

# **CARDIAC MECHANICS**

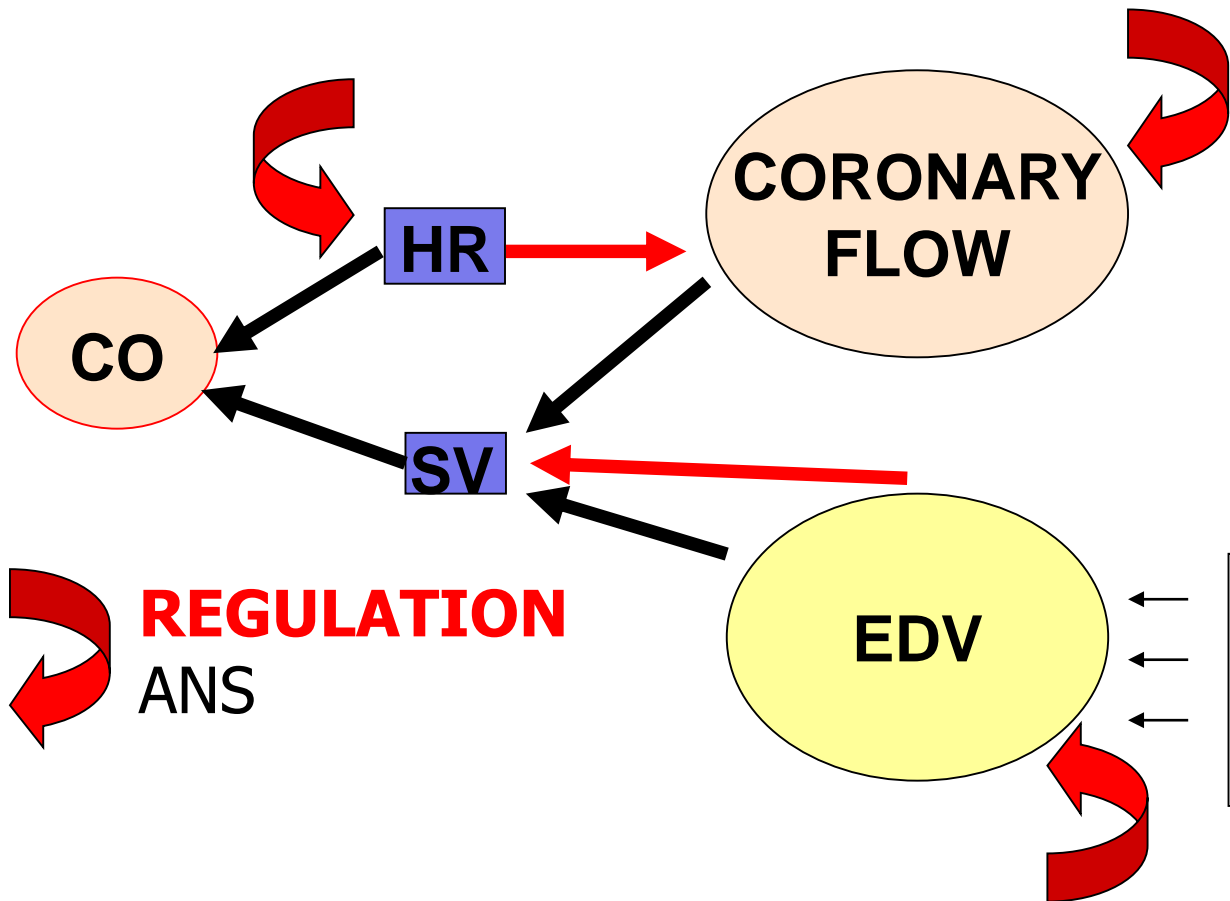
**HEART AS A PUMP**

**CARDIAC CYCLE**

**HEART FAILURE**

# CARDIAC OUTPUT (CO)

$$LV = RV$$



$$CO = HR \times SV \quad 5l/min$$

$$SV = EDV - ESV \quad 70ml$$

$$\text{Ejection fraction} \quad EF = \frac{EDV - ESV}{EDV} \quad >60\%$$

## CONTRACTILITY

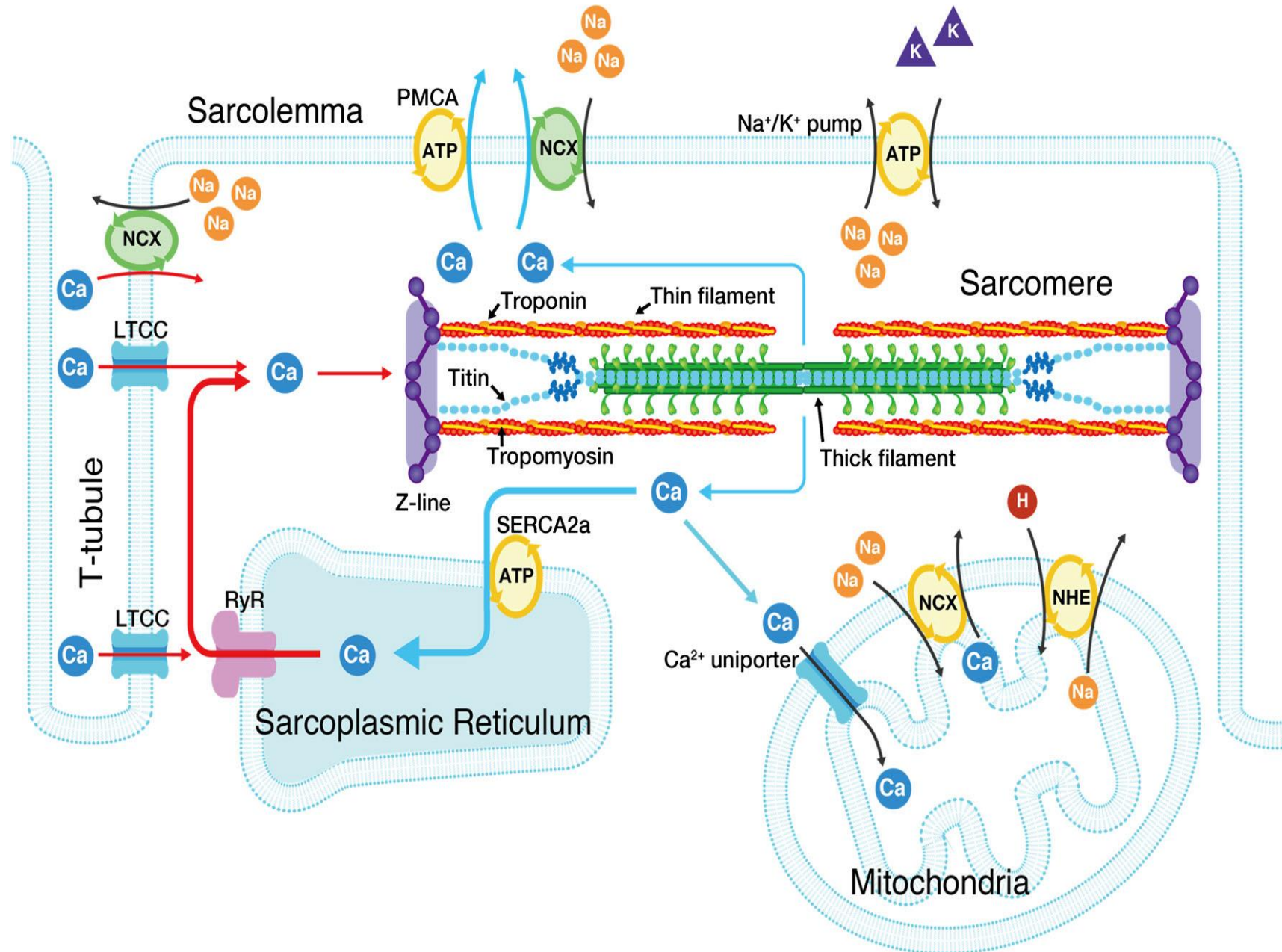
Ability to contract  
Depends on tissue perfusion  
(substrates and oxygen supply for  
ATP production;  $Ca^{2+}$  availability)

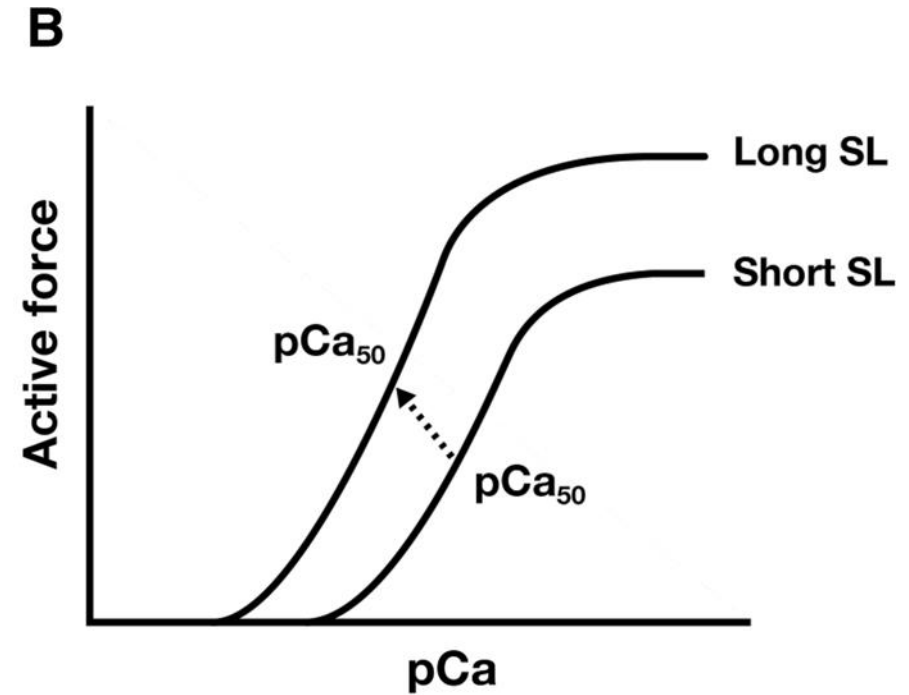
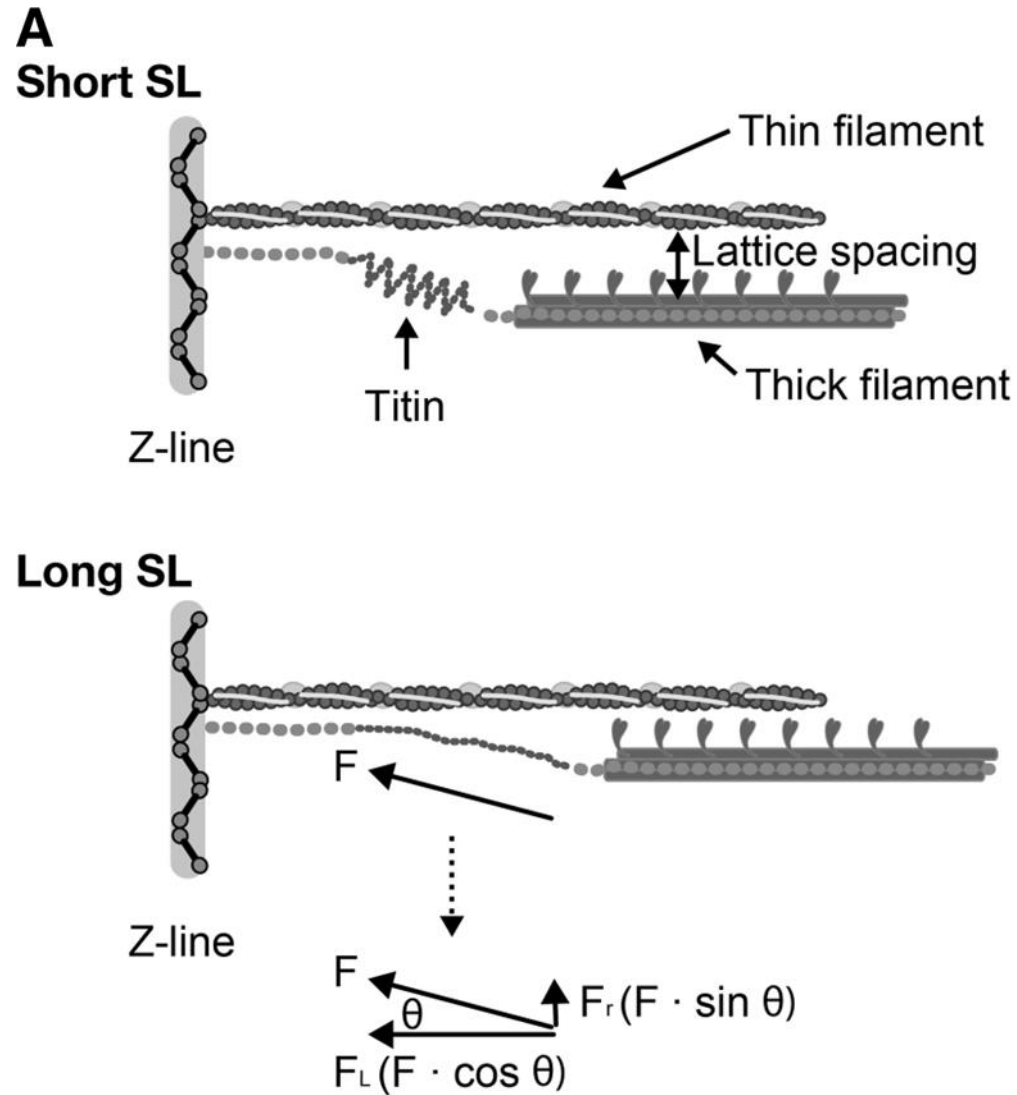
## AUTOREGULATION of cardiac contraction

Heterometric: Starling law

Homeometric: Frequency effect

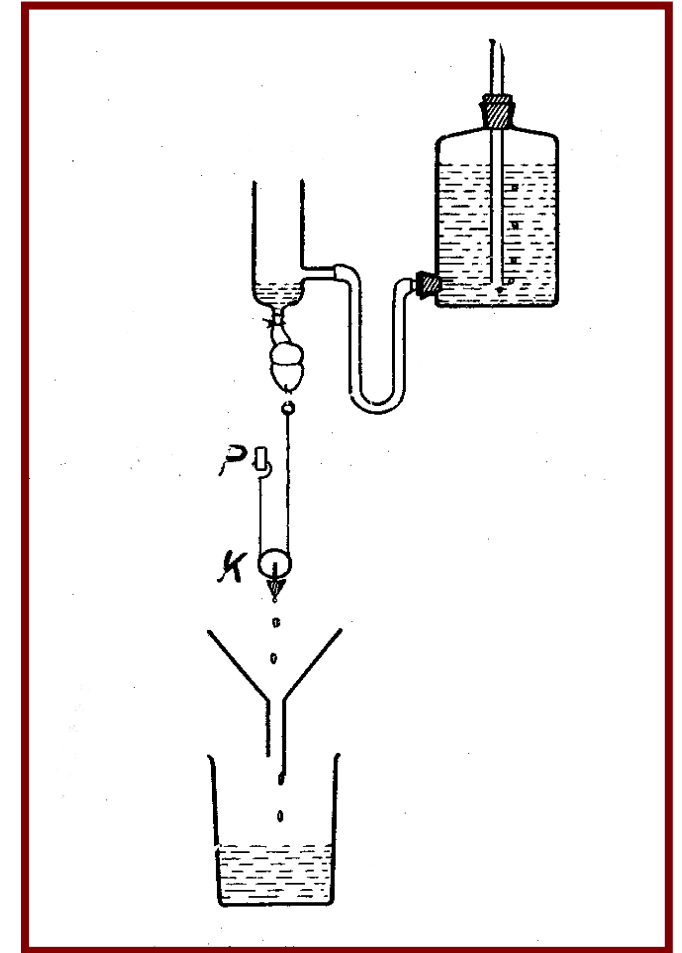
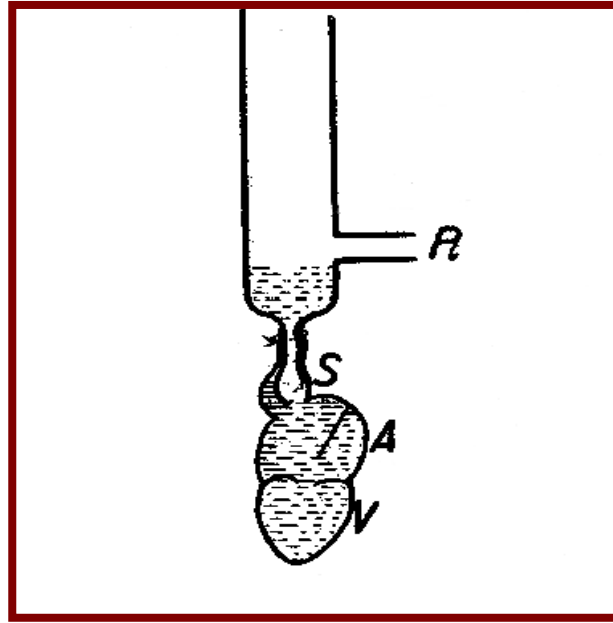
# STARLING LAW







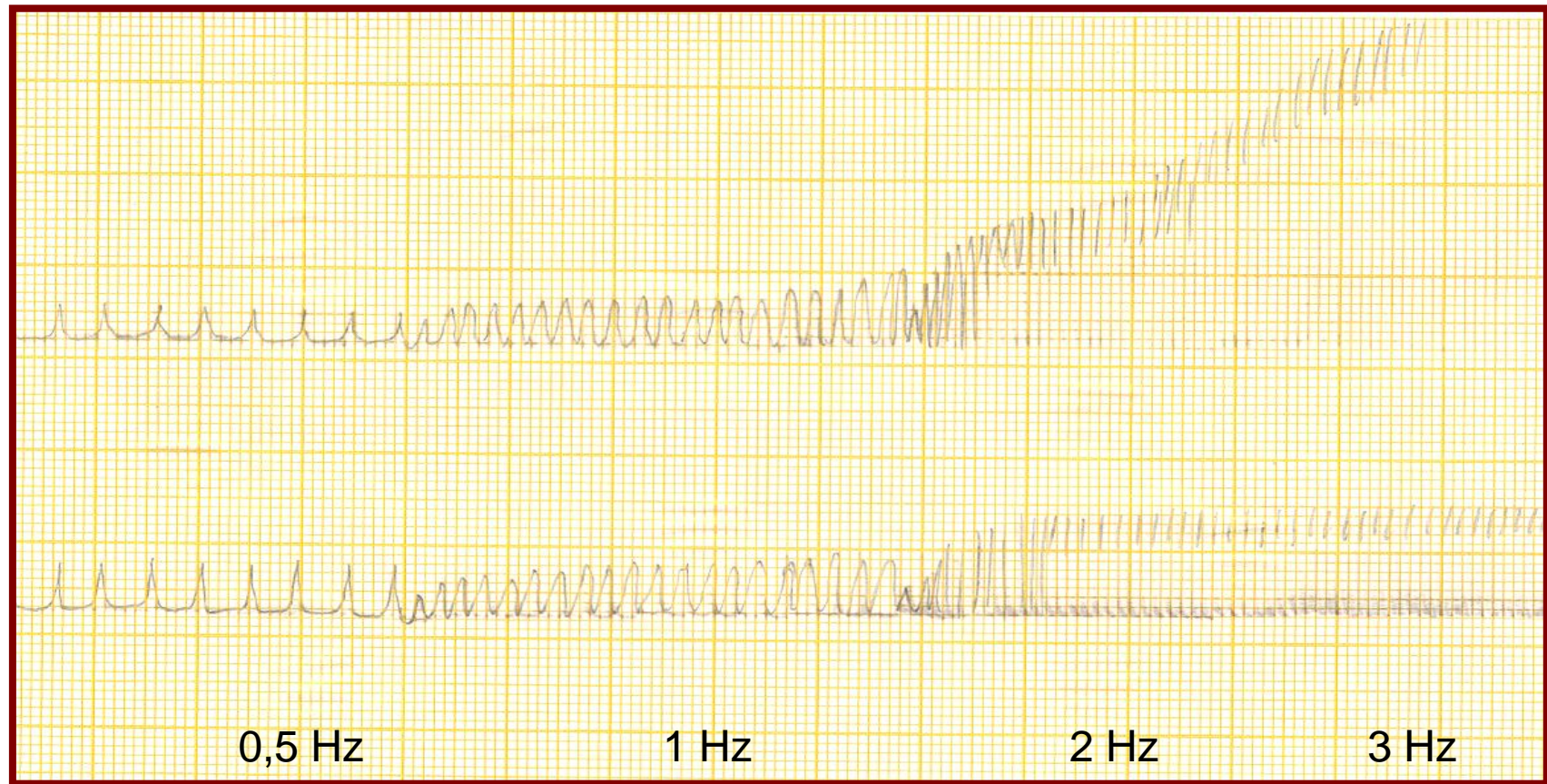
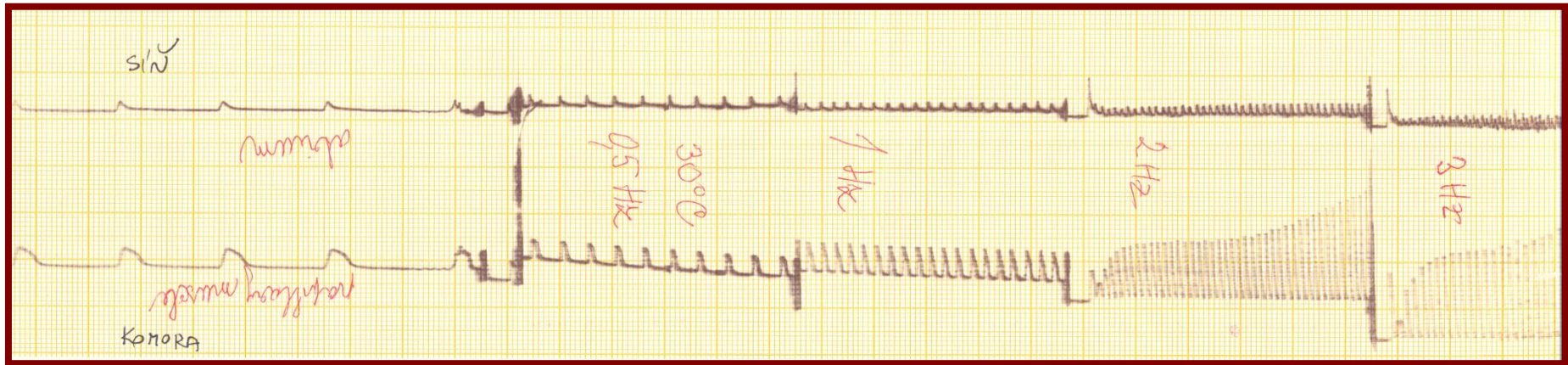
Henry Pickering Bowditch  
(1840 – 1911)



## HOMEOMETRIC AUTOREGULATION (FREQUENCY EFFECT)

During increasing HR (stimulation frequency) the force of developed contraction rises

Ratio between intra- and extracellular calcium concentrations increases



**CARDIAC RESERVE** = maximal CO / resting CO

**4 - 7**

**CORONARY RESERVE** = maximal CF / resting CF

**3,5**

**CHRONOTROPIC RESERVE** = maximal HR / resting HR

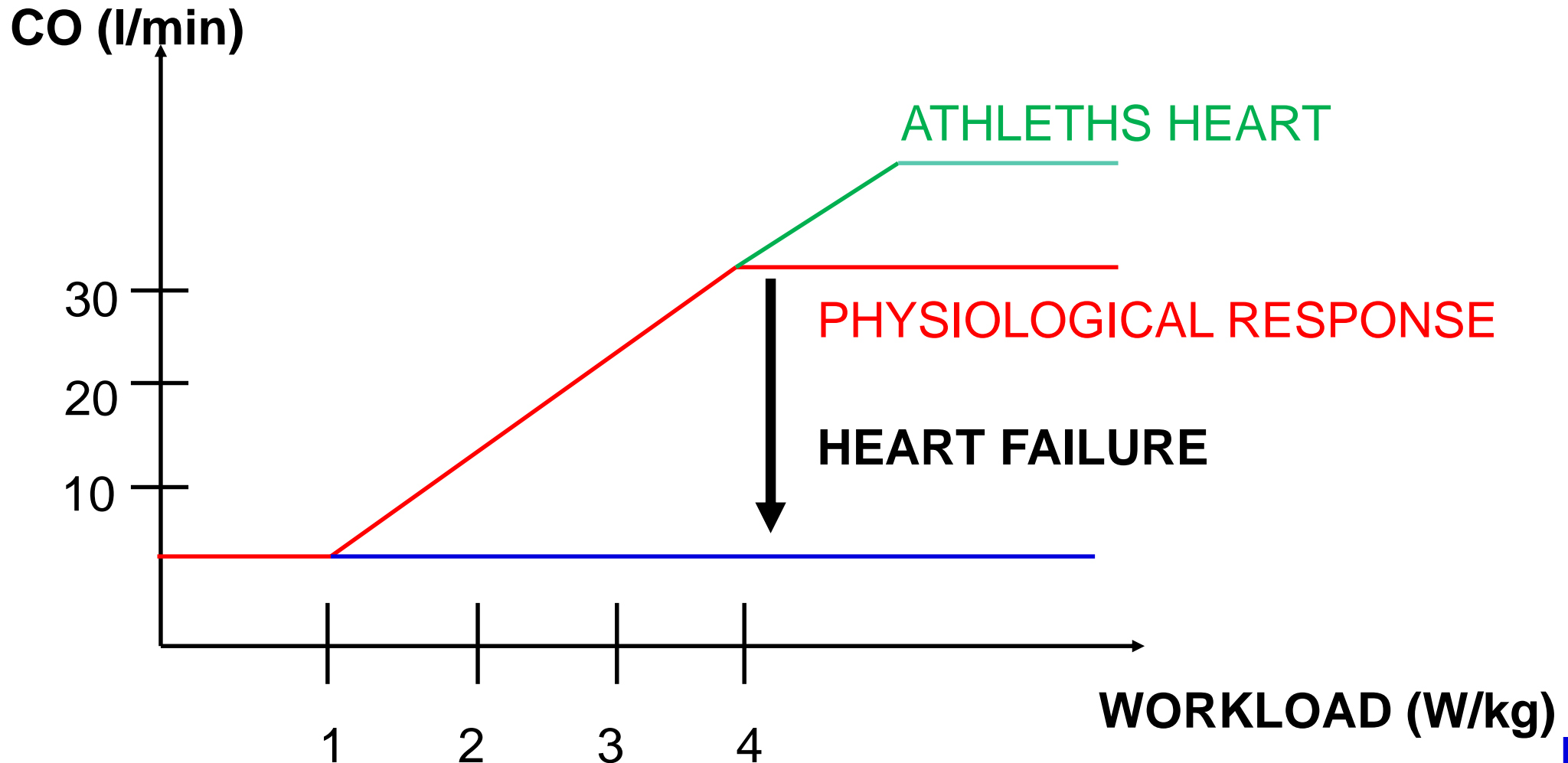
**3 - 5**

**VOLUME RESERVE** = maximal SV / resting SV

**1,5**

CO = cardiac output  
CF = coronary flow  
HR = heart rate  
SV = stroke volume

# CARDIAC RESERVE





## IMPORTANT TERMS

Length-tension relationship (curve)

Minimal length  $l_0$

Passive, active, total force

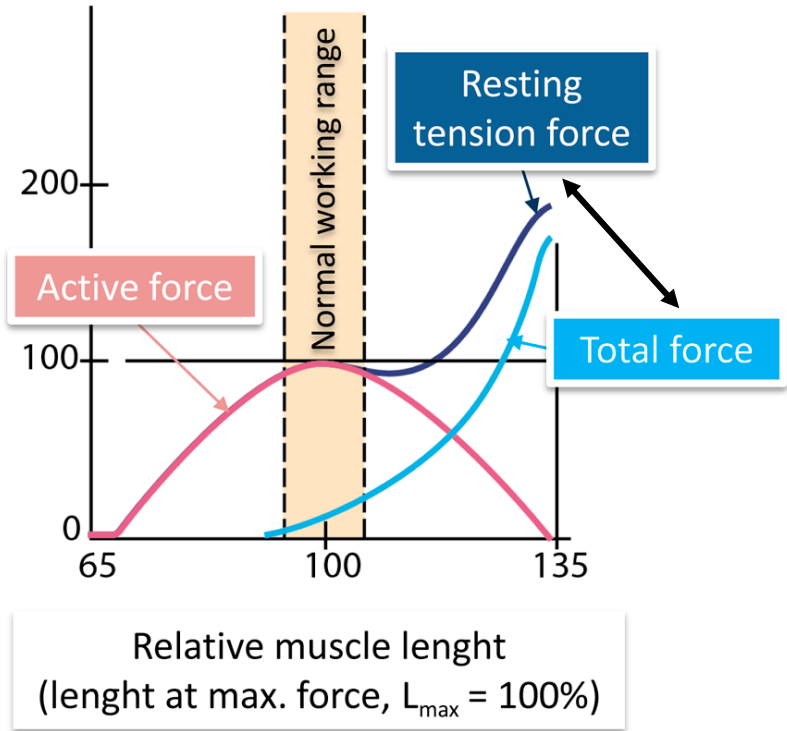
Optimal length

**Isometric, isotonic, auxotonic** contraction

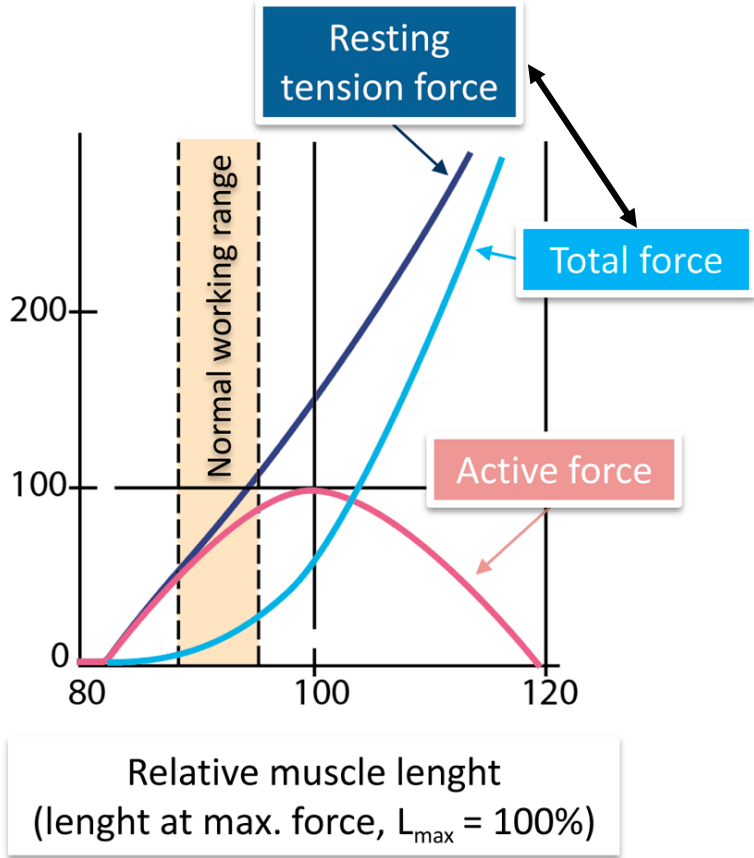
**Autoregulation** of contraction – **heterometric** (Starling)

**Preload, afterload**

# 1. Striated muscle

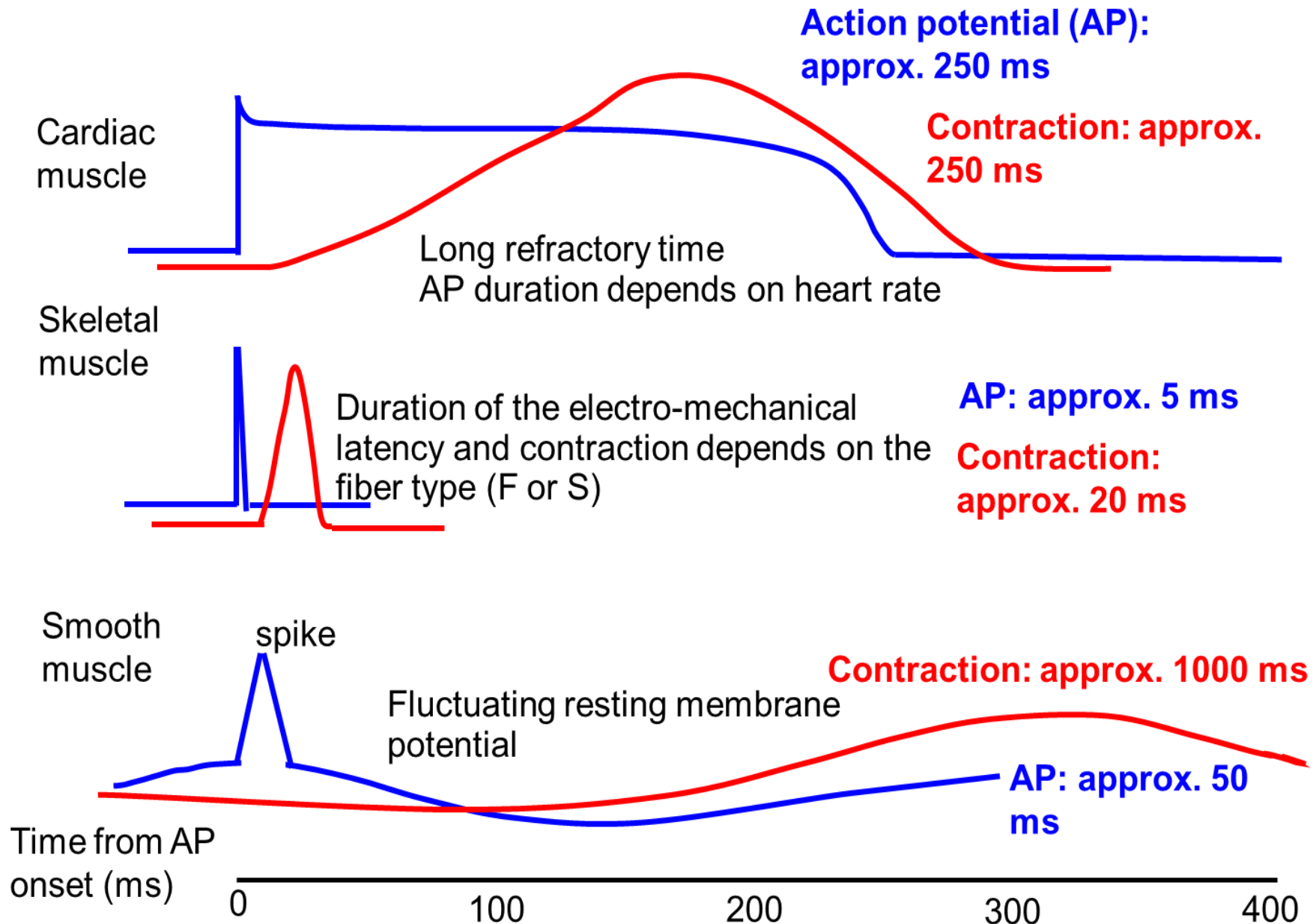


# 2. Cardiac muscle



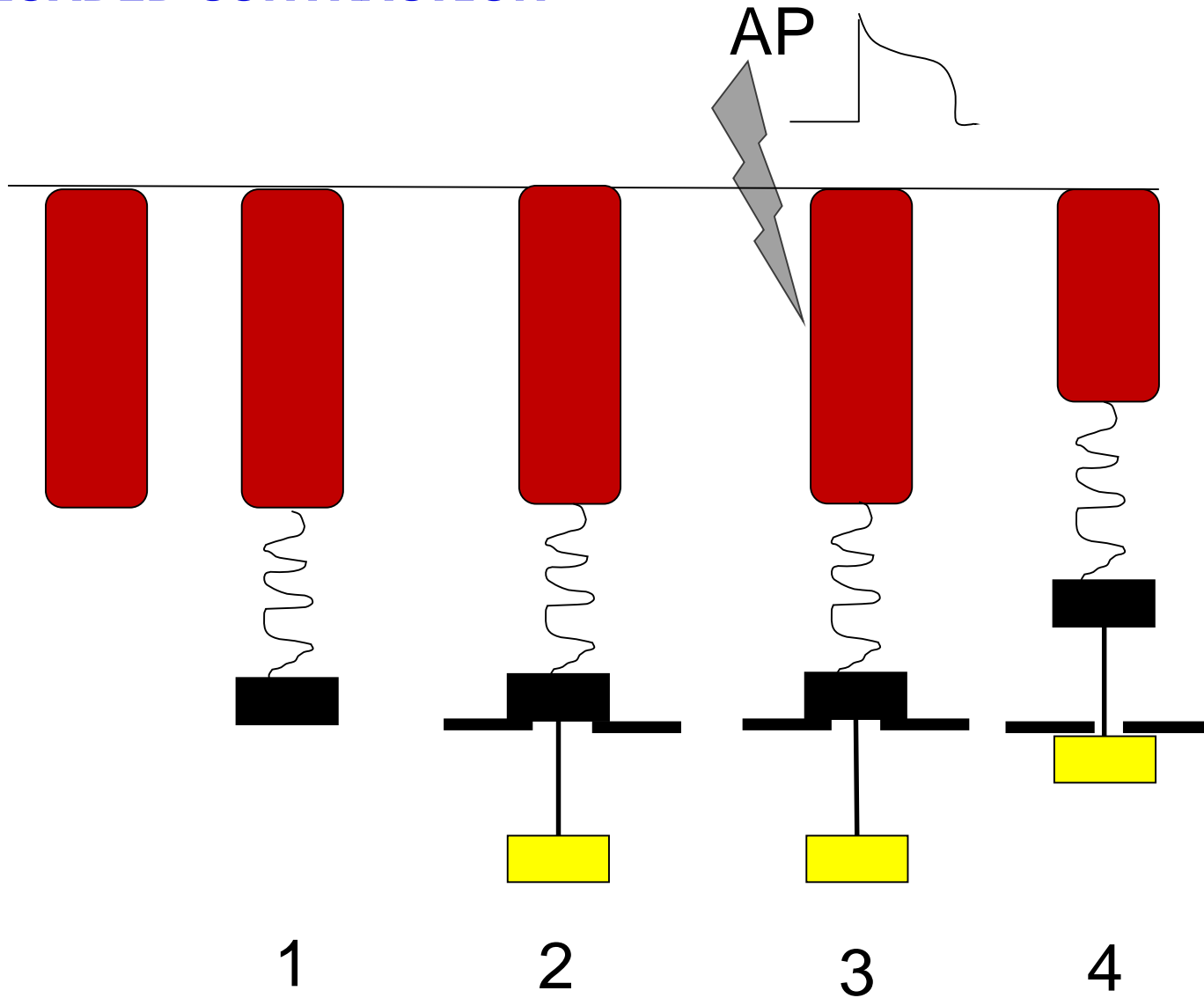
Passive tension, active tension, isometric contraction, isotonic contraction, auxotonic contraction

# Skeletal, cardiac and smooth muscle – action potential and contraction



# AFTERLOADED CONTRACTION

# PRELOAD, AFTERLOAD



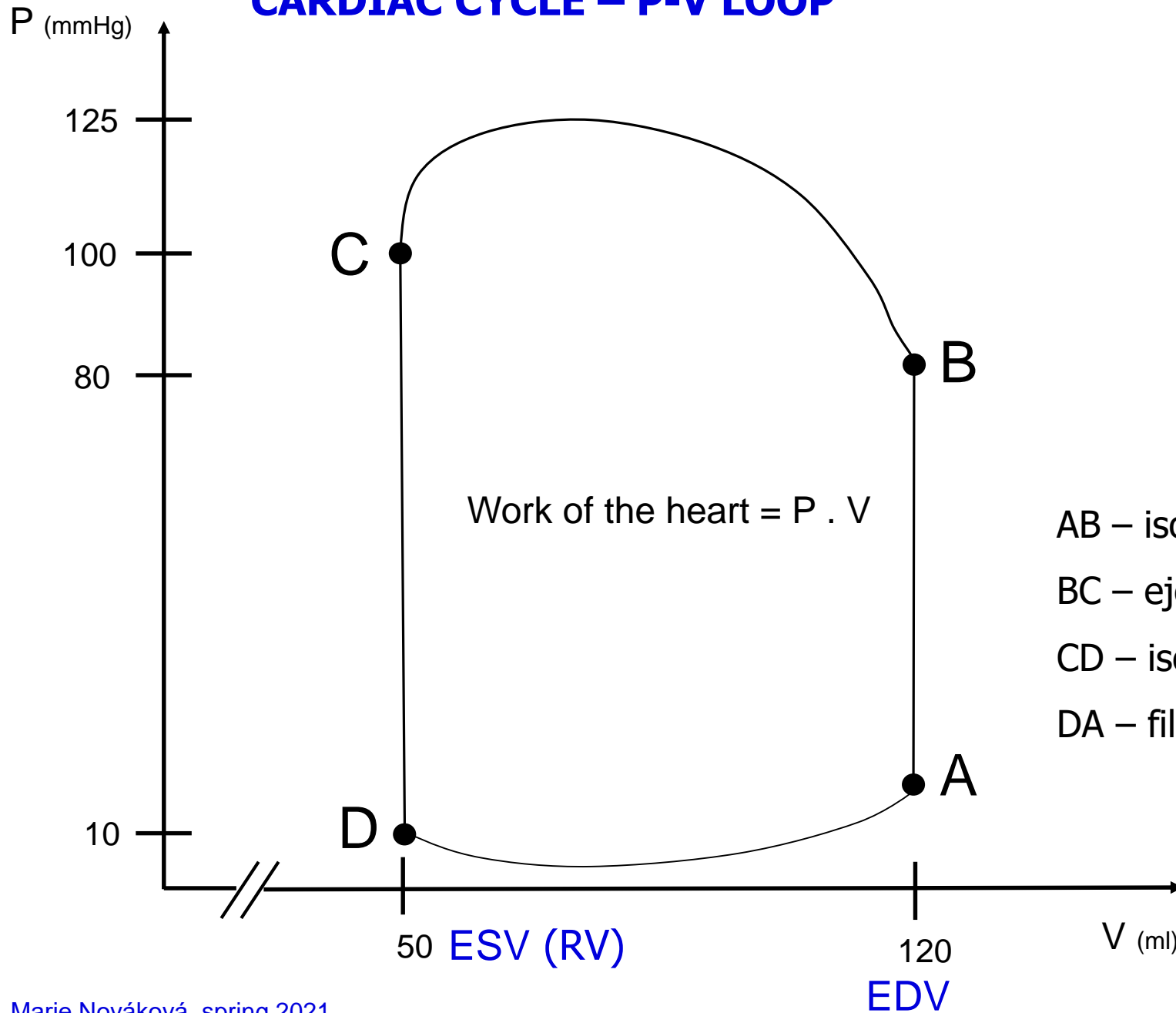
**PRELOAD**  
(~ enddiastolic filling)



**AFTERLOAD**  
(~ pressure which must be developed)



# CARDIAC CYCLE – P-V LOOP



LAPLACE law:

$$T = P \cdot R / h$$

$$\uparrow P = T \cdot \uparrow H / \downarrow r$$

AB – isovolumic contraction

BC – ejection

CD – isovolumic relaxation

DA – filling

$P = \underline{T} \cdot 2h \cdot \underline{r}^{-1}$       **Ventricular filling:** r and T rise, P first falls down, then rises up

(length/tension relationship)

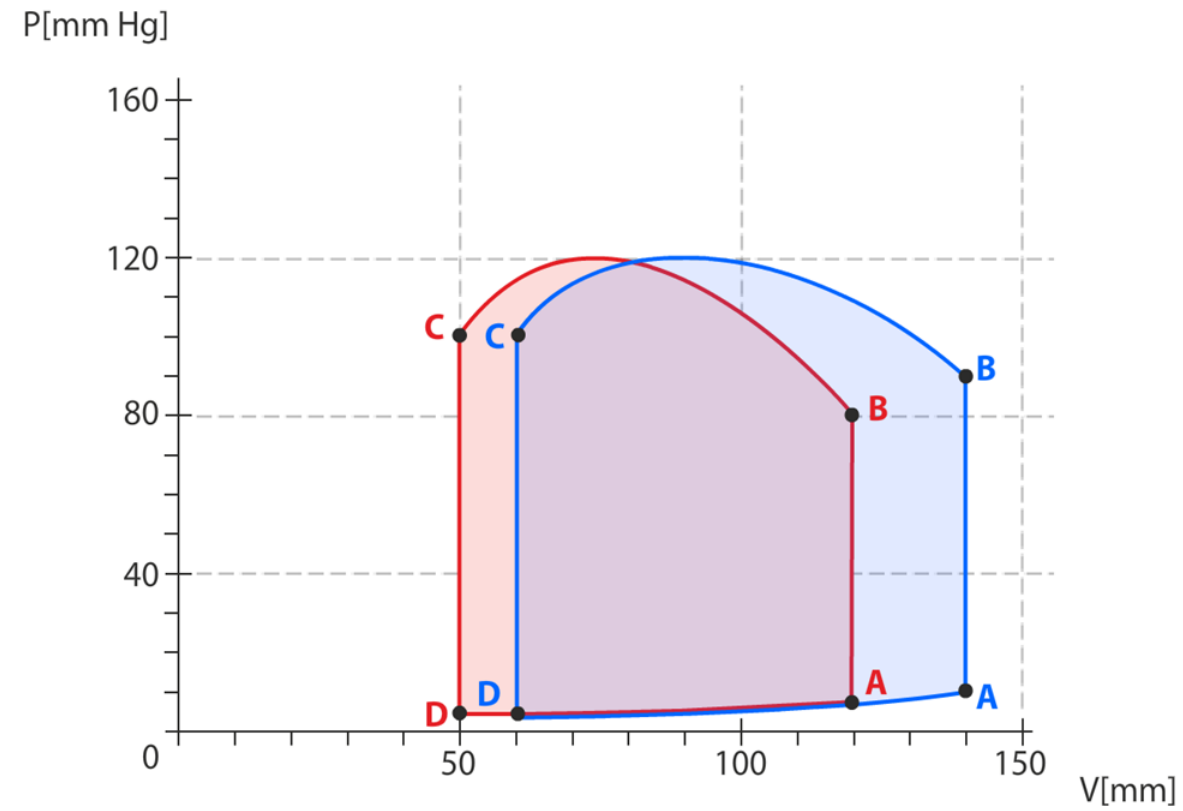
$\underline{P} = \underline{T} \cdot 2h \cdot r^{-1}$       **Isovolumic contraction:** T rises up, valves closed – increase in P

$\underline{P} = T \cdot \underline{2h} \cdot r^{-1}$       **Ejection:** r decreases, h rises, thus P increases (even at the same T)

$P = \underline{T} \cdot 2h \cdot r^{-1}$       **Isovolumic relaxation:** T decreases, valves closed – decrease in P

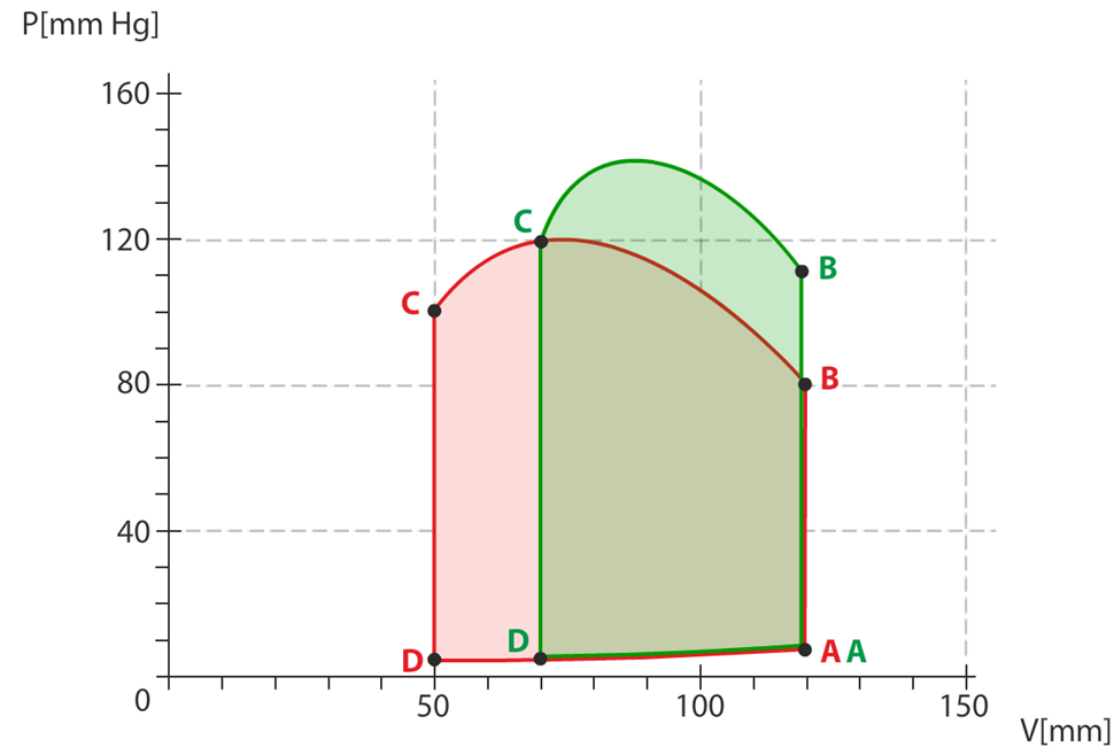
# INCREASED PRELOAD

## MODEL



# INCREASED AFTERLOAD

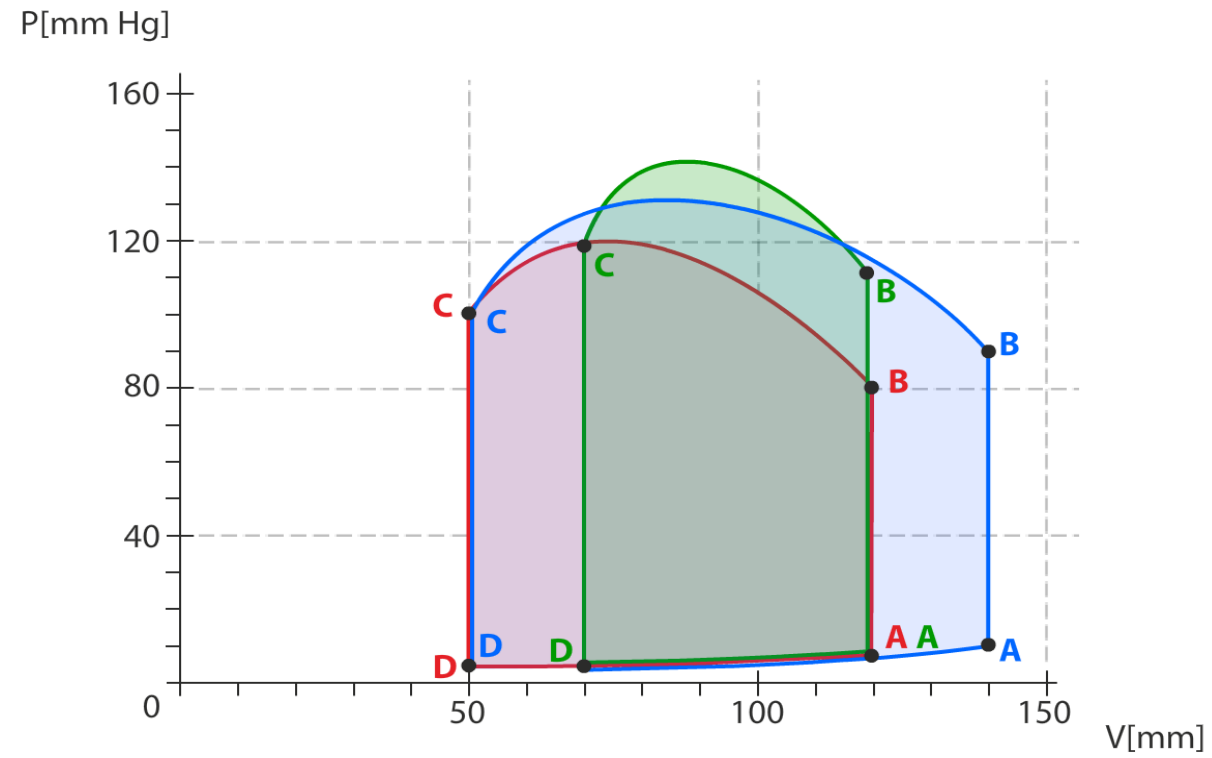
## MODEL





# INCREASED PRELOAD AND AFTERLOAD

## MODEL



# HEART SOUNDS

Caused by vibration of various anatomical structures and event. blood:

- **Closure and stretching of valves**
- Isovolumic contraction of heart muscle (papillary muscles, tendons)
- Turbulent blood flow

**I. – mitral (+ tricuspidal) valve closure**

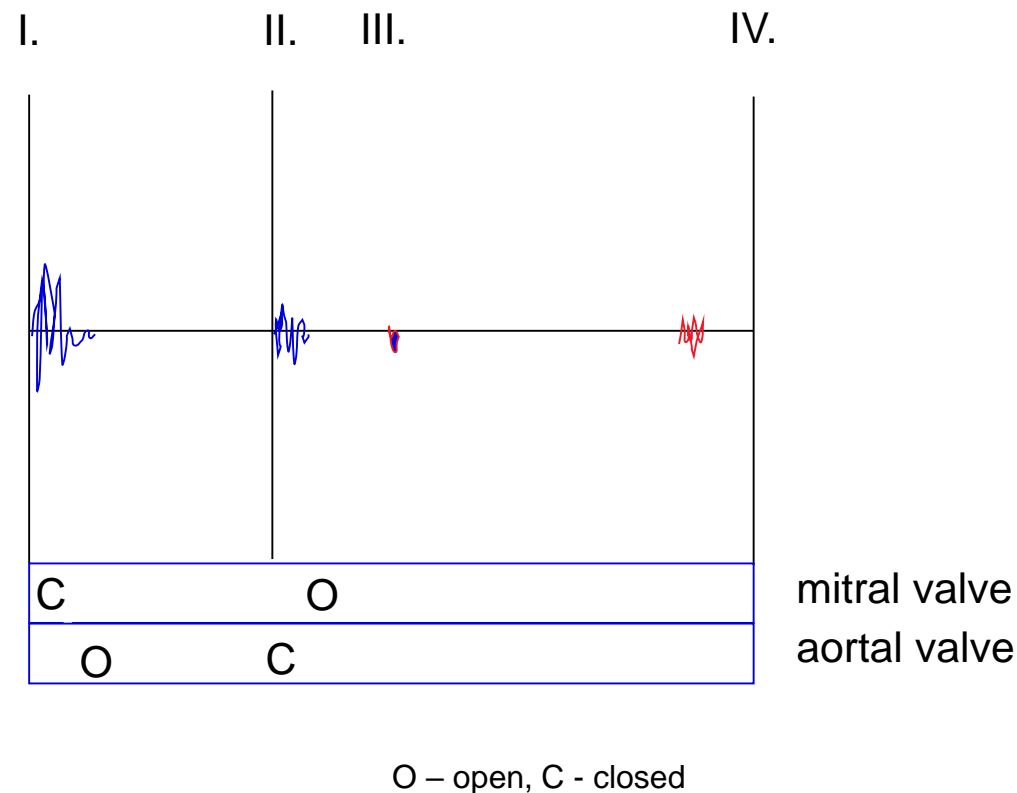
**II. - aortal (+ pulmonary) valve closure**

III. - fast filling of ventricles - **pathological**

IV. - contraction of atria – **mostly pathological**

→ **Vibration of ventricular wall**

**Splitting** of I. or II. sound:  
asynchronous closure of M - T valve (I.)  
or Ao - P valve (II.)  
(inspiration, hypertension....)



# MURMURS – pathological phenomena based on turbulent blood flow

## 1. SYSTOLIC

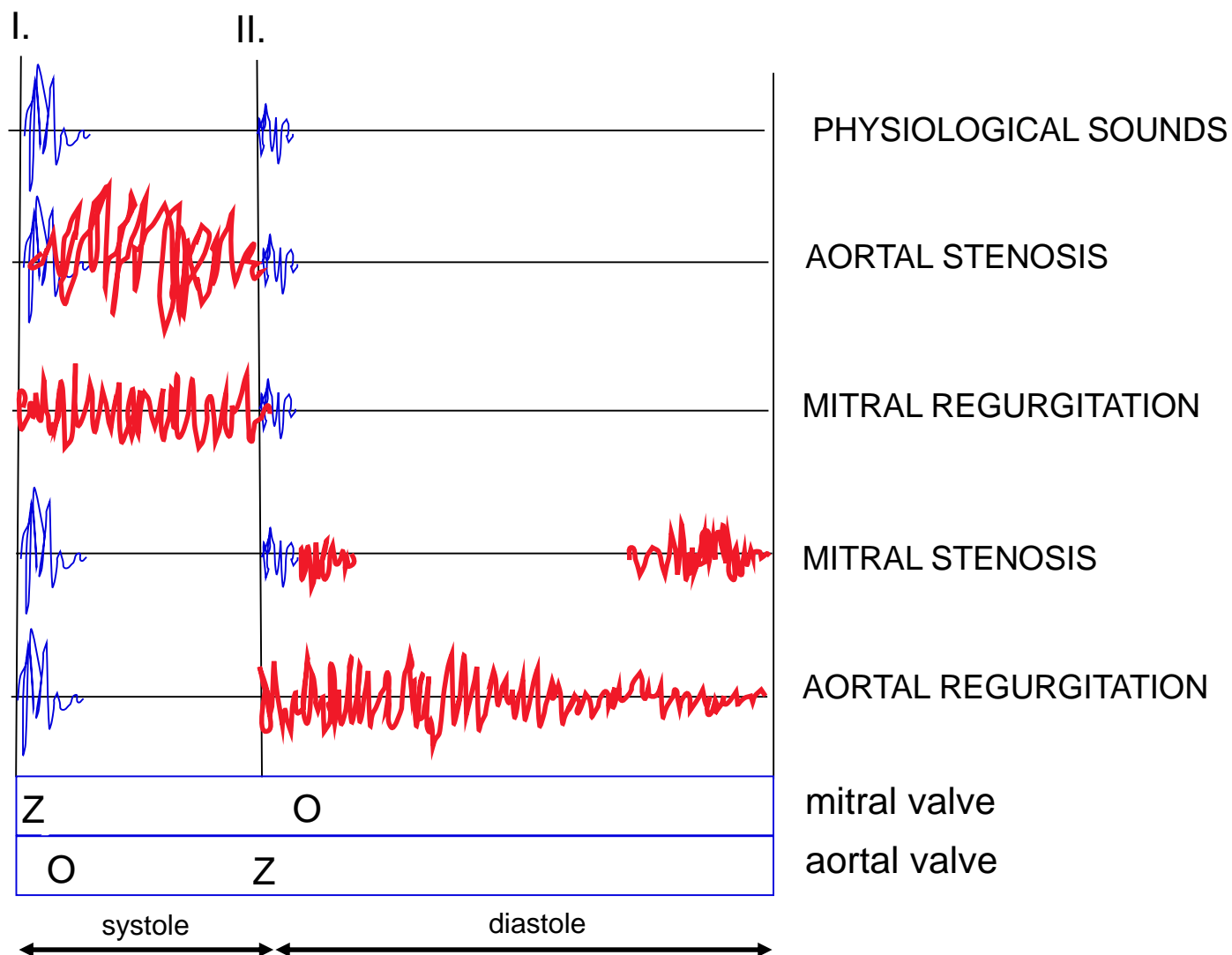
- Stenosis – aortal, pulmonary (1)
- Regurgitation – mitral, tricuspidal (2)

## 2. DIASTOLIC

- Stenosis – mitral, tricuspidal (3)
- Regurgitation – aortal, pulmonary (4)

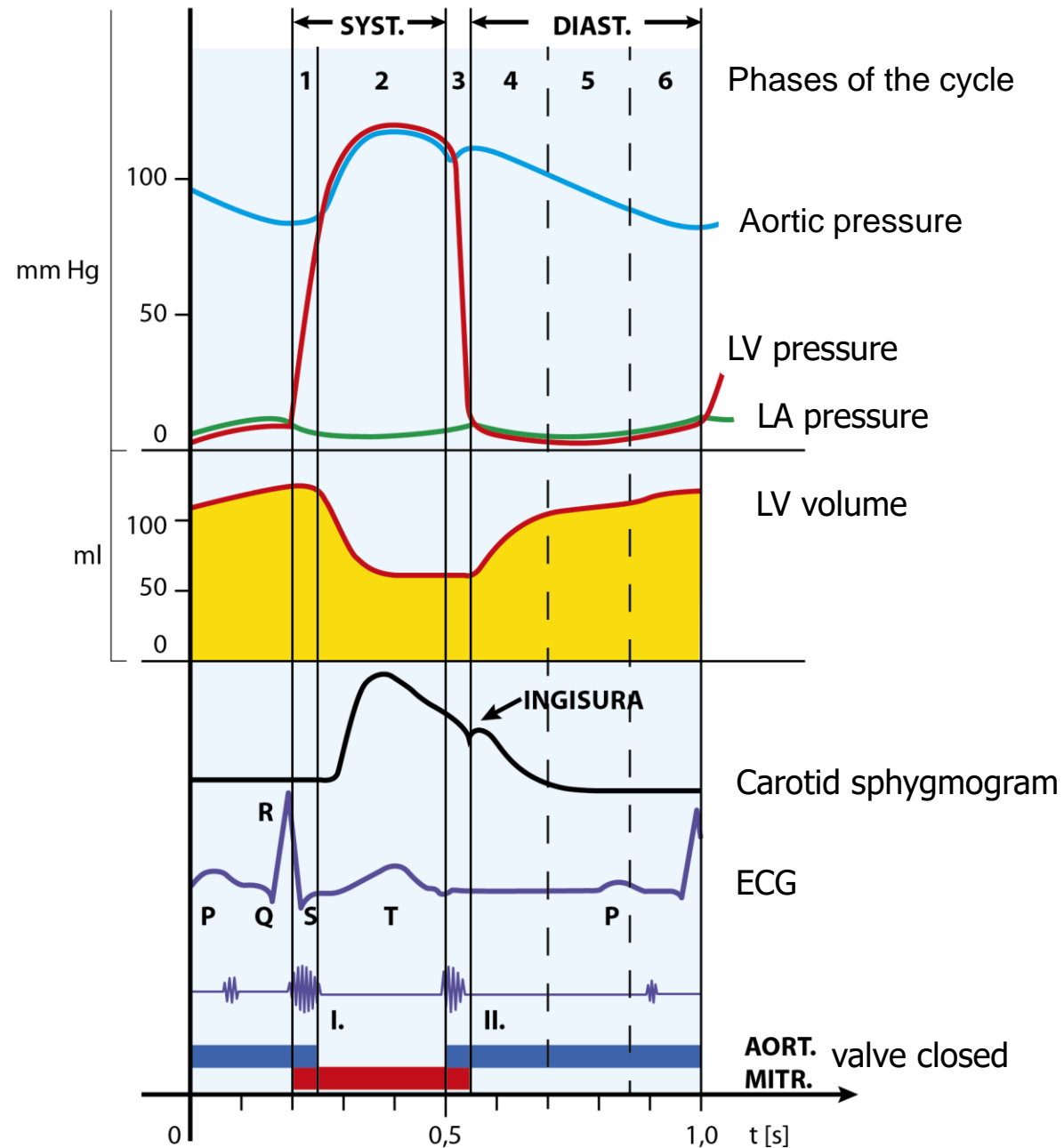
## 3. SUSTAINED:

- Defects of septum



O – open, C - closed

# POLYGRAPHY (polygram)



# HEART FAILURE = loss of cardiac reserve

The heart is not able pump sufficient amount of blood into periphery at normal venous return.

## SYMPTOMS

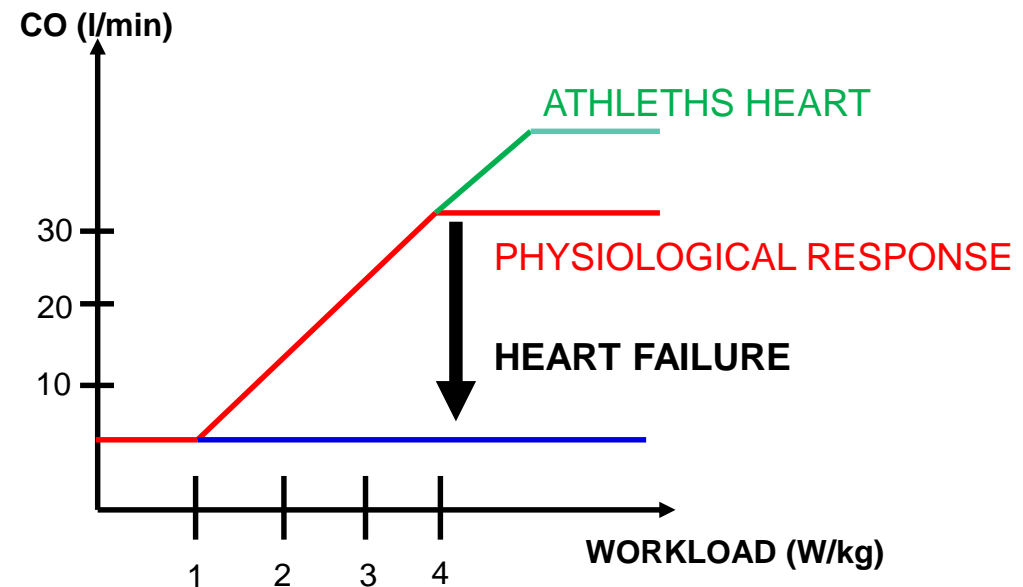
fatigue, oedemas, venostasis, dyspnoea, cyanosis

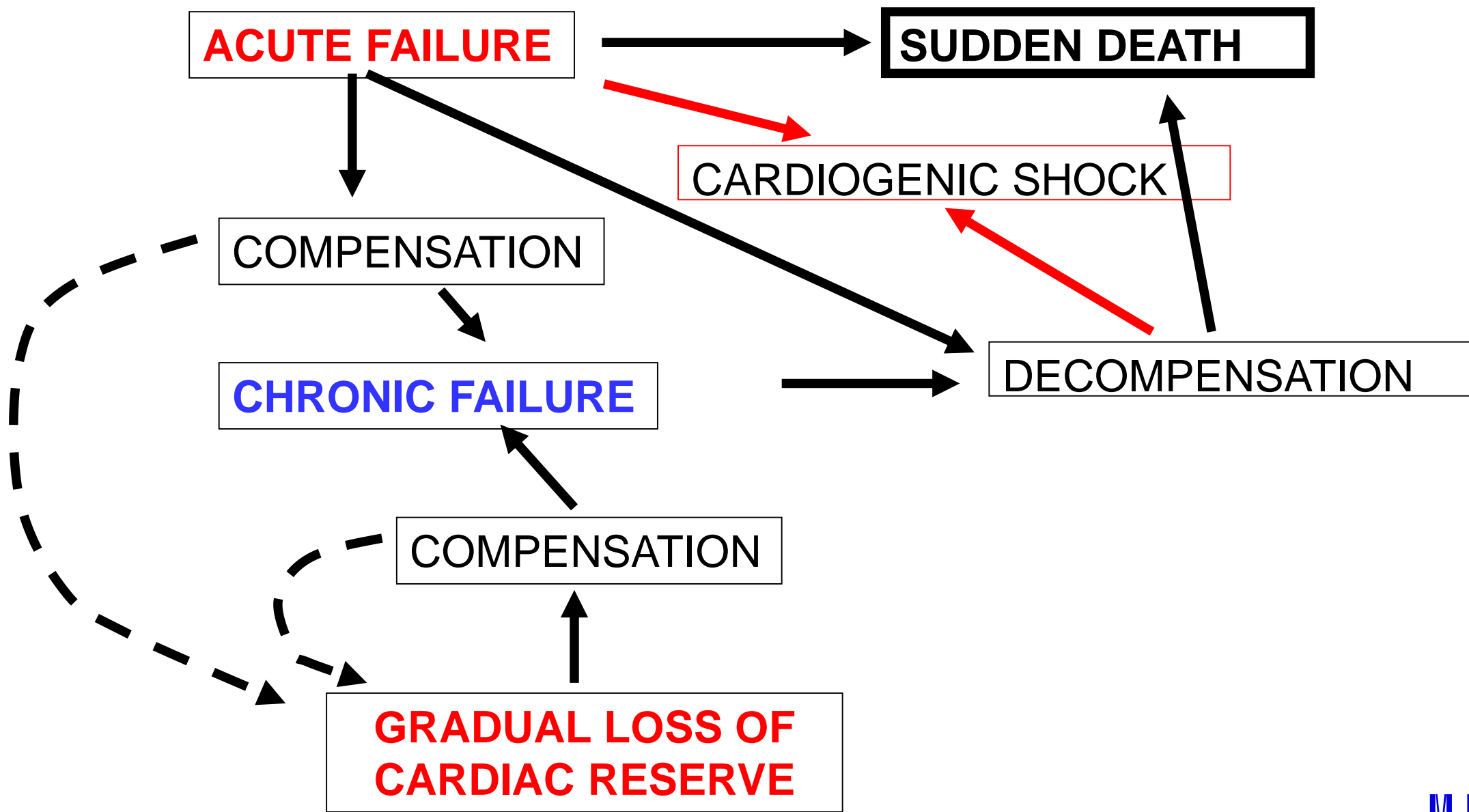
**ACUTE x CHRONIC.**

**COMPENSATED x DECOMPENSATED.**

## MOST FREQUENT CAUSES:

- Severe **arrhythmias**
- **Overload** – *volume* (aortal insufficiency, a-v shunts) or *pressure* (hypertension and aortal stenosis – left overload, pulmonary hypertension and stenosis of pulmonary valve – right overload)
- **Cardiomyopathy**





# HEART FAILURE - COMPENSATION

## BAROREFLEX

**Physiological role:** compensation of decrease in minimal volume of circulating fluids

**Signal:** BP decrease (orthostase, work vasodilatation)

**Sensor:** baroreceptors

**Response:** activation of SAS (increased HR, inotropy, BP)

**Pathological signal:** long-lasting decrease of BP due to heart insufficiency

**Results:** increased energy outcome – **vicious circle**

Ca<sup>2+</sup> - antagonists

β – sympatholytics

## ACTIVATION OF RAAS

**Physiological role:** compensation of loss of circulating fluids (bleeding)

**Signal:** decrease in renal perfusion

**Sensor:** juxtaglomerular system of kidney

**Response:** BP increase (angiotenzin II.), water retention (aldosteron)

**Pathological signal:** decrease in renal perfusion due to heart insufficiency

**Results:** increased preload and afterload, increased energy outcome – **vicious circle**

angiotenzin-converting  
enzyme inhibitors (AT II.  
receptors)

## **DILATATION (STARLING PRINCIPLE)**

**Physiological role:** compensation of momentary right-left differences

**Signal:** orthostase, deep breathing, beginning of exercise

**Pathological signal:** continual blood stasis in the heart

**Results:** increased energy outcome – **vicious circle**

## **HYPERTROPHY**

**Physiological role:** preservation of energetically demanding tension of ventricular wall

**Signal:**  $P = s \cdot 2 h / r$ , intermittent BP increase (athletes heart)

**Response:** concentric remodelling

**Pathological signal:** continual increase of preload or afterload

**Results:** worsening of oxygenation, fibrotisation – **vicious circle**

diuretics

cardiac glycosides (digitalis)