The Dual Circulation of the Lungs and Their Connections

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SUMMARY

The bronchial and pulmonary circulations are outlined and their relevance to certain pathological conditions is conjectured.

INTRODUCTION

The main function of the lung is respiration. Here, the distribution of the blood is just as important as the distribution of the air. It could be said that disturbance in the ventilation/perfusion balance is probably the most common cause of respiratory failure in patients with lung disease (Meyer, 1976). Knowledge of the bronchial and pulmonary circulation, and their different reactions to hypoxia, is thus very important in understanding the way in which certain existing communications between the two systems come into play and the effect thereof under certain pathological conditions.

ANATOMY

Bronchial and pulmonary circulation

The lung is supplied by a small bronchial and a large pulmonary arterial system. The bronchial arteries are usually direct branches from the descending aorta (usually one on the right and two on the left), but there may be variations in their origin. They supply the smooth muscle of the bronchi from the carina to the respiratory bronchioli and also nourish the connective tissue in this area as well as certain parts of the visceral pleura (Last, 1972). The middle segment of the oesophagus, a part of the vagus nerve and the tracheobronchial lymph nodes are also supplied by the bronchial arteries. In addition, there are vasa vasorum in close contact with and supplying the pulmonary arteries.

The bronchial arteries divide into numerous branches before reaching the respiratory bronchioli — the so-called broncho-pulmonary branches. These branches divide further into lobular and eventually into bronchial capillaries which join up with the pulmonary capillaries. Together they drain into the pulmonary veins. The bronchial capillaries usually supply the bronchioli although the pulmonary arteries or even the pulmonary veins may also perform this task (Tobin, 1952). The bronchial arteries are part of the systemic circulation and their blood pressure (120/80 mmHg) is therefore more or less five times greater than that of the pulmonary arteries (25/10 mmHg). The bronchial arteries tend to spiral around the bronchioles and are thus arranged in a dense communicating network. This plexiform arrangement of the bronchial arteries is the reason why the systemic arterial supply to a bronchus is not easily cut off (Liebow, 1949).

The bronchial veins also form a network around the bronchi. The veins which drain the first part of the bronchi (extrapulmonary) drain into the right atrium via the azygos (right) and hemiazygos (left) veins. The rest of the bronchial veins enter the pulmonary veins on their way to the left atrium and their contents make up more or less 2% of the cardiac output (Meyer, 1976). These bronchial venous plexi may anastomose with the venous (pulmonary) capillaries around the alveoli, or they may connect by branches to the pulmonary veins (post-capillary), or they may even anastomose with adjacent parts of the bronchial venous plexi so that the blood may drain upwards into the azygos system (Tobin, 1952).

The pulmonary trunk which gives rise to the pulmonary arteries originates from the right ventricle and bifurcates, sending one branch to each lung. The pulmonary arteries have a rectilinear course roughly parallel to the branches of the respiratory tree. They are true end-arteries, connecting only in the finer capillary networks (Liebow, 1949). The pulmonary arterioles and...
capillaries are short and their diameters relatively wide so that there is a decreased resistance to blood flow and this system is thus one of passive low pressure. The capillaries form a rich plexus around each alveolus and they supply the alveoli with all their needs, excluding oxygen. When a bronchus is obstructed, the pulmonary vessels supplying the poorly ventilated alveoli react to the hypoxia by constricting and shunting the blood to other areas. (The bronchial arteries dilate with hypoxia). Emboli that block small branches of the pulmonary artery provoke a marked rise in pulmonary arterial pressure, although this is absent when larger arteries are blocked (Ganong, 1973), but there is some controversy about this. There are usually two pulmonary veins from each lung, carrying oxygenated blood to the left atrium (Last, 1972). The blood is not 100% saturated with oxygen as there is mixing with venous blood from the bronchial veins.

The pulmonary capillary pressure is about 10 mmHg and the oncotic pressure about 25 mmHg. There is thus an inward-outward pressure gradient of about 15 mmHg which keeps the alveoli free of fluid. If the pulmonary capillary pressure exceeds 25 mmHg (failure of left ventricle), this gradient will shift and oedema and pulmonary congestion will develop (Ganong, 1973).

**Plural fluid**

Plural fluid moves into the pleural cavity under the influence of the parietal plural layer and is absorbed by the visceral layer.

Fluid entering the pleural space depends on the following pressure gradients:

**Parietal plural layer**

Factors causing fluid to enter the pleural space:

- Hydrostatic pressure (in arterial capillary) = 30 cm H₂O
- Oncotic pressure (in capillary) = 34 cm H₂O
- Negative pleural pressure = 5 cm H₂O

Total = 35 cm H₂O

Factors keeping fluid out of the pleural space:

- Oncotic pressure (in pleural fluid) = 8 cm H₂O
- Oncotic pressure (in capillary) = 34 cm H₂O

Total = 26 cm H₂O

The force responsible for moving the fluid into the cavity is thus 35 cm H₂O and the force for absorbing the fluid is only 26 cm H₂O. There is thus a resultant force of 9 cm H₂O in the direction of the pleural cavity.

**Visceral plural layer**

The fluid absorbed by the visceral layer depends on the following pressure gradients:

Factors keeping fluid in pleural space:

- Hydrostatic pressure (in capillary) = 30 cm H₂O
- Oncotic pressure (in arterial capillary) = 30 cm H₂O

Total = 60 cm H₂O

Factors for absorbing fluid:

- Oncotic pressure (in capillary) = 34 cm H₂O
- Oncotic pressure (in pleural fluid) = 8 cm H₂O

Total = 42 cm H₂O

The total force is thus 10 cm H₂O in the direction of the visceral layer and the fluid is absorbed.

**Lung zones**

The transmural pressure of the pulmonary arteries is the result of the perivascular pressure and the intravascular pressure. This resultant force is dependent on gravity; in the upright position the blood pressure is maximum at the lung bases. The vessels are therefore dilated at the bases and narrower at the apices. The capillaries in the apex (Zone I) are thus the first to collapse if the capillary pressure should decrease below that of the alveolar pressure. This does not usually occur under normal conditions, but when there is a fall in arterial pressure (e.g. haemorrhage) or a rise in the alveolar pressure (e.g. the use of positive pressure ventilation) it may happen. In the middle lung regions (Zone II) the arterial capillary pressure is greater than the alveolar pressure and the alveolar pressure is greater than the venous capillary pressure. Therefore it is less likely that the alveolar capillaries in this area will collapse than in Zone I. The blood flow in this area is dependent on the arterial-alveolar gradient (not the usual arterial-venous gradient). In Zone III (bases) no collapse of capillaries takes place, as the capillary arterial pressure is greater than that of the alveolar pressure and the alveolar pressure is smaller than that of the venous capillary pressure. Here the blood flow is dependent on the arterial-venous gradient (Meyer, 1976 and West, 1974).

**Arterio-arterial and arterio-venous communications**

It is generally accepted that there is some overspill of bronchial capillaries into alveolar (pulmonary) capillaries (Last, 1972). However, it has recently also been proved that precapillary anastomoses between bronchial arteries and pulmonary arteries do exist. There are two types of these anastomoses, namely the short (1-2 mm) and narrow (50-100 micron) type which lies peripherally, and the long (10-40 mm), wide (300-400 micron) type which are more centrally situated. These anastomoses have coiled, thick muscle walls which reduce the blood pressure from the bronchial to the pulmonary arterial system (Ganong, 1973). During deep inspiration, the decreased pressure in the pulmonary artery will cause more blood to flow into it from the bronchial arteries through the precapillary arterio-arterial shunts (Pump, 1972).

Obstruction of the bronchial arteries themselves may lead to necrosis of the extrapulmonary bronchi but not of the intrapulmonary bronchi because of their connections with other blood vessels (Tobin, 1952).

Apart from these arterio-arterial anastomoses, there are also arterio-venous connections between the pulmonary arteries and the pulmonary veins. Anatomically it has been seen that there is an arterial loop at the apex of the acinus and that this loop has very thick muscle or elastic tissue in its wall. This segment may be dilated to form glomus-like protrusions or to allow the passage of glass spheres, 200 μ in diameter and many times the accepted diameter of the capillaries, into the pulmonary veins (Tobin, 1966). In the foetus these arterio-venous shunts may play a role in diverting the blood from the pulmonary arteries (and their connections with the bronchial arteries) into the pulmonary veins. The unexpanded alveoli are thus avoided (Tobin, 1952). In normal, resting subjects with no intracardiac shunts, this arterio-venous pulmonary shunt flow averages 1% of the total pulmonary blood flow. When using the Valsalva technique on patients, there is a rise in oxygen saturation during the manoeuvre, and a fall in saturation following release of this technique. The saturation rise average 1.2% in normal people; in patients with cardiac or pulmonary disease this value is even higher, namely 3.9-7.5%. It is supposed that this manoeuvre leads to a fall in transmural pressure in the pulmonary artery with a decreased blood flow in the pulmonary arterial as well as in the arterio-venous shunts. The result is that more blood flows through the capillaries and that more blood becomes oxygenated, which causes the rise in oxygen saturation. This does not, however, always apply to patients with con-
gestive heart failure in whom the Valsalva technique might have no effect on transmural pressure of the pulmonary artery (José and Milner, 1959).

PATHOLOGICAL CONDITIONS

Pulmonary embolism
When a pulmonary artery and vein of a lobe in a dog's lung are both occluded, the vascular pressures within that lobe rise to abnormally high levels (Shedd et al., 1951). The reason for this is that the pulmonary artery responds to the hypoxia by constricting and shunting the blood away from that area, while the bronchial artery (like any other systemic artery) responds by dilating. An extensive opening of the precapillary anastomoses takes place between the bronchial and pulmonary arteries. The high blood pressure of the bronchial arteries is thus transmitted to the pulmonary arteries. The increased hydrostatic pressure of the pulmonary arteries is no longer 11 cm H₂O, but may be 36 cm H₂O or more (Nagaishi, 1972), and the pulmonary arteries supplying the region might have no effect on transmural pressure of the pulmonary artery (Jose and Milner, 1959). The reason for this is that the pulmonary arteries is no longer 11 cm H₂O, but may be 36 cm H₂O or more (Nagaishi, 1972). However, if portal hypertension is present, the pressure in the azygos system will rise and the pressure in the bronchial veins that drain into the azygos system may be increased to such an extent that the pulmonary-bronchial post-capillary venous anastomoses may open. This will lead to an increased hydrostatic pressure in the pulmonary capillaries with decreased absorption of pleural fluid and thus hydrothorax may occur.

Heart failure
Right-sided heart failure is usually secondary to left-sided failure. When the right ventricle begins to fail systemic venous congestion occurs. Most of the bronchial veins drain into the right atrium and the increased pressure in these leads to an increased pressure in the pulmonary veins of the visceral pleura (through their anastomoses). The pleural fluid is not absorbed and a pleural effusion occurs.

CONCLUSION
It is generally accepted that there are connections between the systemic and pulmonary arterial systems in the lungs. These can be at intercapillary or precapillary levels. As regards the precapillary shunts, the mechanisms whereby they open are most important in understanding certain lung conditions, e.g. atelectasis, pulmonary embolus, pneumonia. These mechanisms prevent necrosis of the associated lung parenchyma and explain the phenomenon of local pleural effusions. Post-capillary (veno-venous) anastomoses are responsible for the pleural effusions seen in liver cirrhosis and right-sided heart failure.

Finally, it must be remembered that in all obscure cases of hypertrophy and failure of the right heart, the possibility of widened connections between the bronchial and pulmonary arteries should be considered (Wood and Millar, 1937/8).

References
The stethoscope and its use are described. A current classification of breath sounds, voice sounds and adventitious sounds is presented, and its use in diagnosis outlined. Finally, the value of auscultation to the physiotherapist is discussed.

INTRODUCTION

The invention of the stethoscope by Laennec in 1818 was a major advance in medicine; the instrument allowed diagnosis and assessment of cardio-respiratory disorders to be carried out with precision not previously attainable, and to this day the stethoscope remains an indispensable part of the physician’s armamentarium. For physiotherapists, however, the situation is not so clear cut. A substantial proportion of physicians and surgeons view with deepest suspicion the sight of a physiotherapist auscultating a patient’s chest. In the eyes of these critics it verges on blasphemy for the physiotherapist to own a stethoscope, or to carry one about with insight and understanding. This article is written by many members of the physiotherapy profession, believing auscultation to be very much their business, as a matter of routine. The opposite viewpoint is held by many members of the physiotherapy profession, believing auscultation to be very much their business, and a valuable source of information pertinent to the performance of their work. I do not propose to enter into the pros and cons of this controversy. I consider it beyond dispute, however, that if the stethoscope is to be used by physiotherapists, it should be used correctly, with insight and understanding. This article is written to help achieve this end.

THE STETHOSCOPE

(Greek stethos, the chest; skopeein, to explore).

There are three components of the modern stethoscope. They are the chest piece, the tubing and the binaural.

The chest piece may be of the bell type, diaphragm type, or a combination of these two. The two are really variants of the same principle — that of a damped diaphragm system. The area of skin enclosed by the bell behave as a diaphragm, the tautness of which may be varied by the pressure applied. The firmly applied bell therefore subtends an area of taut skin which behaves in a similar way to the diaphragm chest piece; it filters out low frequency sounds, allowing the higher frequencies to come through. It should be remembered, though, that the volume of sound is related to the area of the chest piece, so that the diaphragm chest piece, being larger, will in general produce a higher amplitude of sound than the firmly applied bell. In contrast, the softly applied bell subtends an area of lax skin which will have a much lower resonant frequency, favouring the transmission of low frequency sounds.

Heart sounds, and some cardiac murmurs, are in the lower frequency range (20 - 115 cycles/sec), so that cardiologists will usually prefer the softly applied bell when auscultating the heart.

Lung sounds, however, and especially abnormal lung sounds, are in the higher frequency range (200 - 2000 cycles/sec), so that use of the diaphragm is generally preferable, although the firmly applied bell could be used. Advantages of the diaphragm include the higher amplitude of sound, as already mentioned, and the easier application over an uneven or bony chest cage, where incomplete contact between skin and the rim of the bell would result in complete loss of sound. The firmly applied bell may be useful on occasion; for example, in confirming the presence of fine adventitious sounds which can sometimes be generated artificially by movement of the diaphragm on the skin surface. The bell is also useful in children, for whom the diaphragm may be inappropriately large, although paediatric stethoscopes are available.

The tubing is of considerable importance in the efficiency of a stethoscope. Sound loss can result from the use of incorrect dimensions or materials. The material should be firm, inert, reasonably thick and polished in its internal bore for maximum transmission. Loss of high frequencies can result if the volume of the system is too large or if the diameter of the tubing is too fine. A good compromise is Tygon tubing, as used in the

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