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# **Pulmonary Blood Flow**

M1 – Cardiovascular/Respiratory Sequence Thomas Sisson, MD



Winter, 2009

## **Objectives**

- The student will know the structure, function, distribution and control of pulmonary blood supply
  - Compare pulmonary and bronchial circulation
  - Compare and contrast pulmonary and systemic circulation
  - Describe and explain the effects of cardiac output and lung volume on pulmonary vascular resistance
  - Describe the effects of hypoxia on pulmonary vascular resistance
  - Describe the effects of gravity of pulmonary blood flow
  - Explain Starling's equation
  - Describe the mechanisms of pulmonary edema

## **Two Circulations in the Lung**

Pulmonary Circulation.
 Arises from Right Ventricle.
 Receives 100% of blood flow.

- Bronchial Circulation.
  - Arises from the aorta.
  - Part of systemic circulation.
  - Receives about 2% of left ventricular output.

## **Bronchial Circulation**



# **Pulmonary Circulation**



United States governmen



PD-INEL Source Undetermined

# **Pulmonary Circulation**





## **Pulmonary Circulation**

- In series with the systemic circulation.
- Receives 100% of cardiac output (3.5L/min/m<sup>2</sup>).
- RBC travels through lung in 4-5 seconds.
- 280 billion capillaries, supplying 300 million alveoli.
  - Surface area for gas exchange =  $50 100 \text{ m}^2$

## **Alveolar Architecture**



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## Functional Anatomy of the Pulmonary Circulation

- Thin walled vessels at all levels.
- Pulmonary arteries have far less smooth muscle in the wall than systemic arteries.
- Consequences of this anatomy the vessels are:
  - Distensible.
  - Compressible.

## **Pulmonary Circulation Pressures**



## **Pulmonary Vascular Resistance**

Vascular Resistance =

input pressure - output pressure

blood flow

PVR = k • mean PA pressure - left atrial pressure cardiac output (index)

mean PA pressure - left atrial pressure = 10 mmHg

mean aorta pressure - right atrial pressure = 98 mmHg

Therefore PVR is 1/10 of SVR

# Vascular Resistance is Evenly Distributed in the Pulmonary Circulation



#### Reasons Why Pressures Are Different in Pulmonary and Systemic Circulations?

#### • Gravity and Distance:

- Distance above or below the heart adds to, or subtracts from, **both** arterial and venous pressure
- Distance between Apex and Base

| Systemic |          | Pulmonary |         |
|----------|----------|-----------|---------|
| Aorta    | 100 mmHg | Main PA   | 15 mmHg |
| Head     | 50 mmHg  | Apex      | 2 mmHg  |
| Feet     | 180 mmHg | Base      | 25 mmHg |

#### Reasons Why Pressures Are Different in Pulmonary and Systemic Circulations?

- Control of regional perfusion in the systemic circulation:
  - Large pressure head allows alterations in local vascular resistance to redirect blood flow to areas of increased demand (e.g. to muscles during exercise).
  - Pulmonary circulation is all performing the same job, no need to redirect flow (exception occurs during hypoxemia).
- Consequences of pressure differences:
  - Left ventricle work load is much greater than right ventricle
  - Differences in wall thickness indicates differences in work load.

#### Influences on Pulmonary Vascular Resistance

Pulmonary vessels have:

- -Little vascular smooth muscle.
- -Low intravascular pressure.
- -High distensiblility and compressiblility.

#### Vessel diameter influenced by extravascular forces:

- -Gravity
- -Body position
- -Lung volume
- -Alveolar pressures/intrapleurql pressures
- -Intravascular pressures

#### **Influences of Pulmonary Vascular Resistance**

- •Transmural pressure = Pressure Inside Pressure Outside.
  - -Increased transmural pressure-increases vessel diameter.
  - –Decreased transmural pressure-decreased vessel diameter (increase in PVR).

-Negative transmural pressure-vessel collapse.



•Different effects of lung volume on <u>alveolar</u> and <u>extraalveolar</u> vessels.

#### Effect of Transmural Pressure on Pulmonary Vessels During Inspiration



## **Effect of Lung Volume on PVR**



## Pulmonary Vascular Resistance During Exercise

 During exercise cardiac output increases (e.g. 5-fold), but with little change in mean pulmonary artery pressure

 How is this possible?

Vascular Resistance =

input pressure - output pressure

blood flow

- ΔPressure= Flow x Resistance
- If pressure does not change, then PVR must decrease with increased blood flow
  - Passive effect (seen in isolated lung prep)
  - Recruitment: Opening of previously collapsed capillaries
  - Distensibility: Increase in diameter of open capillaries.

## **Recruitment and Distention in Response to Increased Pulmonary Artery Pressure**



## **Control of Pulmonary Vascular Resistance**

#### Passive Influences on PVR:

| Influence                        | Effect on PVR                    | Mechanism                               |
|----------------------------------|----------------------------------|---|
| ↑ Lung Volume (above<br>FRC)     | Increase                         | Lengthening and<br>Compression          |
| ↓ Lung Volume (below<br>FRC)     | Increase                         | Compression of<br>Extraalveolar Vessels |
| ↑ Flow, ↑Pressure                | Decrease                         | Recruitment and<br>Distension           |
| Gravity                          | Decrease in Dependent<br>Regions | Recruitment and<br>Distension           |
| ↑ Interstitial Pressure          | Increase                         | Compression                             |
| Positive Pressure<br>Ventilation | Increase                         | Compression and<br>Derecruitment        |

#### Regional Pulmonary Blood Flow Depends Upon Position Relative to the Heart



#### **Gravity, Alveolar Pressure and Blood Flow**

- Pressure in the pulmonary arterioles depends on both mean pulmonary artery pressure and the vertical position of the vessel in the chest, relative to the heart.
- Driving pressure (gradient) for perfusion is different in the 3 lung zones:
  - Flow in zone may be absent because there is inadequate pressure to overcome alveolar pressure.
  - Flow in zone 3 is continuous and driven by the pressure in the pulmonary arteriole – pulmonary venous pressure.
  - Flow in zone 2 may be pulsatile and driven by the pressure in the pulmonary arteriole – alveolar pressure (collapsing the capillaries).

#### Gravity, Alveolar Pressure, and Blood Flow



**PD-INEL** West. Respiratory Physiology: The Essentials 8<sup>th</sup> ed. Lippincott Williams & Wilkins. 2008

Typically no zone 1 in normal healthy person Large zone 1 in positive pressure ventilation + PEEP

# **Gravity Influences Pressure**



(O) Contraction Adrian8\_8 (flickr)

## **Control of Pulmonary Vascular Resistance**

#### Active Influences on PVR:

#### Increase

#### Sympathetic Innervation

 $\alpha$ -Adrenergic agonists

Thromboxane/PGE2

Endothelin

Angiotensin

Histamine

Alveolar Hypoxemia

#### Decrease

Parasympathetic Innervation

Acetylcholine

 $\beta$ -Adrenergic Agents

PGE1

Prostacycline

Nitric oxide

Bradykinin

#### **Hypoxic Pulmonary Vasoconstriction**

- Alveolar hypoxia causes active vasoconstriction at level of precapillary arteriole.
- Mechanism is not completely understood:
  - Response occurs locally and does not require innervation.
  - Mediators have not been identified.
  - Graded response between pO2 levels of 100 down to 20 mmHg.
- Functions to reduce the mismatching of ventilation and perfusion.
- Not a strong response due to limited muscle in pulmonary vasculature.
- General hypoxemia (high altitude or hypoventilation) can cause extensive pulmonary artery vasoconstriction.

## **Barrier Function of Alveolar Wall**

- Capillary endothelial cells:
   permeable to water, small molecules, ions.
   barrier to proteins.
- Alveolar epithelial cells:
  - more effective barrier than the endothelial cells.
  - recently found to pump both salt and water from the alveolar space.



#### **Fluid Movement Due to Osmotic Pressure**



Water moves through the semi-permeable membrane down a concentration gradient to dilute the solute.

## Osmotic Pressure Gradient Can Move Fluid Against Hydrostatic Pressure



## Osmotic Gradient Counteracts Hydrostatic Gradient

- Hydrostatic pressure in the pulmonary capillary bed > hydrostatic pressure in the interstitium
  - hydrostatic pressure drives fluid from the capillaries into the pulmonary interstitium
- Osmotic pressure in the plasma > osmotic pressure in the interstitium
  - osmotic pressure normally would draw fluid from the interstitial space into the capillaries

## **Starling's Equation**



 $\begin{array}{l} \mathsf{Q} = \mathsf{flux} \ \mathsf{out} \ \mathsf{of} \ \mathsf{the} \ \mathsf{capillary} \\ \mathsf{K} = \mathsf{filtration} \ \mathsf{coefficient} \\ \mathsf{Pc} \ \mathsf{and} \ \mathsf{Pi} = \mathsf{capillary} \ \mathsf{and} \ \mathsf{interstitial} \ \mathsf{hydrostatic} \\ & \mathsf{pressures} \\ \pi \mathsf{c} \ \mathsf{and} \ \pi \mathsf{i} = \mathsf{capillary} \ \mathsf{and} \ \mathsf{interstitial} \\ \mathsf{osmotic} \qquad \mathsf{pressures} \\ \sigma = \mathsf{reflection} \ (\mathsf{sieving}) \ \mathsf{coefficient} \end{array}$ 

## Normally Starling's Forces Provide Efficient Protection

- Normal fluid flux from the pulmonary capillary bed is approximately 20 ml/hr.
  - recall that cardiac output through the pulmonary capillaries at rest is ~5 l/min.
  - < 0.0066% leak.
- Abnormal increase in fluid flux can result from:
  - Increased hydrostatic pressure gradient (cardiogenic pulmonary edema).
  - Decreased osmotic pressure gradient (cirrhosis, nephrotic syndrome).
  - Increased protein permeability of the capillary wall (ARDS).

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