDV

↑↑ Myocardial contractility (↑↑ Strength of contraction) Frank-Starling mechanism

## Frank-Starling's Law

- It is the intrinsic relationship between EDV and SV.
- It reflects the ability of the heart to change its force of contraction and therefore SV in response to changes in VR.
- If EDV (preload) is increased, the ventricular fiber length is also increased, resulting in an increased 'tension' of the muscle (i.e SV increases in response to increase of the EDV).



Significance: Frank-Starling's Law

- The greater the stretch of the cardiac muscle the greater would be the force of contraction. i.e.: the energy of contraction is proportional to the initial length of the cardiac muscle fibers (the EDV).
- Why: During stretch, actin & myosin filaments are brought to more optimal degree of sliding, therefore increase force of contraction.
- Within physiologic limits, the heart pumps all the blood that returns to it by the way of the veins.



## Factors affecting preload (EDV)

- **EDV** ↑ with:
  - □ Increased total blood volume.
  - □ Increased venous return.
  - □ Increased venous tone.
  - □ Increased skeletal muscle pump (exercise).
  - □ Increased negative intrathoracic pressure.
  - □ Stronger atrial contraction.

Preload and Afterload

- EDV↓ with:
  - Standing.
  - Decreased venous return.
  - □ Increased intrapericardial pressure.
  - **Decreased ventricular compliance.**

- Indices of left ventricular preload: • Left ventricular end-diastolic volume (LVEDV).
- Left ventricular end-diastolic pressure (LVEDP).



End-Systolic Volume (ESV)

- *ESV* = The volume of blood remaining in the ventricle at the end of systole.
- $\uparrow$  ESV  $\rightarrow \downarrow$  stroke volume
- $\downarrow \text{ESV} \rightarrow \uparrow \text{stroke volume}$
- <u>ESV is determined by:</u>
   1. Cardiac contractility
   2. Afterload



## Determinants of ESV

### **<u>1- Cardiac contractility</u>**.

- $\uparrow \uparrow$  contractility  $\rightarrow \downarrow \downarrow ESV \rightarrow \uparrow \uparrow SV \& CO$
- $\downarrow \downarrow$  contractility  $\rightarrow \uparrow \uparrow ESV \rightarrow \downarrow \downarrow SV \& CO$

Variation in the Contractility of the Heart



(Preload) Stretching of the Myocardium

## Determinants of ESV....Cont.

### 2- Afterload:

- It is expressed as tension which must be developed in the wall of ventricles during systole, i.e the load the heart needs to overcome to open the semilunar valves and eject blood to aorta/pulmunary artery.
- Left ventricular afterload represents the force that the muscle must generate to eject the blood into the aorta.
- Right ventricular afterload represents the force that the muscle must generate to eject the blood into pulmonary artery.



## Determinants of ESV....Cont.

- When the aortic pressure (afterload) increases, the velocity of shortening of the LV myocardial fibers ↓. Hence, the LV eject less blood → ↓ SV →↑ ESV.
- The opposite is true when LV afterload is reduced.

#### increased afterload = reduced contraction



## Factors affecting Afterload

- Afterload  $\uparrow$  by any factor that restricts blood flow
  - •Vasoconstriction **\** the peripheral vascular resistance  $\rightarrow \uparrow \uparrow ESV \rightarrow \downarrow \downarrow SV \& CO$
  - •Vasodilatation  $\downarrow \downarrow$  the peripheral vascular resistance  $\rightarrow$  $\downarrow \downarrow ESV \rightarrow \uparrow \uparrow SV \& CO$
- Causes of 1 left ventricular afterload:
  - 1- Aortic stenosis
  - 2- Arterial hypertension.
  - 3- Vasoconstriction (the peripheral vascular resistance).
- Causes of ↑ right ventricular afterload:
  - 1- Pulmonary stenosis
  - 2- Pulmonary hypertension.
  - 3- Pulmonary vasoconstriction.



Normal arteriolar tone

#### Vasoconstriction



(increased contraction of circular smooth muscle in the arteriolar wall, which leads to increased resistance and decreased low through the vessel

#### Vasodilation

(decreased contraction of circular smooth muscle in the arteriol wall which leads to decreased resistance and increased flow through the vessel





Explanation: an increase in afterload decreases the velocity of fiber shortening. This reduces the rate of volume ejection so that more blood is left within the ventricle at the end of systole  $\rightarrow \uparrow$  ESV.

Summary of the Factors Influencing Stroke Volume





## Inotropic, Chronotropic & Dromotropic

### CONTRACTILITY

- Positive inotropic agent =  $\uparrow$  in contractility.
- Negative inotropic agent =  $\downarrow$  in contractility

### HEART RATE

- Positive chronotropic effect =  $\uparrow$  in HR
- Negative chronotropic effect =  $\downarrow$  in HR

### **CONDUCTION VELOCITY**

- Positive dromotropic effect =  $\uparrow$  in conduction velocity
- Negative dromotropic effect =  $\downarrow$  in conduction velocity

## The Heart Rate (HR)

- Normal heart rate is regular sinus rhythm= 60-100 beats/min
- > 100 beats/min  $\rightarrow$  Tachycardia
- < 60 beats/min  $\rightarrow$  Bradycardia
- Since the CO = SV X HR,  $\uparrow$  heart rate  $\rightarrow \uparrow$  CO.
- At heart rates up to about 180, ventricular filling is adequate as long as there is enough venous return,
- However, at very high heart rates, filling may be compromised to such a degree that CO falls.



## Frequency-force relationship

 The heart rate has an influence on cardiac contractility as well (frequencyforce relationship).

Increasing heart rate increases contractility.

- This frequency dependency of force generation is probably due to 

   Ca<sup>2+</sup>
   availability to the contractile proteins within cardiac myocytes as a result
   of:
  - The ↑No. of depolarizations/min → more frequent plateau phases → more Ca<sup>2+</sup> entry.
  - ↑ The magnitude of Ca<sup>2+</sup> current
     → ↑ the intracellular Ca<sup>2+</sup> stores.
  - Both effects enhance Ca<sup>2+</sup> release & uptake by the sarcoplasmic reticulum.



## Regulation of Heart Rate

#### 1- Autonomic nervous system

- Sympathetic nerves increase HR (and the contractility) through noradrenaline release (or adrenaline), which stimulates heart β1- receptors. E.g during stress, crisis or low blood pressure
- Parasympathetic nerves (vagus nerve) slow HR (but has little inotropic action). Through Ach release that stimulates muscarinic (M2) receptors.







Under normal conditions the SA node is under continuous vagal tone causing its basal firing rate to decrease (Vagal Tone)

Regulation of Heart Rate.....Cont.

### 2- Physical factors:

- Age: Resting HR is faster in fetus and then gradually decreases throughout life.
- Gender: HR is faster in females (72-80 beats/min) than in males (64-72 beats/min).
- Temperature: Heat increases HR as occurs in high fever. Cold has the opposite effect.
- Exercise: Increases HR through sympathetic nervous system.

### 3- Hormones and drugs

- Epinephrine and thyroxin increase HR.
- Increased Ca<sup>2+</sup> level in blood causes prolonged contraction
- Reduced Ca<sup>2+</sup> level in blood depress the heart.

Regulation of Heart Rate.....Cont.

#### 4- Blood volume

- Atrial Reflex (Bainbridge reflex)
- This reflex adjusts heart rate in response to venous return.
- Increase blood volume, stimulates stretch receptors in right atrium.
- This triggers increase in heart rate through increased sympathetic activity.





## Regulation of Cardiac Output by Autonomic Nerves





## Summary of the Factors Affecting Cardiac Output



# Pathological Low Or High Cardiac

- Causes of low CO:
- Low VR e.g hemorrhage.
- **Reduced contractility e.g. heart failure**
- Tachyarrhythmias e.g. atrial fibrillation and ventricular tachycardia
- □ Marked bradycardia e.g., complete heart block.

Causes of high CO:

- □ Hyperthyroidism: the increase in the CO is due to the high metabolic rate → vasodilatation →↑ CO to 50% of control.
   □ AV fistulas.
- **Fever.**
- Anaemia.
- Anxiety.

## Measurement of Cardiac Output

13°C

I3°C

- 2-Dimensional Echocardiography
- Ultra-fast computer tomography
- Fick's principle

2-Dimensional Echocardiography

- Records real-time changes in ventricular dimensions during systole and diastole.
- It thus computes the SV, which when multiplied by HR, the CO will be calculated.



Ultra-fast Computer Tomography



Can measure changes in ventricular diameter at several depths to estimate changes in ventricular volume.





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**Fick's principle** 

It assumes that the amount of oxygen consumed = the amount of oxygen delivered by the arterial blood minus the amount of oxygen in venous blood of the organ.



# Measurement of Cardiac Output: Fick's method

 $CO (L/min) = \frac{Total O_2 consumption}{AO_2 - VO_2}$ 

 $AO_2$  = Arterial  $O_2$  concentration

 $VO_2$  = Mixed  $O_2$  venous concentration



