# Pathophysiology of circulatory shock

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# Shock - definition

- Severe tissue hypoperfusion resulting in low supply of oxygen to the organs
- Systemic hypotension (of various causes) is present
- $P = Q \times R$
- Q ~ CO = SV × f
- CO depends on
  - a) cardiac function
  - b) venous return ( $\rightarrow$ preload)
- R systemic resistance (mostly arterioles) afterload

### Cardiac function and venous function





# Vascular resistance

× R [kg.s<sup>-1</sup>.m<sup>-4</sup>]: can be obtained from Hagen-Poiseuill law: R = 8 ×  $\eta$  × d /  $\pi$  × r<sup>4</sup>,where:

> η = viscosity d = length of the segment

r = radius



# Preload and afterload in heart

- Law of Laplace for wall tension in a hollow sphere:  $\sigma = \frac{P \times r}{2h}$ , where:
  - P....pressure inside the sphere
  - r....inner radius
  - h....wall thickness



- Preload wall tension (N.m<sup>-2</sup> = Pa force per area) before the systole
  - The main factor is venous return  $\rightarrow$  filling of cardiac ventricles
- Afterload increase in wall tension during the systole
  - The main factor is a peripheral resistance, or pulmonary vascular resistence in a case of the right ventricle
- Preload is higher in the right ventrikle, afterload is higher in the left one



# Phases of shock

- Compensation of initiating cause
- Decompensation
- Refractory shock

### Compensatory mechanisms and their limits

- Activation of sympathetic nervous system (tens of seconds)
- Activation of RAAS (cca I hour)
- Vasoconstriction (if possible) but it leads into lower blood supply
- Vasodilatation in some tissues (esp. myocardium)
- Positively inotropic effect of SNS (if possible) but at cost of higher metabolic requirements of the heart
- Increased heart rate but CO decreases in high HR (>150 bpm)
- Keeping circulating volume by lower diuresis – but at cost of acute renal failure
- Shift to anaerobic metabolism but at cost of ↓ ATP a ↑ lactate (acidosis)
- Increased respiratory rate (but shallow breathing results in ↑ relative deadspace)
- Shift of saturation curve of hemoglobin to right (<sup>1</sup>2,3-DPG)
- Hyperglycemia but there is decreased utilization of Glc in the periphery





# Decompensated shock

- ↓ BP
- $\downarrow$  diuresis
- Brain hypoperfusion involvment of mental functions
- Acrocyanosis
- Tachypnea
- "Golden hour"

# Shock at cellular level

- Mitochondrial dysfunction (result of hypoxia) lower production of ATP
- ↑ ROS production by dysfunctional mitochondria
- Failure of ion pumps (e.g. Na/K ATP-ase →↑intracelular Ca<sup>2+</sup>)
- Activation of Ca<sup>2+</sup> -dependent proteases
- Lysosomal abnormalities release of lysosomal proteases
- ↓ intracelular pH, ↑ lactate
  - <sup>o</sup> promote hyperpolarization of muscle cells by opening K<sup>+</sup> channels  $\rightarrow \downarrow Ca^{2+}$  entry  $\rightarrow \downarrow$  smooth muscle cell and cardiomyocyte contraction



# **Refractory shock**

- Vicious circles
  - I) Vasodilatation  $\leftrightarrow$  hypoperfusion
  - Endothelial cells contain two isoforms of nitric oxid synthase constitutive (eNOS) and inducible (iNOS)
  - In lasting hypoxia of endothelial cells there is increased iNOS activity (primarily physiological mechanism)
  - ↑NO increases vasodilation and hypoperfusion
  - Lactate acidosis  $\rightarrow$  hypotension (lactate prognostic factor)
  - 2) Myocardial hypoxia  $\leftrightarrow$  lower contractility
  - $^\circ\,$  Lower myocardial perfusion leads into  $\downarrow CO$ , which further reduces coronary flow
  - Myocardium does not benefit from the shift of Hb saturation curve efficiency of  $O_2$  extraction is already at its maximum
  - 3) Brain hypoperfusion  $\leftrightarrow \downarrow$  SNS activity
  - Lower perfusion of vasomotor centre leads first into SNS hyperactivity, which is then followed by its supression
  - That leads into  $\downarrow$  brain perfusion

### Other vicious circles in refractory shock

### Vicious cycle of shock

\* SIRS (systemic inflammation) \* DIC (systemic activation of coagulation)



Source: Brunicardi FC, Andersen DK, Billiar TR, Dunn DL, Hunter JG, Matthews JB, Pollock RE: Schwartz's Principles of Surgery, 9th Edition: http://www.accessmedicine.com Copyright © The McGrav-Hill Companies, Inc. All rights reserved.

### Systemic Inflammatory Response Syndrome (SIRS)

- Systemic activation of immune mechanisms
- Causes:
  - infections (sepsis)
  - Shock caused by non-infectious causes (diffuse tissue damage in hypoxia)
  - Non-compatible blood transfusions
  - Radiation syndrome (esp. GIT form)

# Disseminated intravascular coagulopathy (DIC)

- Systemic exposure to thrombin
- Two phases:
  - Formation of microtrombi (with local ischemia)
  - 2) Bleeding as a result of consummation of coagulation factors
- Consequence of the vessel wall damage
- Moreover, slower blood flow contributes to the extent of coagulation reactions
- DIC is especially frequent in septic shock



# Signs of shock (benchmark)

- systolic BP < 90 mmHg
- mean BP < 65 mmHg
- lactate > 4 mmol/l
- diuresis < 0.5 ml/kg/h
- often:
  - CI (= CO/body surface area) < 1.8 (not in septic shock)
  - HR > 100/min (not in shock with bradycardia, neurogenic shock)

# Forms of shock

- a) Hypovolemic ("cold and dry") shock low circulating volume, low preload
- b) Distributive ("warm and dry") shock low resistance, low afterload, CO might be increased
- c) Cardiogennic ("cold and wet") shock low CO in bad cardiac function, fluid congestion
- d) Obstructive shock low preload of one ventricle in normovolemia and subsequent lowering of CO + congestion – pathophysiology similar to cardiogennic shock (but congestion occurs in one half of the circulation)

## Cardiac and venous function in shock



- Hypovolemic shock: compensation by the vasoconstriction and cardiac mechanisms (but: CO is limited by low venous return)
- Distributive shock: compensation by cardiac mechanisms (vasoconstriction is usually impossible)
- Cardiogennic (and obstructive) shock: compensation by vasoconstriction

# "Interests" of the heart and perfused tissues

- Hypovolemic shock  $\downarrow$  preload
- Distributive shock ↓ afterload
- From the heart's viewpoint, ↓ preload and ↓ afterload ere advantageous, regarding the blood supply to key organs they may be linked to circulatory failure (shock states) the cause is, however, an extracardiac insult → ↓ preload or ↓ afterload (or both polytrauma)
  - But: heart must ensure its own perfusion
- Cardiac causes of shock
  - ↓ inotropy
  - $\downarrow$  lusitropy
  - ∘ ↓ HR

### Muscular work of the heart – P-V diagram:





volume

## P-V diagram in the right ventricle



# P-V diagram during changes of preload or afterload



# Inotropy and lusitropy

- ↑ inotropy (,,ability to contract") of the heart – shifts the endsystolic P-V curve up
- - The relaxation process is ATPdependent – as well as it is enabled by pumping out the cytosolic Ca<sup>2+</sup>
- ↓ inotropy or lusitropy decrease an area of P-V diagram (i.e. the cardiac work decreases – compensation by RAAS and SNS linked to an increase of preload and afterload follows)



Limit of Frank-Starling mechanism (active muscular force decreases)



Passive contraction by elastic fibres (relaxation ability decreases)

# Hypovolemic shock - causes

- Acute bleeding
- Burns, trauma
- Rapid development of ascites
- Acute pancreatitis
- Severe dehydratation
  - Vomiting, diarrhoea
  - Excessive diuresis (e.g. in diabetes insipidus)



# Acute blood loss

- Circulatory disorder (SBP < 100 mmHg, HR > 100/min) following the loss of 15% of circulating volume, shock in 30% of circulating volume
- Immediate priorities are to maintain the tissue perfusion (crystalloids, colloids) and to stop bleeding (if possible), then blood derivates (erythrocytes + plasma + thrombocytes)



## Distributive shock - causes

- Anafylactic shock
- Anafylactoid shock
  - Mediators of mast cells, but without lgE
  - E.g. snake venoms, radiocontrasts
- Septic shock
  - Role of bacterial lipopolysaccharides
  - Bacterial toxins
  - IL-I,TNF- $\alpha$  stimulate synthesis of PGE<sub>2</sub> and NO
- Neurogennic shock
  - Vasodilatation as a result of vasomotoric centre (or its efferent pahways) impairment

## Development of anaphylactic reaction

- Sensibilization of Th- and B-cells and IgE production
- Opsonization of basophils a mastocytes
  - $^\circ\,$  IgE binds to FccR (I a II)
- IgE-mediated degranulation of the mast cell and basophils following the repeated contact with an antigen
  - mediator release
    - primary (stored)– HISTAMIN (dominantly H<sub>1</sub> receptors)
    - secondary (newly formed) PG, LTA, PAF, bradykinin, cytokines, ...
  - efects
    - vazodilatation, SMC contraction (incl. bronchoconstriction), <sup>1</sup>capillary permeability, chemotaxis, <sup>1</sup>mucus secretion, platelet aggregation

### Anaphylactic and anaphylactoid reaction

### **Anaphylaxis**

- Severe, systemic, potentially life-threatening reaction following systemic exposition to an allergen
- Medication, food, insects, allergen extracts, latex
- Manifestation
  - mucous membrane, derm: erythema, exanthema, pruritus, oedema
  - resp. system: acute rhinitis, nasal obstruction, sneezing, irritation to cough, breathing problems, foreign body sensation in throat
  - GIT: vomitus, colic, diarrhoea
  - CV system: palpitation, tachycardia, hypotension, arrhythmia
  - urogenital system: urine incontinence
  - CNS: consciousness disorders, spasms
- Anaphylactoid reaction:
  - Participation of mast cell mediators, but without IgE
  - IgG, immune complexes, anaphylatoxins (C3a, C5a), myorelaxants, opiates, contrast matters, snake venoms...



# SIRS and sepsis



# Cardiogennic shock - causes

- Myocardial infarction
- Arrhythmias
- Valvular disease (e.g. rupture of papillary muscles)
- Decompensation of heart failure in dilated/restrictive cardiomyopathy, amyloidosis
- Overload by catecholamines ("tako-tsubo syndrome" apical akinesia + basal hyperkinesia)



- Rupture of ventricular septum
- Obstructive shock e.g. cardiac tamponade, massive pulmonary embolism, aortic dissection

### "Backward" acute heart failure – X-ray



Pulmonary oedema

Bilateral pleural effusion

# Organ complications in shock

- Lungs
  - ARDS
- Liver
  - necrosis of hepatocytes
- GIT
  - stress ulcer
  - $^\circ$  Damage of intestinal mucosa by ischemic necrosis  $\rightarrow$  sepsis
- Kidneys
  - Acute renal failure in vasoconstriction of a. afferens
  - Acute tubular necrosis during ischemia

### Adult Respiratory Distress Syndrome (ARDS – "shock lung")

- Result of lung inflammation in SIRS, pulmonary infections, aspiration of gastric juice, drowning
- Exsudative phase (hours): cytokine release, leukocyte infiltration, pulmonary edema, destruction of type l pneumocytes
- Proliferative phase: fibrosis, ↑ dead space, proliferation of type II pneumocytes
- Reparative phase: ↓ inflammation, ↓ edema, continuing fibrosis, in most cases permanent restrictive diseases



# Multiorgan dysfunction syndrome (MODS)

- Functional disorder of more organs at once (lungs, liver, GIT, kidneys, brain, heart)
- It can develop after initial insult (days or weeks)
- Hypermetabolism, catabolic stress
- Can both preceed or result from SIRS (primary vs. secondary MODS)
- Dysfunction  $\rightarrow$  failure

# Persistent MODS as an adaptation?

- $\downarrow$  mitochondria in tissues
- ↓ T3
- Analogy of hibernating myocardium (here, also ↓ of contractile apparatus and energy consumption)
- Gene expression similar to hibernating animals
- Later functional improvement is possible

# General principles of treatment

- Treatment of underlying cause
- Positively inotropic drugs, vasopressors (e.g. catecholamines – but: they can worsen the situation in obstructive shock)
- Colloid solutions, crystalloid solutions (but: there is a risk of oedema in cardiogenic shock)
- O<sub>2</sub>
- i.v. corticoids (anaphylaxis, SIRS?)
- ATB (septic shock)
- Mechanic circulation support (cardiogenic shock)
- Anti-shock position (?)

# Crystalloid x colloid solutions

- Crystalloid ionic solutions (best normochloremic)
  - They do not induce allergic reactions or alter coagulation
- Colloids high molecular weight compounds (hydroxyethylstarch, gelatine, albumin)
  - Fluid distribution points more to intravascular compartment
    - But less than is expected theoretically damaged glycocalyx defines water reabsorption



# Mechanical circulatory support



# ECMO: Kaplan-Meier curves



www.jtcvs.org/article/S0022-5223(18)30906-1/fulltext

# Trendelenburg ("anti-shock") position

- 15-30°
- ↑ Venous return
- After collapse
- Inefficient in the long term
- Central venous catheter insertion (circulatory support administration)
- Worsens pulmonary ventilation

