

CARDIAC MECHANICS
HEART AS A PUMP
CARDIAC CYCLE
HEART FAILURE

CARDIAC OUTPUT (CO)

LV = RV

REGULATION

CORONARY

 $CO = HR \times SV$

5l/min

SV = EDV - ESV

70ml

>60%

Ejection fraction

EF = EDV - ESV / EDV

Venous return Compliance Aortal pressure

CONTRACTILITY

Ability to contract
Depends on tissue perfusion
(substrates and oxygen supply for
ATP production; Ca2+ availability)

AUTOREGULATION of cardiac contraction

Heterometric: Starling law

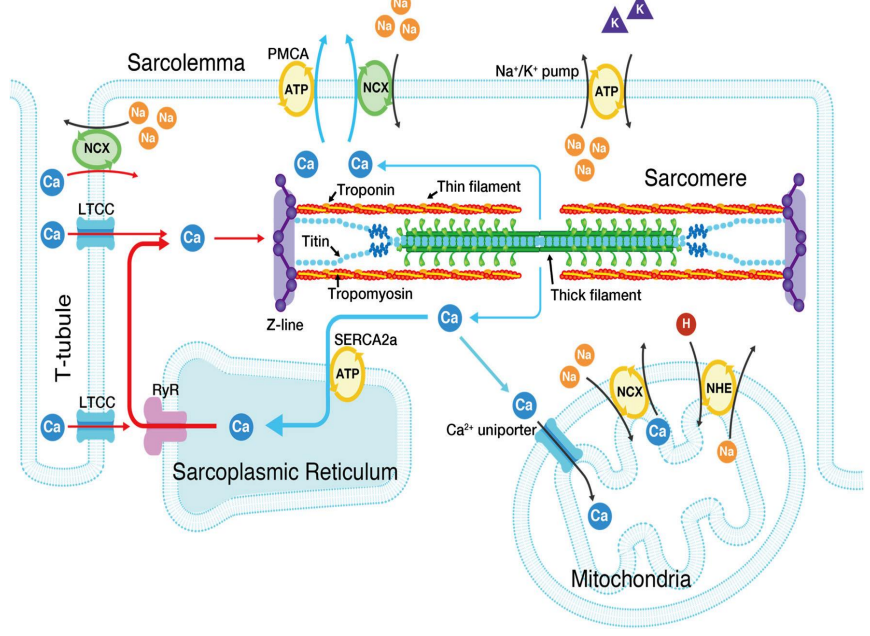
Homeometric: Frequency effect

EDV

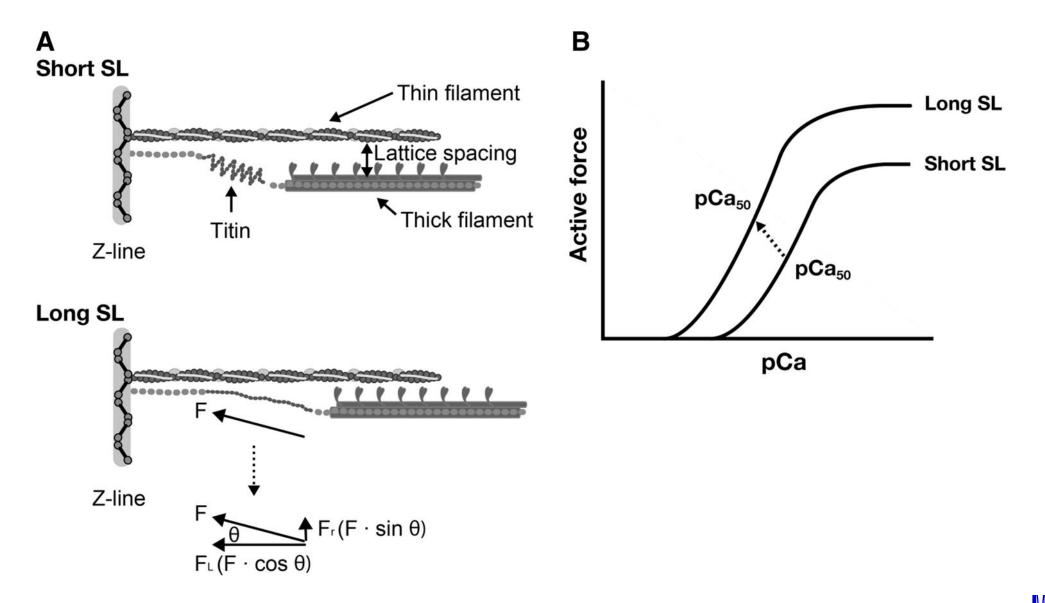


ANS

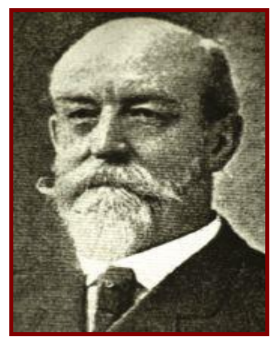
STARLING LAW



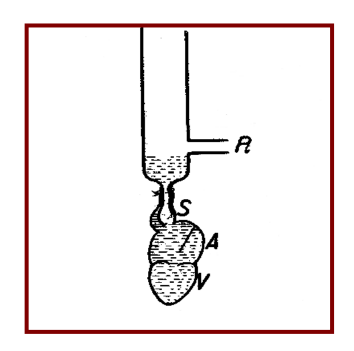


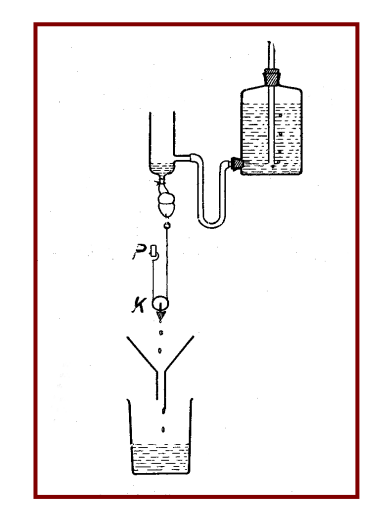






Henry Pickering Bowditch (1840 – 1911)





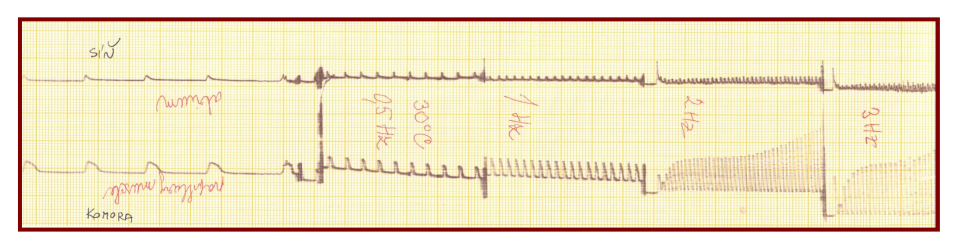
HOMEOMETRIC AUTOREGULATION

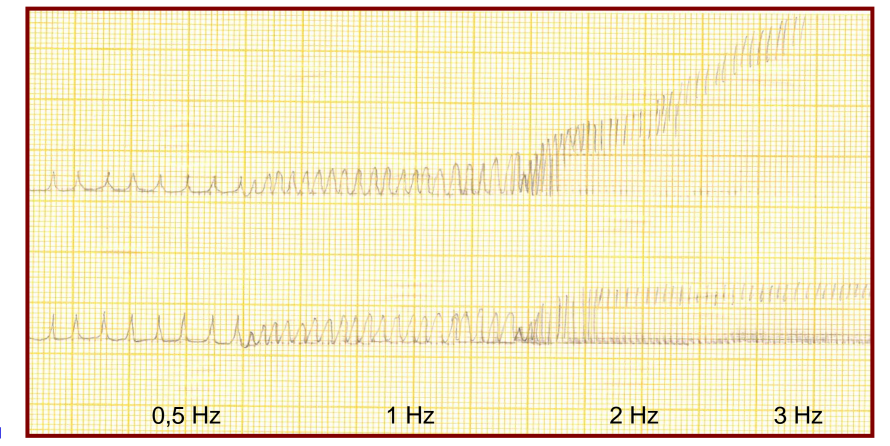
(FREQUENCY EFFECT)

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

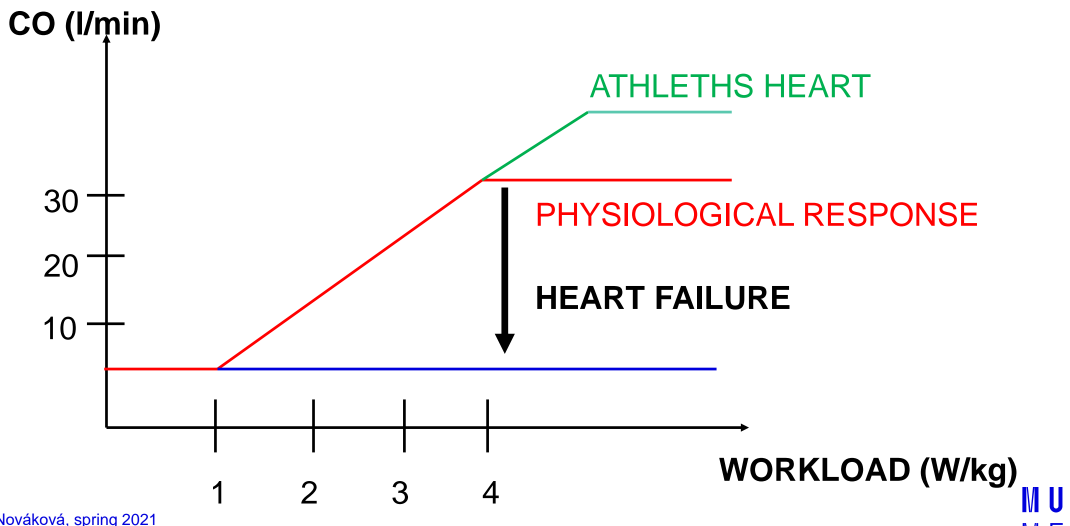
CF = coronary flow

HR = heart rate

SV = stroke volume



CARDIAC RESERVE



IMPORTANT TERMS

Length-tension relationship (curve)

Minimal length l₀

Passive, active, total force

Optimal length

Isometric, isotonic, auxotonic contraction

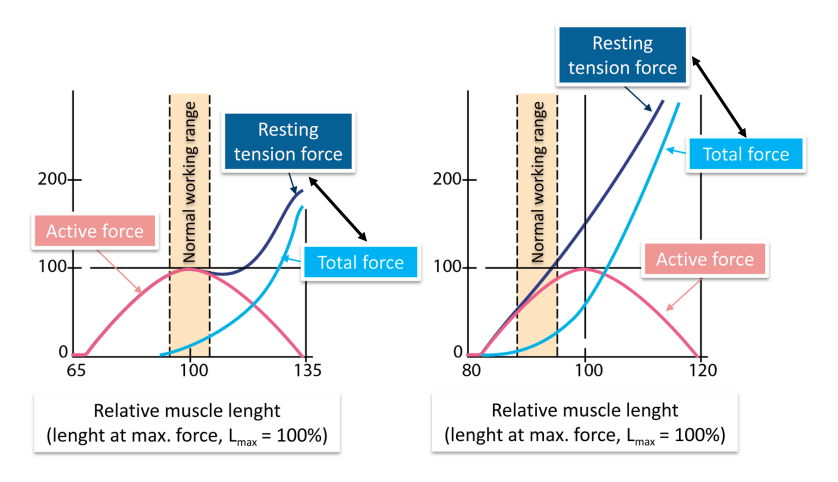
Autoregulation of contraction – **heterometric** (Starling)

Preload, afterload



1. Striated muscle

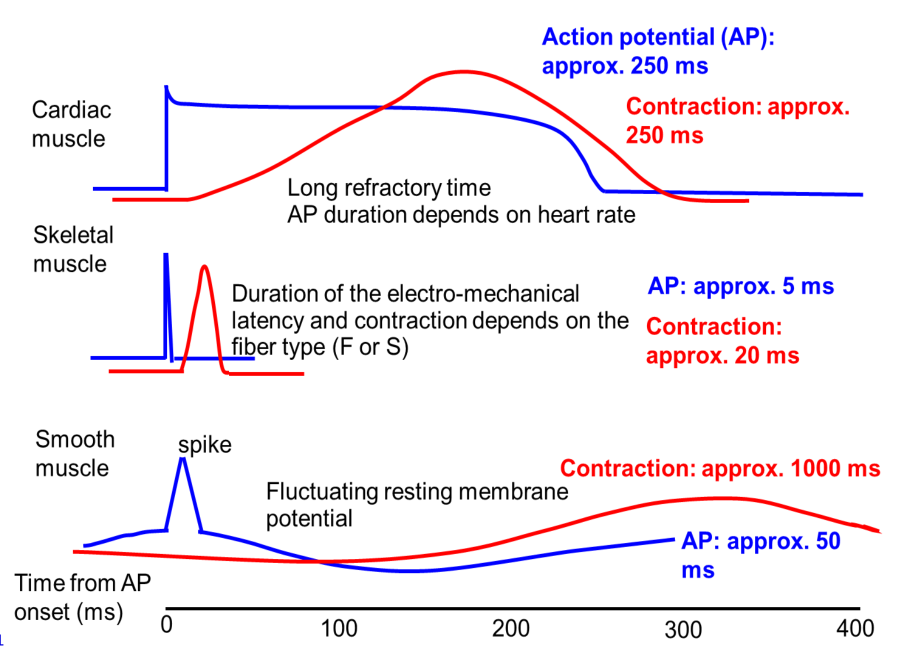
2. Cardiac muscle

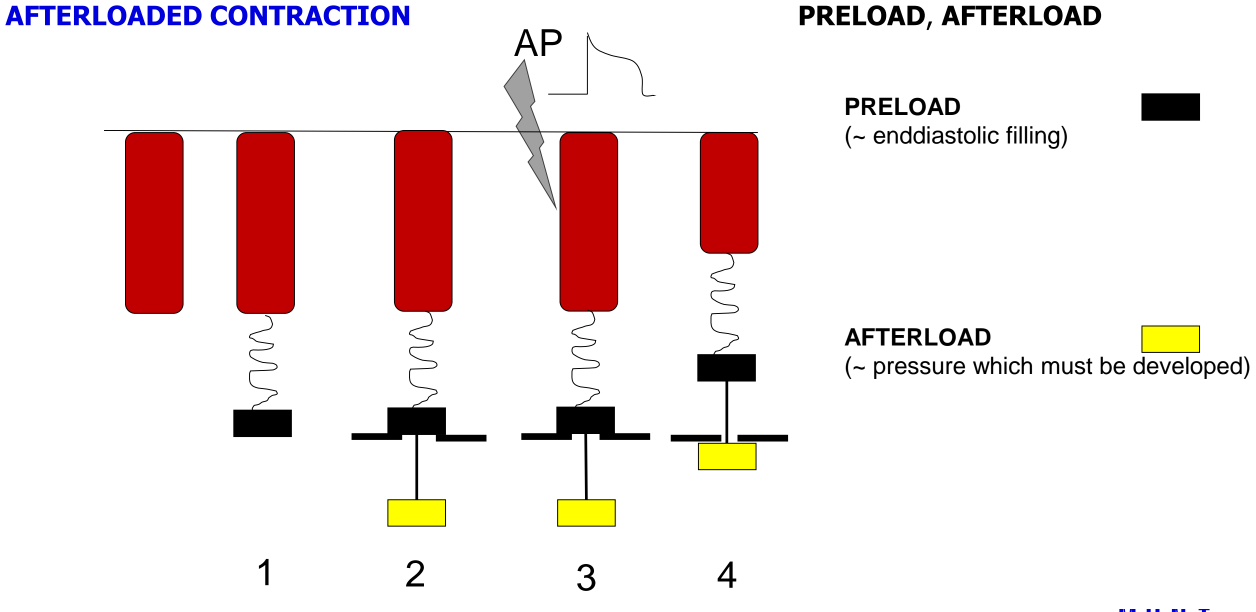


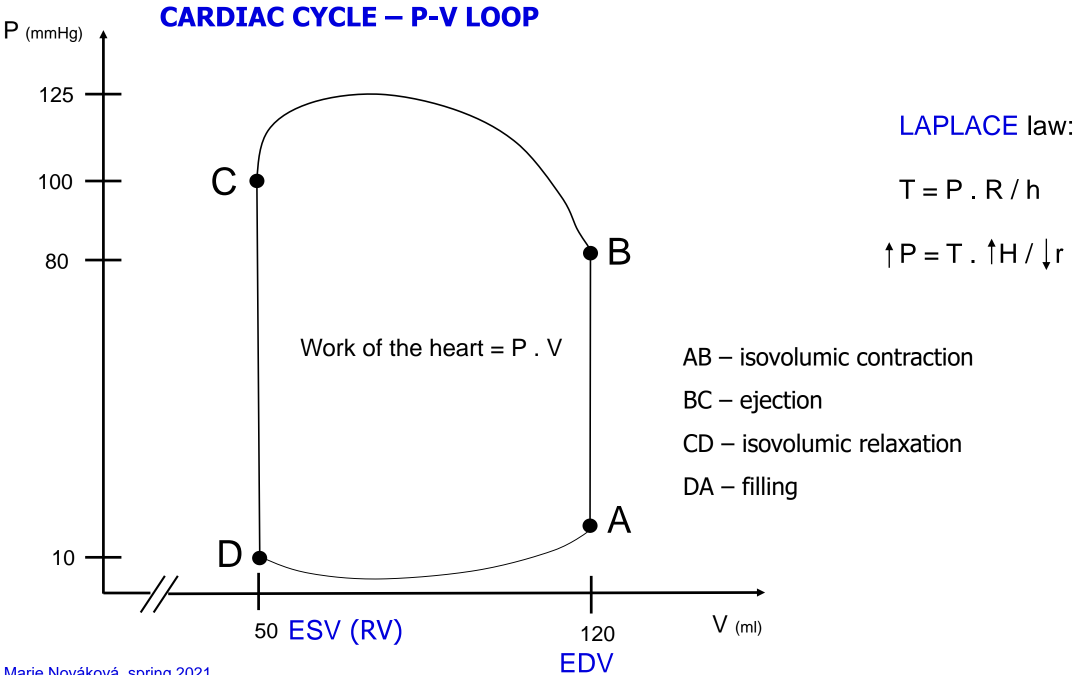
Passive tension, active tension, isometric contraction, isotonic contraction, auxotonic contraction



Skeletal, cardiac and smooth muscle – action potential and contraction







 $P = \underline{T}$. 2h $\underline{r-1}$ Ventricular filling: r and T rise, P first falls down, then rises up (length/tension relationship)

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

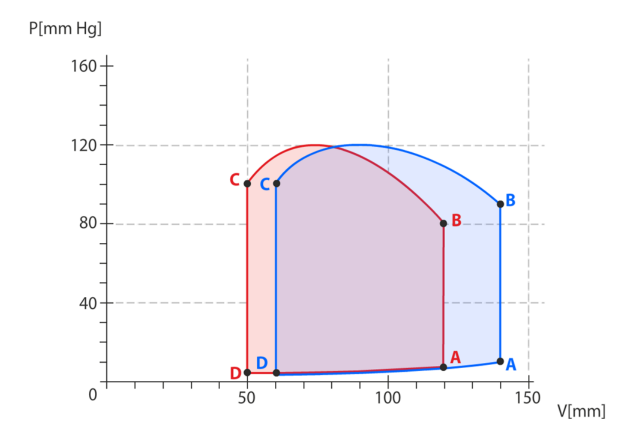
P = T.2h.r⁻¹ Ejection: r decreases, h rises, thus P increases (even at the same T)

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



INCREASED PRELOAD

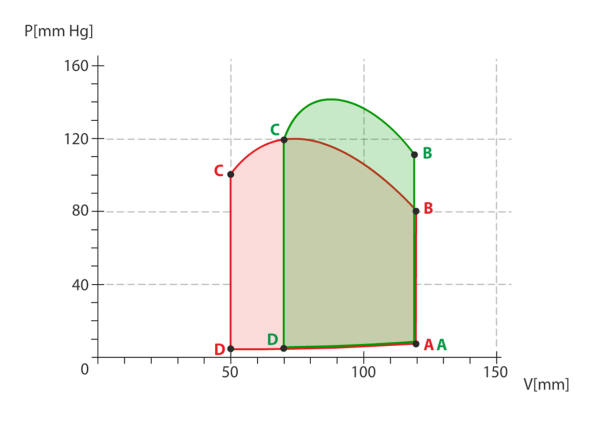
MODEL





INCREASED AFTERLOAD

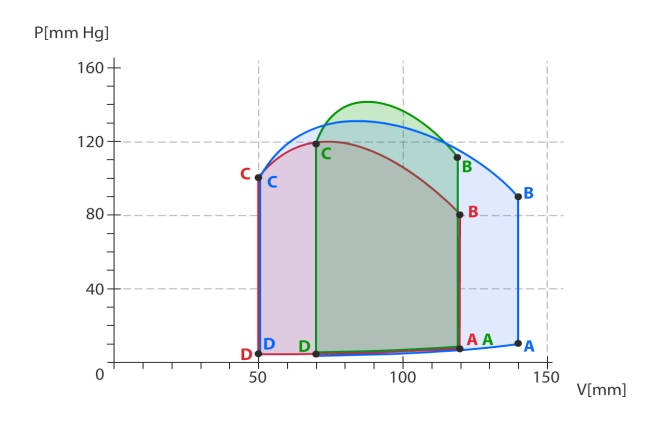
MODEL





INCREASED PRELOAD AND AFTERLOAD

MODEL





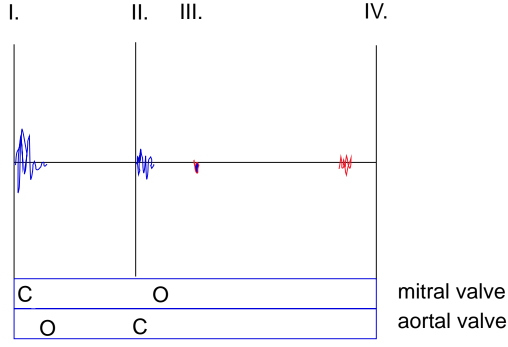
HEART SOUNDS

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....)



O - open, C - closed



MURMURS - pathological phenomena based on turbulent blood flow

1. SYSTOLIC

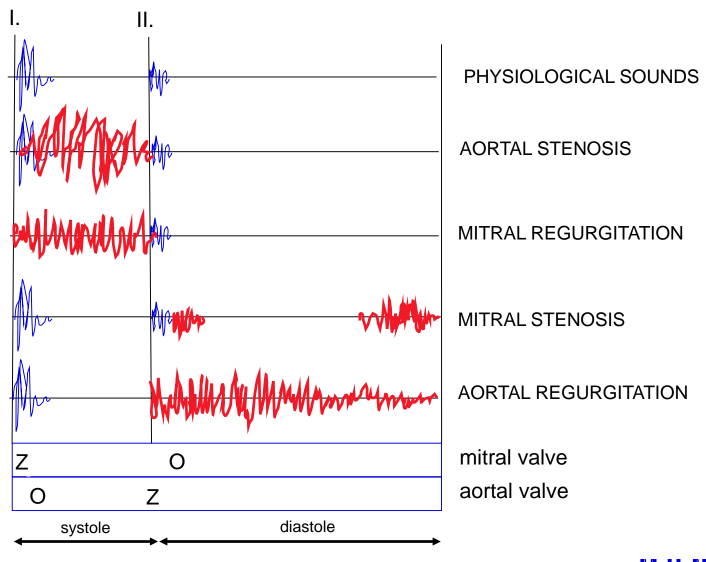
- Stenosis aortal, pulmonary (1)
- Regurgitation mitral, tricuspidal (2)

2. DIASTOLIC

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

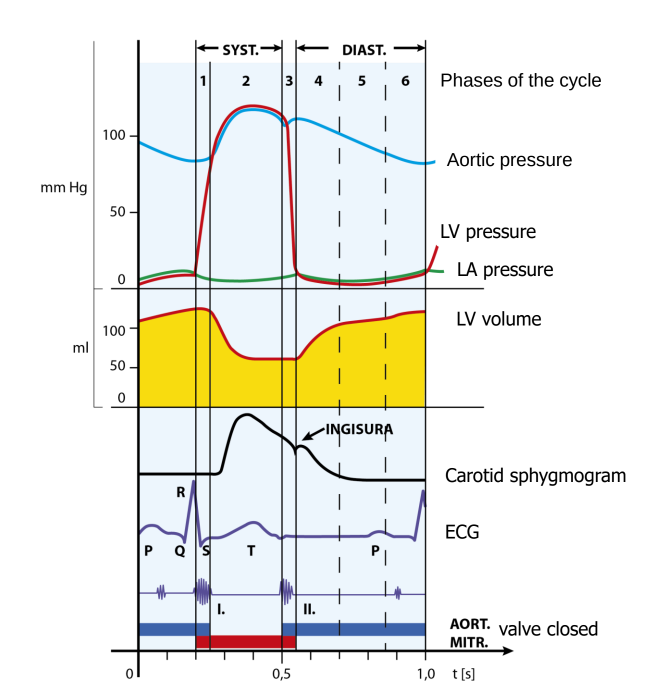
3. SUSTAINED:

Defects of septum





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

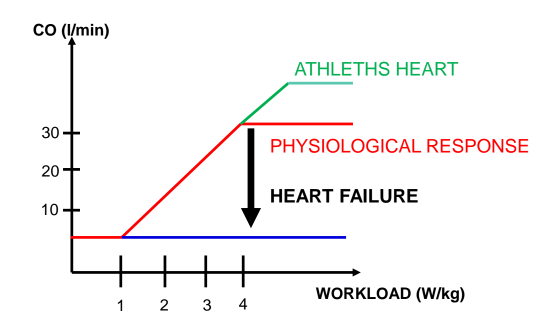
fatigue, oedemas, venostasis, dyspnoea, cyanosis

ACUTE x CHRONIC.

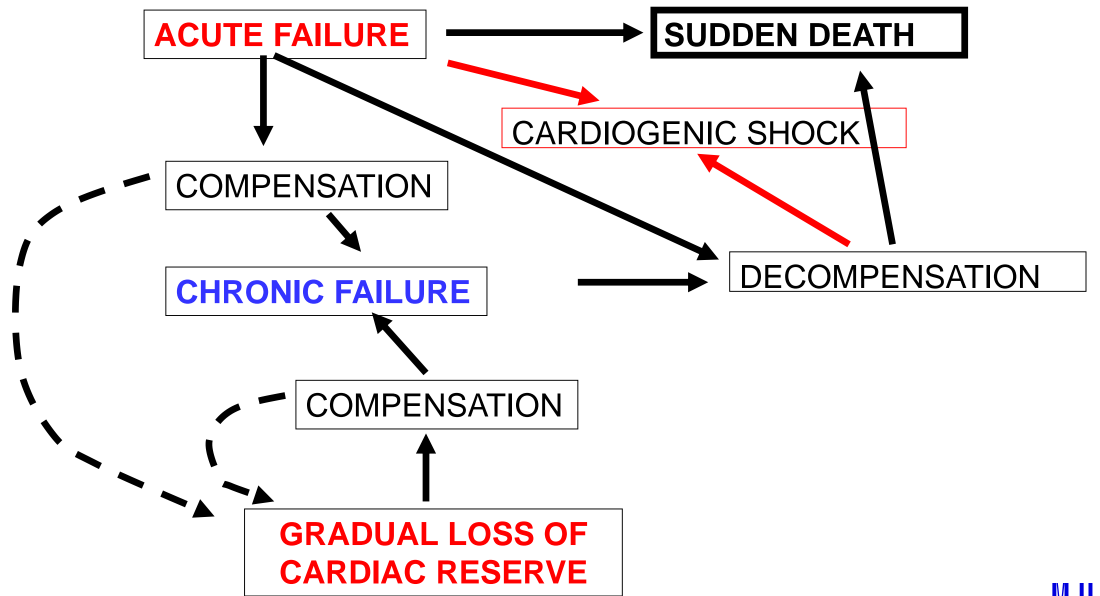
COMPENSATED x DECOMPENSATED.

MOST FREQUENT CAUSES:

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

Ca²⁺ - antagonists

 β – sympatolytics

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)

