RESPIRATORY PHYSIOLOGY – Part 2
ANAESTHESIA TUTORIAL OF THE WEEK 160

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QUESTIONS

Before continuing, try to answer the following questions. The answers can be found at the end of the article, together with an explanation.

1. The oxyhaemoglobin dissociation curve is shifted to the left by:
   a) an increase in arterial PCO2
   b) acidosis
   c) a decrease in 2,3 DPG
   d) carbon monoxide
   e) a fall in temperature

2. Anatomy of the respiratory tract
   a) the alveoli humidify gas
   b) generations 1-18 are known as the conducting zone of the lungs
   c) there are 25 bronchial generations
   d) bifurcation of the bronchi occurs at T4
   e) the lungs are lined with transitional epithelium

3. Dead space
   a) is defined as the volume of gas which does take part in gas exchange
   b) physiological dead space is the same as alveolar dead space
   c) anatomical dead space is measured by the Bohr equation
   d) dead space is usually about 45% of tidal volume
   e) occurs with pneumonia

4. Pulmonary vascular resistance
   a) is high at high lung volumes
   b) is low at low lung volumes
   c) can cause heart failure
   d) is calculated using PCWP
   e) is decreased by nitrous oxide

5. Diffusion of gases
   a) is increased with a decrease in surface area
   b) is indirectly proportional to the concentration gradient
   c) is not governed by Grahams law
   d) is directly proportional to the thickness
   e) is decreased according to Beers law
6. Gas transport in the blood
   a) there is about 15ml of oxygen per 100ml of oxygenated blood
   b) oxygen CO2 is mainly transported as carbamino compounds
   c) CO2 is carried best by oxygenated haemoglobin
   d) More oxygen is carried dissolved in blood than CO2
   e) CO2 is transported in the blood by 3 main routes

PERFUSION OF THE LUNG

Blood flow to and around the lung is similar to any other organ but at much lower pressures than the systemic system (see diagram). The blood vessels in the lungs continually branch and get consistently smaller very like the branching of the airways. The pulmonary arteries whose walls are very thin in comparison to that of the arteries in the main circulation feed the lung up to the level of the terminal bronchioles and then split into the capillary bed. The capillaries have great capability to distend thus enhancing gas exchange and reservoir action. Once the red blood cells have become oxygenated the capillary bed is drained into venules which then join to form the pulmonary veins. It is the ability of the blood vessels to distend and be recruited which allows the pressures in the pulmonary system to stay low despite very high blood flow.

The pulmonary arteries only supply blood flow and oxygen to the lungs and must have the ability to accept huge blood volumes at times. The low pulmonary pressures are important to minimise the work of the right heart.

Figure 6: The pressures within the main and pulmonary circulatory systems
PULMONARY VASCULAR RESISTANCE

\[ PVR = 80 \times (MPAP - PCWP) / CO \]

**Acute and chronic lung disease**

Both can cause an increase in PVR. Chronic PVR can lead to right sided heart failure

**Lung volumes**

PVR depend on the alveolar and extra-alveolar resistances. Both low and high lung volumes will increase PVR, it is at its lowest at FRC.

Alveolar vessels or pulmonary capillaries will become overly stretched longitudinally at high lung volumes and therefore their resistance will increase.

With extra – alveolar vessels resistance will increase at low lung volumes since they are not encouraged to expand by lung parenchyma.

PVR can also be altered through the distension and recruitment of pulmonary vessels which have a large ability to increase their capacitance.

**Hypoxic pulmonary vasoconstriction**

In response to episodes of low partial pressure of alveolar oxygen the lung will divert its own blood supply to areas of properly ventilated lung tissue. This is important in matching and maintaining ventilation to perfusion in the lung. HPV is also obtunded by the general anaesthetic inhalational agents, it is relatively preserved by TIVA.

**Metabolic substances**

Endothelin – released from the lung endothelial cells in response to tissue damage – is a very potent vasoconstrictor

Prostacyclin – a naturally occurring vasodilator

Nitric oxide – Also known as endothelin derived relaxing factor – a potent vasodilator

**Perfusion throughout the lungs**

Taking all of the above into account it is easy to understand then why blood flow is not always even throughout the lung. Gravity plays a large part in directing blood flow by setting up a hydrostatic pressure gradient which is higher at the base of the lung than at the top. Blood is preferentially directed to the lung bases.

As well as the affect of gravity on perfusion and ventilation, the differing pressures within the alveoli, arteries and venous systems heavily influences the outcome. The relationship between these factors describes West zones.

West zones within the lung are 3 vertically split zones (in the upright subject) which explain how alveoli, arterial and venous pressures differ in each zone and thus affect perfusion and ventilation throughout the lung.

In zone 1 the alveolar pressure may exceed that of the arterial and venous pressure and thus little perfusion will occur as the vessels collapse, this then leads to dead space.

In zone 2 the arterial pressure will exceed that of the alveoli pressure but not the venous pressure.

In zone 3 both the arterial and venous pressures exceed the alveoli pressure.
**Key points**

- The lung is a low pressure system
- Low pulmonary pressure minimises the work of the right heart
- The pulmonary vessels distend and recruit to maintain low pressures
- Very high and very low lung volumes increase pulmonary vascular resistance
- Hypoxic pulmonary vasoconstriction diverts blood flow away from poorly ventilated regions of lung
- HPV is obtunded by anaesthesia

**Clinical points**

- Ventilatory strategies can prevent increases in pulmonary pressures e.g. low tidal volumes, permissive hypercapnia
- TIVA does not affect HPV

**GAS EXCHANGE**

The partial pressure of oxygen that is inhaled from our natural environment through normal inhalation is not maintained at the same partial pressure by the time it reaches the alveoli and indeed the mitochondria. The process by which this decrease in partial pressure occurs is called the oxygen cascade.

The air surrounding us – how much oxygen does it contain?

Dry atmospheric air gas – 21% of 100KPa

Therefore 21 KPa or 160mmHg

However as gas is inspired it is diluted by water vapour which reduces the partial pressure of oxygen

Water vapour – 6.3 KPa/ 47mmHg

\[
\text{PO}_2 = 0.21 \times (760-47) = 149\text{mmHg}
\]

\[
\text{PO}_2 = 0.21 \times (100-6.3) = 19.8\text{KPa}
\]
When the gas reaches the alveoli the partial pressure of oxygen will again decrease as some oxygen is absorbed and CO2 is excreted. The partial pressure at this point in the oxygen cascade can be determined by using the alveolar gas equation.

\[ \text{PAO}_2 = \text{P}O_2 - \frac{\text{PACO}_2}{\text{RQ}} \]

The RQ stands for respiratory quotient and is normally 0.8. It is determined by the amount of CO2 produced/oxygen consumed.

\[ \text{PAO}_2 = 0.21 \times 5/0.8 = 14 \text{KPa} \ (106 \text{mmHg}) \]

Again when the gas reaches the arterial blood a further small drop in partial pressure will have occurred as blood known as venous admixture with a lower oxygen content mixes with the oxygenated alveolar blood.

Venous admixture is made up of blood that has passed through poorly ventilated regions of lung and thus has a lower O2 partial pressure. Venous admixture is also composed of venous blood which has drained the lungs and left side of the heart. This blood is known as true shunt and drains directly into the left side of the heart.

Extraction of oxygen from this blood then causes the end capillary oxygen partial pressure to be 6-7KPa (40 - 50mmHg)

In the mitochondria the PO2 varies hugely from 1-5KPa (7.5-40mmHg)

This provides us with an explanation for the following graph, the oxygen cascade.

![Figure 3](image)

**Figure 8:** Diffusion of gases from alveoli to blood – the oxygen cascade

**Diffusion of Gases Across the Alveolar Membrane**

The speed and ease of diffusion are controlled by the laws of diffusion.

*Fick’s law* of diffusion states that gas transfer across a membrane is directly proportional to the concentration gradient.

*Graham’s law* states that diffusion of a gas is inversely proportional to the square root of the molecular weight of the molecule.

Other factors which increase diffusion:
- Large surface area
- Thin membrane
- High solubility
The following equation incorporates the important factors

Diffusion is proportional to A/T. D. (P1 – P2)

A = Area
T = Thickness
D = Diffusion constant
P1 – P2 = Concentration gradient

Diffusion in the lungs can be limited in the presence of disease states e.g pulmonary oedema and thickening of the alveolar membrane in pulmonary fibrosis.

**Oxygen transport**

Oxygen is carried in 2 forms in the blood:

- Oxygen combined to haemoglobin (97%)

Haemoglobin molecule consists of 2 alpha and 2 beta chains; each chain is formed from an iron–porphyrin molecule - haem. Each haemoglobin molecule can bind 4 oxygen molecules (20ml oxygen per 100ml blood) or 15ml oxygen per 100ml in venous blood

- Oxygen dissolved in the blood – this accounts for a minimal amount (0.3ml per dl)

The amount dissolved obeys Henrys’ law – amount is proportional to the partial pressure 0.023ml per KPa per 100ml blood

**Oxygen content in the blood**

Total content of oxygen in the blood can be calculated from the Oxygen flux equation:

\[
\text{Flux} = (CO \times Hb \times \text{Saturation} \times \text{Huffners constant}(1.39)) + (0.023 \times PO2)
\]

**Oxygen dissociation curve**

- Sigmoid shaped curve relating the fact that binding of oxygen to the haemoglobin molecule is a cooperative process
- Describes the relationship of saturation of haemoglobin with oxygen at varying partial pressures
- Be aware of the P50 –(point at which Hb is 50% saturated)
- Decreasing pH, increasing temperature, 2,3-DPG and CO2 tension will cause a right shift of the curve
- Increased pH, and reduction in CO2 tension, temperature and 2,3-DPG produce a left shift of the curve
- If a right shift occurs the Hb molecule is more likely to offload oxygen to the tissues
- In a left shifted situation the Hb is less likely to release oxygen to the tissues
FIGURE 9: The Oxyhaemoglobin dissociation curve

2,3-DPG
This molecule binds to deoxygenated Hb – it reduces the affinity of haemoglobin for oxygen and therefore ensures offloading of oxygen to the tissues.

The Bohr Effect
This describes the affect that CO\textsubscript{2} has on influencing the release of oxygen to the tissues. On entering red blood cells the following reaction occurs:

\[
\text{CO}_2 + \text{H}_2\text{O} \leftrightarrow \text{H}_2\text{CO}_3 \leftrightarrow \text{H}^+ + \text{HCO}_3^- 
\]

An increase in H\textsuperscript{+} will cause an acidosis and therefore encourage the release of oxygen from Hb. In the lungs where the CO\textsubscript{2} is being removed, the alkalosis will encourage the uptake of oxygen.

Oxygen Delivery (DO\textsubscript{2})
This can be calculated as follows:

Cardiac output X arterial O\textsubscript{2} content

Carbon dioxide transport
Carbon dioxide is carried in the blood in 3 ways:

- As bicarbonate – 90%
- As dissolved CO\textsubscript{2} – 5%
- As carbamino compounds – 5%
Carbamino compounds are formed by the reaction of the CO$_2$ with terminal amino groups of proteins and side chains of arginine and lysine. Haemoglobin is essential for this process to occur since it has 4 amino groups per molecule. Albumin also provides amino groups but only 1 per molecule.

**The Haldane effect**

This phenomenon refers to the increased ability of blood to carry CO$_2$ when haemoglobin is deoxygenated. Deoxyhaemoglobin is 3.5 times more effective than oxyhaemoglobin in forming carbamino compounds.

**Figure 10**: Carbon dioxide transport in blood

**Ventilation perfusion Relationships**

As mentioned earlier any discrepancy between ventilation and blood flow in the lung will result in V/Q mismatch and potentially dangerous irregularities in gas exchange. If flow of blood to the lung units is to match that of ventilation to the same unit then the ratio of ventilation to perfusion should be in a ratio of 1:1

If the lung is being underventilated but perfused as normal then we say that the V/Q ratio is <1
If the lung is underperfused then the V/Q is >1

Even in a normal lung the V/Q ratio is not uniformly 1 throughout the lung as perfusion and ventilation both have favoured parts of the lung.

Differences between the apices and bases of the lungs

At the apices there is less ventilation than the bases as alveoli are already very stretched however there is proportionally less perfusion therefore the overall V/Q ratio is higher compared to the base of the lung.

Blood flow is directly affected by gravity and naturally has a tendency to flow to the bases of the lungs thus V/Q ratios towards the lower segments of the lung are usually greater than 1.

The vertical change in V/Q ratios in the lung is because although both ventilation and perfusion increase from top to bottom of the lung, perfusion increases much quicker than ventilation. Thus the V/Q ratio at the top of the lung is 3.3 whereas at the bases it is around 0.6. See the following illustration (figure 11).
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**Figure 5-8.** Distribution of ventilation and blood flow down the upright lung (compare Figures 2-7 and 4-7). Note that the ventilation-perfusion ratio decreases down the lung.

**Figure 5-9.** Result of combining the pattern of ventilation-perfusion ratio inequality shown in Figure 5-8 with the effects of this on gas exchange as shown in Figure 5-7. Note that the high ventilation-perfusion ratio at the apex results in a high $P_{O_2}$ and low $P_{CO_2}$ there. The opposite is seen at the base.

**Figure 11:** Ventilation perfusion relationships in the lungs
Alveolar – Arterial P0₂ gradient

The value for the A-a gradient gives the clinician some idea about the amount of VQ mismatch and shunt that is present in the lungs. A typical normal value would be around 0.5-1 KPa (5mmHg) though values up to around 15mmHg may be accepted.

It is calculated as PAO₂ – PaO₂. The PAO₂ is calculated using the alveolar gas equation.

Shunt

True shunt refers to a VQ = 0. That is to say that blood has passed through areas of the lung where no ventilation is occurring. As discussed earlier VQ mismatch is also referred to as shunt. Blood passes through areas of the lung which are poorly ventilated ie VQ<1.

Physiological shunt refers to the amount of venous admixture which is directly added to main circulatory blood without having passed through the oxygenating mechanism of the lung. Blood from the bronchial veins draining the lung parenchyma and the thebesian veins draining the cardiac muscle represent the physiological shunt (around 5% of cardiac output.)
The shunt equation allows calculation of the amount of shunt present in an individual subject.

The Shunt Equation:

\[
\frac{Q_s}{Q_t} = \frac{Cc'_{O_2} - Ca_{O_2}}{Cc'_{O_2} - Cv_{O_2}}
\]

Figure 12: A visualization of the shunt equation

The amount of oxygen leaving the lungs is Qt x CaO₂. This is equal to the shunted blood flow plus the oxygen content from the lung which would be (Qs x CvO₂) + (Qt-Qs) x CcO₂. (shunt flow x mixed venous O2 content + pulmonary capillary flow x pulmonary capillary O2 content).

When these equations as rearranged it provides the classic shunt equation:

\[
\frac{Q_s}{Q_t} = \frac{Cc'O_2 - CaO_2}{Cc'O_2 - CVO_2}
\]
Key points

- The amount of oxygen inhaled is greatly different from the amount reaching the mitochondria
- The oxygen cascade describes the fall in oxygen as it passes from the air to the mitochondria
- Diffusion of gas is affected by molecular size, surface area and thickness of diffusion barrier and concentration gradient of the gas
- Oxygen content in the blood can be calculated by the flux equation
- CO2 is mainly carried as bicarbonate in the blood
- Deoxygenated haemoglobin is more effective at carrying CO2 than oxygenated haemoglobin

Clinical points

- CO, Hb saturation and haemoglobin concentration are the prime factors affecting oxygen delivery
- These factors can be manipulated by the anaesthetist to optimise a patient pre-operatively
- Insertion of haemodynamic monitoring, fluid therapy, transfusion, supplemental oxygen therapy and the correct use of inotropes are the key components in optimisation strategies to improve oxygen delivery

ANSWERS TO MCQS

1. FFTTT
The oxyhaemoglobin dissociation curve is shifted to the left with a decrease in C0₂, 2,3 DPG, temperature and an alkalosis. Carbon monoxide prevents haemoglobin from releasing oxygen to the tissues therefore it causes a shift in the curve to the left.

2. FFFFTF
The nose and mouth act to filter and humidify gases as they flow into the lungs. There are 23 bronchial generations, 1-16 are the conducting zone and 17-23, the respiratory zone. The lungs are lined with columnar ciliated epithelium.

3. FFFFT
Dead space is the volume of gas which does not take part in gas exchange. It is classified into Anatomical, avelore and physiological which is a combination of both and can account for about 30% of tidal volume. Anatomical dead space is measured via Fowlers method and Physiological by the Bohr equation.

4. TFTTF
PVR (pulmonary vascular resistance) = 80 x (MPAP - PCWP) / CO and is minimised at FRC and high at both low and high lung volumes. If it becomes a chronic problem it can lead to right sided lung failure. It is decreased by nitric oxide and prostacyclin - both have vasodilatory effects.

5. FFFFF
Diffusion = Area/Thickness . (P1- P2) Diffusion has been best described by Fick and Grahams law.

6. FFFFF
There is 15ml of oxygen on venous blood and 20ml per 100ml in oxygenated blood. O2 is carried mainly by haemoglobin whereas CO2 is transported as bicarbonate, caramino compounds or dissolved in blood. The Haldane effect describes deoxygenated blood carrying CO2 better than oxygenated blood due to its effectiveness in forming carbamino compounds.
WEB LINKS

www.frca.co.uk

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