Effects of alveolar dead-space, shunt and $\dot{V}/\dot{Q}$ distribution on respiratory dead-space measurements

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Background. Respiratory dead-space is often increased in lung disease. This study examines the effects of increased alveolar dead-space ($V_{D\text{alv}}$), pulmonary shunt, and abnormal ventilation perfusion ratio ($\dot{V}/\dot{Q}$) distributions on dead-space and alveolar partial pressure of carbon dioxide ($P_{\text{ACO}_2}$) calculated by various methods, assesses a recently published non-invasive method (Koulouris method) for the measurement of Bohr dead-space, and evaluates an equation for calculating physiological dead-space ($V_{D\text{phys}}$) in the presence of pulmonary shunt.

Methods. Pulmonary shunt, $\dot{V}/\dot{Q}$ distribution and $V_{D\text{alv}}$ were varied in a tidally breathing cardiorespiratory model. Respiratory data generated by the model were analysed to calculate dead-spaces by the Fowler, Bohr, Bohr–Enghoff and Koulouris methods. $P_{\text{ACO}_2}$ was calculated by the method of Koulouris.

Results. When $V_{D\text{alv}}$ is increased, $V_{D\text{phys}}$ can be recovered by the Bohr and Bohr–Enghoff equations, but not by the Koulouris method. Shunt increases the calculated Bohr–Enghoff dead-space, but does not affect Fowler, Bohr or Koulouris dead-spaces, or $V_{D\text{phys}}$ estimated by the shunt-corrected equation if pulmonary artery catheterization is available. Bohr–Enghoff but not Koulouris or Fowler dead-space increases with increasing severity of $\dot{V}/\dot{Q}$ maldistribution. When alveolar $PCO_2$ is increased by any mechanism, $P_{\text{ACO}_2}$ calculated by Koulouris’ method does not agree well with average alveolar $PCO_2$.

Conclusions. Our studies show that increased pulmonary shunt causes an apparent increase in $V_{D\text{phys}}$, and that abnormal $\dot{V}/\dot{Q}$ distributions affect the calculated $V_{D\text{phys}}$ and $V_{D\text{alv}}$, but not Fowler dead-space. Dead-space and $P_{\text{ACO}_2}$ calculated by the Koulouris method do not represent true Bohr dead-space and $P_{\text{ACO}_2}$ respectively, but the shunt-corrected equation performs well.

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Keywords: computers; lung, shunting; ventilation, deadspace; ventilation, ventilation–perfusion

Symbols used in the paper

$P_{\text{ACO}_2}$=ideal alveolar carbon dioxide partial pressure; $P_{\text{ACO}_2}$=arterial carbon dioxide partial pressure; $P_{\text{ACO}_2}$=volume-weighted average of $PCO_2$ in all the alveolar compartments in the model; $P_{\text{CO}_2}$=mixed expired carbon dioxide partial pressure; $P_{\text{ECO}_2}$=end-tidal partial pressure of carbon dioxide; $Q_{\text{S}}/Q_{\text{T}}$=pulmonary shunt fraction; $V_{D\text{alv}}$=alveolar dead-space; $V_{D\text{anat}}$=anatomical dead-space in the model; $V_{D\text{BE}}$=Bohr–Enghoff dead-space calculated by substituting $P_{\text{ACO}_2}$ for $P_{\text{ACO}_2}$ in Bohr equation; $V_{D\text{bohr}}$=Bohr dead-space; $V_{D\text{con}}$=Bohr–Enghoff dead-space corrected for shunt; $V_{D\text{ETT}}$=dead-space calculated by substituting $P_{\text{CO}_2}$ for $P_{\text{ACO}_2}$ in Bohr equation; $V_{D\text{fowler}}$=Fowler dead-space; $V_{D\text{K}}$=dead-space calculated by Koulouris method; $V_{D\text{phys}}$=physiological dead-space (total dead-space in the model);

$\dot{V}/\dot{Q}$=ventilation to perfusion ratio; $V_T$=tidal volume; $V_{T\text{alv}}$=alveolar tidal volume ($V_T–V_{D\text{fowler}}$).

Introduction

Respiratory dead-space measurements have been used in determining surfactant efficacy in surfactant-depleted lungs, diagnosing pulmonary embolism, providing useful prognostic information early in the course of acute respiratory distress syndrome, predicting successful extubation in infants and children, and separating patients with asthma from patients with emphysema with the same degree of airways obstruction. All these clinical applications depend on accurate measurements of respiratory dead-space.

Bohr dead-space ($V_{D\text{bohr}}$) is a function of ideal alveolar partial pressure of carbon dioxide ($P_{\text{ACO}_2}$). Because $P_{\text{ACO}_2}$
is difficult to estimate, Enghoff\textsuperscript{8} substituted arterial partial pressure of carbon dioxide ($P_{\text{ACO}_2}$) for $P_{\text{ACO}_2}$, giving rise to the Bohr–Enghoff dead-space (Appendix A), usually referred to as physiological dead-space ($V_{\text{Dphys}}$). Alveolar dead-space\textsuperscript{9} is commonly defined as the difference between $V_{\text{Dphys}}$ and the anatomical dead-space ($V_{\text{D}_{\text{Fowler}}}$), which is estimated by a method proposed by Fowler\textsuperscript{10} (Appendix A). Disadvantages of the Bohr–Enghoff method are that it is invasive and cannot be used breath-by-breath when $P_{\text{ACO}_2}$ is changing rapidly. Recently, Koulouris and colleagues\textsuperscript{11} reported a new non-invasive method to calculate Bohr dead-space and $P_{\text{ACO}_2}$ based on an analysis of the expired carbon dioxide volume vs expired tidal volume curve from a single expiration (Appendix A). This technique is apparently simple and non-invasive, but has not been validated independently.

Shunt reduces the overall efficiency of gas exchange and results in arterial blood gas tensions closer to those of mixed venous blood, thus increasing the measured apparent physiological dead-space by increasing $P_{\text{ACO}_2}$\textsuperscript{12}. A method for correcting dead-space measurements for the effects of shunt has been reported by Kuwabara and Duncalf\textsuperscript{13} (Appendix A) but its validity has not been demonstrated.

Inhomogeneity of ventilation/perfusion ($\dot{V}/\dot{Q}$) ratio increases the measured alveolar dead-space by two mechanisms. Firstly, the venous admixture is increased from lung regions with low $\dot{V}/\dot{Q}$ ratio; secondly, lung units with high $\dot{V}/\dot{Q}$ ratio contribute to physiological dead-space.

Although series dead-space can be altered easily in studies in vivo, it is difficult to control changes in alveolar deadspace and the $\dot{V}/\dot{Q}$ distribution. Thus the effects of changes in $\dot{V}/\dot{Q}$ distribution on measures of respiratory dead-space have not been studied systematically.

The aims of this study were to assess the method of Koulouris and colleagues\textsuperscript{11} for calculating Bohr deadspace and alveolar $PCO_2$, to demonstrate the validity of the correction proposed by Kuwabara and Duncalf\textsuperscript{13} for calculating physiological dead-space in the presence of pulmonary shunt, and to evaluate the effects of varying alveolar dead-space, pulmonary shunt and abnormal $\dot{V}/\dot{Q}$ distributions on $P_{\text{ACO}_2}$ and dead-space calculated by five different methods.\textsuperscript{7,8,10,11,13}

\section*{Methods}

\textbf{The computer model}

We used a comprehensive, mathematical, tidally breathing computer model of the cardiorespiratory system.\textsuperscript{14} The model incorporates a 15-compartment approximation to Weibel’s lung model\textsuperscript{15} and simulates diffusive and convective transport and storage of gases in the lungs. The branching airways comprise 11 anatomical dead-space compartments and terminate in one unperfused and three perfused alveolar compartments (Fig. 1). Carbon dioxide is stored in two lung tissue compartments.\textsuperscript{16} Ventilation and $\dot{V}/\dot{Q}$ heterogeneity can be simulated by varying the inspired gas distribution and the fraction of the cardiac output perfusing the three perfused alveolar compartments. A variable right-to-left shunt is provided. The model simulates alveolar-capillary diffusion; intraventricular and intravascular mixing; variable transport delays; intravascular storage; and carbon dioxide and oxygen storage, production and consumption in eight anatomically appropriate body compartments. Molar quantities of oxygen, nitrogen and carbon dioxide are conserved in the model. Non-linear blood gas dissociation curves\textsuperscript{17} include the Haldane and Bohr effects. The tissue dissociation curves are after Farhi and Rahn\textsuperscript{18} and Cherniack and Longobardo.\textsuperscript{19}

The respiratory flow waveform was selected to match a mechanically ventilated subject: constant inspiratory flow followed by exponential expiratory flow. The model is implemented in Matlab and Simulink (Mathworks, Natick, MA, USA) and has been verified against published human data.\textsuperscript{14} Additional validation is described in Appendix B.

\section*{Part 1. Effect of the ratio of physiological dead-space to tidal volume on respiratory dead-space measurements}

The model was configured to simulate a 70 kg healthy adult male subject with respiratory parameters, as shown in Table 1. The ratio of the tidal volume that reaches alveolar dead-space ($Alv$ DS in Fig. 1) to the alveolar tidal volume was set in turn to 1, 10, 20, 30, 40 and 50%. Alveolar tidal volume ($VTalv$) is the volume of fresh inspired gas that reaches the alveoli. Corresponding ratios of physiological dead-space to tidal volume in the model were 30, 36, 43, 51, 58 and 65%, respectively. The $\dot{V}/\dot{Q}$ ratios in the three ventilated and perfused lung compartments were the same, and decreased as the dead-space increased. Total anatomical dead-space and total alveolar volume were kept constant. The volumes of the alveolar dead-space compartment and the associated anatomical dead-space compartments in the model were increased in proportion to the alveolar dead-space ventilation. The volumes of the perfused alveoli and their associated anatomical dead-space compartments were decreased in proportion to the decreasing ventilation directed to those alveoli. Under each condition, the model was run for 7200 s simulation time to achieve steady-state $PCO_2$ and $PO_2$ in mixed venous blood, and alveolar and body compartments. The data were sampled at 100 Hz and stored for off-line analysis. The last complete respiratory cycle in each 2-h run was analysed. The simulated subject was assumed to be anaesthetized and paralysed and $P_{\text{ACO}_2}$ was allowed to reach unphysiological levels.

\section*{Part 2. Effect of pulmonary shunt on respiratory dead-space measurements}

In this part of the study the ratio of alveolar dead-space to alveolar tidal volume was kept constant at 1%, corresponding to a ratio of physiological dead-space to tidal volume of
30%. The tidal volume in the alveolar dead-space was 5.3 ml. The pulmonary shunt was set in turn to 2, 10, 20, 30, 40 and 50% of the total pulmonary blood flow and the other respiratory parameters were the same as in Part 1 (Table 1).

V/Q ratios of all the perfused compartments were the same, and increased as the shunt increased. Under each condition the model was run for 7200 s of simulation time to reach steady state $P_{CO_2}$ and $P_{O_2}$. Data from the last complete respiratory cycle in each 2-h run were recorded and analysed.

**Part 3. Effect of V/Q ratio heterogeneity on respiratory dead-space measurements**

The ratio of alveolar dead-space to alveolar tidal volume and the pulmonary shunt were set to 1% and 2% respectively. The minute alveolar ventilation of 5.30 litre min$^{-1}$ and the pulmonary perfusion of 5.40 litre min$^{-1}$ were unevenly distributed to the three ventilated and perfused alveolar compartments to create $V/Q$ values of 0.1, 1.0 and 10 to simulate patients with chronic obstructive pulmonary disease.$^{20}$ The percentage perfusion to the middle $V/Q$ compartment ($V/Q=1$) was set in turn to 98, 78, 58, 38 and 18% of pulmonary blood flow to simulate increasing severity of $V/Q$ mismatch, and the ventilation and perfusion of all three ventilated and perfused compartments were calculated by the method described in Appendix C. The model was run for 7200 s simulation time at each setting to reach steady-state $P_{CO_2}$ and $P_{O_2}$ in mixed venous blood and alveolar and body compartments, and the respiratory

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**Table 1** Model parameters

<table>
<thead>
<tr>
<th>Respiratory parameters</th>
<th>Value</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tidal volume</td>
<td>0.75</td>
<td>litre</td>
</tr>
<tr>
<td>Respiratory rate</td>
<td>10</td>
<td>bpm</td>
</tr>
<tr>
<td>I:E ratio</td>
<td>1:2.3</td>
<td>–</td>
</tr>
<tr>
<td>Airway temperature</td>
<td>35</td>
<td>°C</td>
</tr>
<tr>
<td>FRC</td>
<td>2.2</td>
<td>litre</td>
</tr>
<tr>
<td>$P_{CO_2}$</td>
<td>35</td>
<td>%</td>
</tr>
<tr>
<td>Instrument dead-space</td>
<td>50</td>
<td>ml</td>
</tr>
<tr>
<td>Airway dead-space</td>
<td>170</td>
<td>ml</td>
</tr>
<tr>
<td>Pulmonary shunt</td>
<td>2</td>
<td>%</td>
</tr>
<tr>
<td>Cardiac output</td>
<td>5.29</td>
<td>litre min$^{-1}$</td>
</tr>
<tr>
<td>Hb</td>
<td>15</td>
<td>g dl$^{-1}$</td>
</tr>
</tbody>
</table>
parameters were measured and analysed. Log standard deviations of the perfusion distributions were calculated for each condition.

Data analysis

\( P_{\text{aco2}} \), fluctuated during respiration and was therefore averaged over a complete respiratory cycle. \( P_{\text{aco2}} \) is the volume-weighted average of the \( P_{\text{co2}} \) in the three perfused and ventilated alveolar compartments averaged over one respiratory cycle. \( V_{\text{dphys}} \) was calculated by the equal area method (Appendix A). \( V_{\text{dBohr}} \) (Parts 1 and 2 of this study) and \( V_{\text{DBE}} \) (Parts 1, 2 and 3 of this study) were calculated according to Equations 1 and 2 (Appendix A) respectively. Bohr–Enghoff dead-space corrected for the effects of shunt (\( V_{\text{DCorr}} \)) was calculated using Equation 3 (Appendix A). The calculation of Bohr dead-space by the method of Koulouris and colleagues\(^{11} \) (\( V_{\text{DK}} \)) is described in Appendix A. An estimate of the Bohr dead-space was also calculated by substituting end-tidal carbon dioxide partial pressure (\( P_{\text{etco2}} \)) for \( P_{\text{aco2}} \) (Equation 4, Appendix A). We refer to this dead-space as \( V_{\text{DBE}} \). The physiological dead-space of the model (\( V_{\text{dphys}} \)) was calculated as follows:

\[
V_{\text{dphys}} = V_{\text{dAnat}} + (V_T - V_{\text{dAnat}}) F_{\text{Dalv}}
\]

where \( V_{\text{dAnat}} \) is the volume of the anatomical dead-space compartments in the model and \( F_{\text{Dalv}} \) is the fraction of the tidal volume that enters the parallel dead-space compartment.

Results

Part 1. Effect of the ratio of physiological dead-space to tidal volume on respiratory dead-space measurements

The \( V/Q \) ratios in the three ventilated and perfused alveolar compartments and the arterial partial pressures of carbon dioxide and oxygen resulting from increased alveolar dead-space are shown in Table 2. Increased alveolar dead-space affects \( P_{\text{aco2}} \), more than \( P_{\text{atco2}} \). Figure 2a shows simulated and calculated \( P_{\text{co2}} \) as functions of the ratio of physiological dead-space to tidal volume. \( P_{\text{aco2}} \) and \( P_{\text{aco2}} \), predicted by the model, do not differ greatly and increase monotonically with increasing physiological dead-space fraction. \( P_{\text{etco2}} \) and the associated measured values (\( P_{\text{etco2}} \) and \( P_{\text{aco2}} \)) decrease slightly with increasing physiological dead-space. The maximum difference between \( P_{\text{etco2}} \) and \( P_{\text{aco2}} \) is less than 0.25 kPa (2 mmHg) under all conditions studied.

Figure 2b shows calculated dead-space to tidal volume ratios as functions of true physiological dead-space to tidal volume ratio. \( V_{\text{dBohr}} \) and \( V_{\text{DBE}} \) increase with increasing physiological dead-space. Both \( V_{\text{dBohr}} \) and \( V_{\text{DBE}} \) overestimate the model physiological dead-space when alveolar dead-space is small but slightly underestimate physiological dead-space when the model alveolar dead-space is large.

| Table 2 | \( V/Q \) ratios of perfused alveolar compartments and steady state arterial blood gas tensions in parts 1 and 2 of this study. \( V/Q \) is the ventilation to perfusion ratio for the three ventilated and perfused compartments with equal \( V/Q \) |
|---------|-----------------|-----|-----|-----|-----|-----|-----|-----|-----|
| \( V_{\text{dAnat}}/V_T \) | 1% | 10% | 20% | 30% | 40% | 50% |
| \( V/Q \) | 0.95 | 0.87 | 0.77 | 0.67 | 0.58 | 0.48 |
| \( P_{\text{aco2}} \) (kPa) | 4.7 | 5.1 | 5.6 | 6.2 | 7.0 | 8.1 |
| \( P_{\text{atco2}} \) (kPa) | 23.1 | 23.4 | 22.9 | 22.2 | 21.2 | 19.9 |

<table>
<thead>
<tr>
<th>Shunt</th>
<th>2%</th>
<th>10%</th>
<th>20%</th>
<th>30%</th>
<th>40%</th>
<th>50%</th>
</tr>
</thead>
<tbody>
<tr>
<td>( V/Q )</td>
<td>0.95</td>
<td>0.88</td>
<td>0.78</td>
<td>0.68</td>
<td>0.58</td>
<td>0.49</td>
</tr>
<tr>
<td>( P_{\text{aco2}} ) (kPa)</td>
<td>4.7</td>
<td>4.8</td>
<td>4.9</td>
<td>5.0</td>
<td>5.1</td>
<td>5.4</td>
</tr>
<tr>
<td>( P_{\text{atco2}} ) (kPa)</td>
<td>23.9</td>
<td>17.1</td>
<td>13.0</td>
<td>10.2</td>
<td>8.1</td>
<td>6.5</td>
</tr>
</tbody>
</table>
VDK, VDFowler and VDET are approximately independent of alveolar dead-space, shunt and V/Q ratio heterogeneity and lie between approximately 29 and 35% under all conditions studied (Figs 2–4).

**Part 2. Effect of pulmonary shunt on respiratory dead-space measurements**

The ventilation and perfusion to the three ventilated and perfused alveolar compartments and the arterial partial pressure of carbon dioxide and oxygen when the shunt is increased are shown in Table 2. Shunt affects $\text{PaO}_2$ more than $\text{PaCO}_2$. $\text{PaCO}_2$ increases with increasing pulmonary shunt, while $\text{PeCO}_2$, $\text{PAKCO}_2$ and the volume-averaged $\text{PAkCO}_2$ and $\text{PAKxCO}_2$ fall slightly (Fig. 3A). The effect of pulmonary shunt on the partial pressure of carbon dioxide in the alveoli and in the arterial blood is less marked than the effect of directing similar proportions of tidal volume to alveolar dead-space.

The Bohr–Enghoff dead-space increases with increasing pulmonary shunt (Fig. 3B). $\text{VDcorr}$ and $\text{VDBohr}$ are greater than $\text{VDFowler}$ but smaller than $\text{VDK}$ and are approximately independent of shunt.

**Part 3. Effect of $V/Q$ ratio heterogeneity on respiratory dead-space measurements**

The perfusion and ventilation of each compartment, the respiratory variables and the arterial partial pressures of carbon dioxide and oxygen are shown in Table 3. In the $V/Q$ ratio heterogeneity study it was not appropriate to calculate Bohr dead-space due to the variation of $\text{PCO}_2$ among the ventilated and perfused alveoli. $\text{PaCO}_2$ and the $\text{PCO}_2$ in each individual alveolar compartment increase with increasing heterogeneity of the $V/Q$ ratio (Fig. 4A). The $\text{PCO}_2$ of each alveolar compartment is inversely related to the $V/Q$ ratio of the compartment. The $\text{VDcorr}/\text{VT}$ ratio increases from 30.5% at optimal $V/Q$ distribution to 64.9% when 78% of the pulmonary perfusion is distributed to the compartments with $V/Q$ of 0.1 and 10 (Fig. 4B).

**Discussion**

This study found that the Koulouris method for calculating Bohr dead-space overestimates slightly when alveolar
dead-space is small but substantially underestimates Bohr dead-space when alveolar dead-space is increased. VDK appears to be no better than VDET as an estimate of Bohr dead-space under the conditions in this study. The Koulouris method for calculating alveolar P\textsubscript{CO\textsubscript{2}} does not respond to changes in the P\textsubscript{CO\textsubscript{2}} of ventilated and perfused alveoli caused by increasing alveolar dead-space. As respiratory dead-space has been widely studied and used in anaesthesia and in emergency and intensive care medicine, it is important that new methods for dead-space measurement should be validated independently before they are used clinically. Bohr dead-space as calculated by the Koulouris method is not accurate and thus the Koulouris method is not validated by this study.

Pulmonary embolism results in lung units that are poorly perfused but maintain approximately normal ventilation.\textsuperscript{25} Although pulmonary embolism is a complex pathological entity with mixed presentation of shunt and dead-space, the increased pulmonary dead-space in Part 1 of this study approximately simulates the main features of gas exchange in patients with pulmonary embolism. The Bohr–Enghoff dead-space accurately follows the increase in the model alveolar dead-space, while the Fowler dead-space is unaffected (Fig. 2b). Hence, calculated alveolar dead-space, one of the diagnostic markers of pulmonary embolism,\textsuperscript{23} increases. Dead-space calculated by the Koulouris method does not correlate well with alveolar dead-space or Bohr dead-space, suggesting that the Koulouris method cannot contribute to the diagnosis of pulmonary embolism.

Pulmonary shunt increases V\textsubscript{BE} but does not affect V\textsubscript{Fowler} or V\textsubscript{Bohr}. The equation of Kuwabara and Dunn\textsuperscript{13} (Equation 3, Appendix A) calculates model dead-space correctly in the presence of substantial shunt. Use of this correction equation requires pulmonary artery catheterization due to the need for measurement of shunt, mixed venous P\textsubscript{CO\textsubscript{2}} and P\textsubscript{aCO\textsubscript{2}}.

The invalidity of the Koulouris method can be demonstrated theoretically. The Bohr dead-space equation (Equation 1, Appendix A) assumes that expired gas emanates from two compartments: a perfused alveolar compartment and an unperfused dead-space. The Bohr equation makes no assumptions regarding the sequence in which gas from the two compartments is expired. In contrast, the Koulouris method (Fig. A1b in Appendix A) assumes that an expiration comprises two sequential volumes: a dead-space (V\textsubscript{DK}=ia) containing a volume of carbon dioxide [V\textsubscript{CO\textsubscript{2}(d)=ay] is expired first at a mean carbon dioxide concentration of Fd=ya/ia, and the remaining carbon dioxide (ce') is assumed to be expired in a subsequent volume ae of alveolar gas at end-tidal carbon dioxide concentration F\textsubscript{CO\textsubscript{2}}=ce'/ae=yada. In Figure A1b in Appendix A the lines ia and id represent V\textsubscript{DK} and V\textsubscript{DET} respectively. Hence the line da, which represents the difference between V\textsubscript{DK} and V\textsubscript{DET}, can be expressed as V\textsubscript{CO\textsubscript{2}(d)/Fe\textsubscript{CO\textsubscript{2}}. V\textsubscript{DK} and V\textsubscript{DET} are therefore related by:

\[
V\textsubscript{DK}=V\textsubscript{DET}+V\textsubscript{CO\textsubscript{2}(d)}/Fe\textsubscript{CO\textsubscript{2}}
\]

where V\textsubscript{CO\textsubscript{2}(d)} is the volume of carbon dioxide expired in V\textsubscript{DK}. Similarly, it can be shown that P\textsubscript{A\textsubscript{CO\textsubscript{2}}} and P\textsubscript{E\textsubscript{CO\textsubscript{2}}} are related by:

\[
P\textsubscript{A\textsubscript{CO\textsubscript{2}}}/P\textsubscript{E\textsubscript{CO\textsubscript{2}}}=V\textsubscript{CO\textsubscript{2}}/(V\textsubscript{CO\textsubscript{2}}–V\textsubscript{CO\textsubscript{2}(d)})
\]

where V\textsubscript{CO\textsubscript{2}} is the total amount of carbon dioxide expired in one breath. It can be concluded that V\textsubscript{DK} and P\textsubscript{A\textsubscript{CO\textsubscript{2}}} are always larger than V\textsubscript{DET} and P\textsubscript{E\textsubscript{CO\textsubscript{2}}} respectively.

Alveolar P\textsubscript{CO\textsubscript{2}} increases during expiration and reaches a peak shortly after end-expiration. In vivo, P\textsubscript{E\textsubscript{CO\textsubscript{2}}} is dominated by end-expiratory alveolar gas and can be greater than P\textsubscript{A\textsubscript{CO\textsubscript{2}}, particularly during exercise or when tidal volumes are large.\textsuperscript{24–26} Fletcher and colleagues\textsuperscript{27} found zero or negative arterial-to-end-tidal PC\textsubscript{O\textsubscript{2}} differences in 12% of non-pregnant patients during anaesthesia in which large tidal volumes and low respiratory rates were used. In the method of Koulouris and colleagues,\textsuperscript{11} the relationship between V\textsubscript{DK} and V\textsubscript{DET} is fixed and independent of tidal volume, respiratory rate and alveolar dead-space, so end-tidal P\textsubscript{CO\textsubscript{2}} can never exceed P\textsubscript{A\textsubscript{CO\textsubscript{2}}}.

In this study the simulated subject was assumed to be anaesthetized and paralysed and P\textsubscript{A\textsubscript{CO\textsubscript{2}}} was allowed to rise as high as 8.1 and 9.2 kPa when alveolar dead-space and V/Q ratio heterogeneity respectively were increased. In a study of COPD patients with severe V/Q heterogeneity, Conti and colleagues\textsuperscript{28} reported P\textsubscript{A\textsubscript{CO\textsubscript{2}}} values as high as 11.6 kPa in mechanically ventilated patients. Breen and colleagues\textsuperscript{29} studied the effects of large pulmonary embolism on carbon dioxide kinetics and physiological dead-space. They found that after 70 min of occlusion of a large pulmonary artery, the P\textsubscript{A\textsubscript{CO\textsubscript{2}}} increased from 5.5 to 7.3 kPa and P\textsubscript{E\textsubscript{CO\textsubscript{2}}} decreased by 13% of baseline value while

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**Table 3 Ventilation and perfusion parameters of perfused alveolar compartments in Part 3 of this study**

<table>
<thead>
<tr>
<th>Perfusion to high and low V/Q compartments (%)</th>
<th>Pa\textsubscript{a} (kPa)</th>
<th>Pa\textsubscript{co2} (kPa)</th>
<th>Log sd</th>
<th>High: V/Q=10</th>
<th>Middle: V/Q=1.0</th>
<th>Low: V/Q=0.1</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>23.9</td>
<td>4.7</td>
<td>0.012</td>
<td>0.0080</td>
<td>5.27</td>
<td>7.7±7</td>
</tr>
<tr>
<td>20</td>
<td>14.8</td>
<td>5.4</td>
<td>0.42</td>
<td>0.0099</td>
<td>4.21</td>
<td>0.098±9</td>
</tr>
<tr>
<td>40</td>
<td>10.9</td>
<td>6.2</td>
<td>0.54</td>
<td>0.20</td>
<td>3.13</td>
<td>0.20±1.96</td>
</tr>
<tr>
<td>60</td>
<td>8.6</td>
<td>7.4</td>
<td>0.60</td>
<td>0.30</td>
<td>2.05</td>
<td>0.029±2.94</td>
</tr>
<tr>
<td>80</td>
<td>6.6</td>
<td>9.2</td>
<td>0.61</td>
<td>0.39</td>
<td>0.97</td>
<td>0.39±3.92</td>
</tr>
</tbody>
</table>

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physiological dead-space increased from 31 to 52%. Our simulation results are consistent with the in vivo observations of Conti and colleagues\(^{28}\) and Breen and colleagues.\(^{29}\)

We used a mathematical model for this study to facilitate the controlled variation of alveolar dead-space, anatomical dead-space, pulmonary shunt and \(V/Q\) ratio distribution, which are difficult to change prospectively in in vivo studies. A computer model study also avoids the confounding effects associated with biological variations and measurement errors. The main limitations of our computer model include the lumped approximation of the respiratory tree and alveoli, the approximations used to estimate diffusion and convection in the airways and the assumption of equal respiratory time constants and consequent simultaneous emptying of alveolar compartments. The model does not automatically redistribute ventilation or perfusion when these parameters are perturbed and does not simulate hypoxic pulmonary vasoconstriction. These approximations and limitations may affect the shape of the expirogram and hence the calculated dead-space and \(PCO_2\) values. We expect, however, that the limitations of the model affect only the magnitude of the results, not their form or direction.

In conclusion, our simulation results suggest that while the physiological dead-space is estimated well by the Bohr–Enghoff equation when alveolar dead-space and \(V/Q\) ratio distribution vary, respiratory dead-space and alveolar carbon dioxide partial pressure calculated by the Koulouris method do not represent the true Bohr dead-space or alveolar carbon dioxide partial pressure. Increasing pulmonary shunt can cause an apparent increase in \(VD_{phys}\) and abnormal \(V/Q\) distributions affect calculated \(VD_{phys}\) and \(VD_{adv}\), but not Fowler dead-space. The equation suggested by Kuwabara and Duncalf\(^{13}\) for the calculation of dead-space in the presence of shunt performs well, but requires invasive measurements.

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**Appendix A. Dead-space calculation methods and symbols**

**Fowler dead-space**

On an expired carbon dioxide concentration vs expired volume curve (Fig. A1A), a straight line is fitted to the alveolar plateau between 60 and 90% of expired volume by linear regression.\(^{10\,22}\) A vertical line is drawn from the regression line to the \(x\)-axis to divide the rapidly rising part of the carbon dioxide expirogram into two equal areas \((p'\text{ and } q')\). The intersection of the perpendicular line and the \(x\)-axis is the anatomical or Fowler dead-space.\(^{10\,22}\)

**Bohr dead-space**

Bohr dead-space \((VD_{Bohr})\) is the dead-space calculated by the original Bohr equation:\(^{7\,21}\)

\[
VD_{Bohr}=VT\left(1-\frac{PE_{CO_2}}{PA_{CO_2}}\right) \quad (1)
\]

Where \(PE_{CO_2}\) is the mixed expired carbon dioxide partial pressure and \(PA_{CO_2}\) is the ideal alveolar carbon dioxide partial pressure. In Parts 1 and 2 of this study, \(PA_{CO_2}\) is the volume-weighted average of the \(PCO_2\) in the three perfused and ventilated alveolar compartments averaged over one respiratory cycle.

**Bohr–Enghoff dead-space**

Because of the controversy concerning the definition and estimation of \(PA_{CO_2}\),\(^{21}\) Enghoff\(^8\) suggested substituting \(PA_{CO_2}\) for \(PA_{CO_2}\) in the Bohr equation. The dead-space so calculated is termed the Bohr–Enghoff dead-space \((VD_{BE})\):

\[
VD_{BE}=VT\left(1-\frac{PE_{CO_2}}{PA_{CO_2}}\right) \quad (2)
\]

**Shunt correction method**

In the presence of right-to-left shunt, venous blood mixes with pulmonary capillary blood and raises \(PA_{CO_2}\), thus increasing the difference between \(PE_{CO_2}\) and \(PA_{CO_2}\). Hence the Bohr–Enghoff dead-space calculated by Equation 2 is increased. Kuwabara and Duncalf\(^{13}\) applied simple mass balance principles and derived an equation to estimate a corrected physiological dead-space \((VD_{corr})\) in the presence of right-to-left shunt.

\[
VD_{corr}/VT=1-\frac{PE_{CO_2}}{PV_{CO_2}-(PV_{CO_2}-PA_{CO_2})}/\left(1-\frac{Q_s}{Q_T}\right) \quad (3)
\]

where \(Q_s/Q_T\) is the shunt fraction, and \(PV_{CO_2}\) is the mixed venous \(PCO_2\).

**\(VD_{ET}\)**

Dead-space calculated by using \(PE_{CO_2}\) in place of \(PA_{CO_2}\) in the Bohr equation (Equation 1) is termed \(VD_{ET}\) in this paper:

\[
VD_{ET}=VT\left(1-\frac{PE_{CO_2}}{PE'_{CO_2}}\right) \quad (4)
\]

**Koulouris dead-space**

Figure A1B shows expired carbon dioxide volume vs expired volume. Line \(cb\) is drawn such that areas \(p\) and \(q\) are equal. Point \(d\) is chosen such that the slope of line \(cd\) is end-tidal concentration. The volume \(de\) is the volume the expired carbon dioxide would occupy at end-tidal carbon dioxide concentration (total volume of carbon dioxide divided by end-tidal carbon dioxide concentration), hence \(id\) represents \(VD_{ET}\) as calculated by Equation 3 above. Line \(dx\) is perpendicular and intersects \(cb\) at \(x\). Line \(xy\) is parallel to the \(x\)-axis
and intersects \( cd \) at \( y \). Line \( ya \) is perpendicular and intersects the \( x \)-axis at \( a \). According to Koulouris and colleagues, lines \( ee' \) and \( e'c \) represent the quantities of carbon dioxide expired in the dead-space and alveolar gas respectively. The line \( ae \) represents alveolar tidal volume, line \( ia \) represents Bohr dead-space and the slope of line \( ac \) is the alveolar concentration of carbon dioxide.

**Appendix B. Validation of the cardiorespiratory model**

The model was validated in a clinical study and by comparison of predicted \( P_{CO_2} \) with published measurements.

Figure A2A shows the realistic airway, arterial, alveolar and mixed venous \( P_{CO_2} \) changes with time in the tidally breathing model. The model simulated a 70-kg male subject who was mechanically ventilated with a tidal volume of 9 ml kg\(^{-1}\) at a respiratory rate of 10 bpm. Other parameters are the same as shown in Table 1. Both arterial and alveolar \( P_{CO_2} \) fluctuate during the respiratory cycle but arterial \( P_{CO_2} \) lags alveolar \( P_{CO_2} \). Also demonstrated is that alveolar \( P_{CO_2} \) peaks shortly after the end of expiration. The average mixed venous and arterial \( P_{CO_2} \) are in agreement with the literature.\(^{30,31}\)

Figure A2B shows the dynamic change of measured and predicted \( P_{ECO_2} \) following changes in tidal volume and frequency. After ethics committee approval and informed
consent from the patient, a 59-yr-old 77-kg male patient undergoing vascular surgery was studied. The tidal volume and respiratory rate were set to: 950 and 15; 750 and 7; 500 and 6; 850 ml and 10 bpm respectively and in sequence. The $P_{E\,CO_2}$ was recorded for each expiration. The model was set to simulate a 75-kg male subject undergoing mechanical ventilation with the same combinations of tidal volume and respiratory rate. The results show that the model simulation closely represents the dynamic change of $P_{E\,CO_2}$ recorded from the patient (Fig. A2B).

Figure A2C shows the model’s prediction of $P_{a\,CO_2}$ and $P_{E\,CO_2}$ and comparison with a published study. In that study,
12 patients were ventilated with tidal volumes of 10, 7.5, 5 and 2.5 ml kg$^{-1}$ and respiratory rates of 10, 13, 21 and 40 bpm respectively. Each setting was maintained for 10 min and $P_{ACO2}$ and $P_{ECO2}$ were measured and recorded. The model was set to simulate a 75-kg male subject ventilated with the same tidal volumes and respiratory rates as the patients. Other parameters are the same as in Table 1. The anatomical dead-space was adjusted as a function of tidal volume. At each setting, the model was run for 10 min and the $P_{ACO2}$ and $P_{ECO2}$ were recorded. The average absolute error of the model predictions was 0.94% of the mean and 9.1% of the SD for measured $P_{ACO2}$, and 4.4% of the mean and 28.4% of the SD for measured $P_{ECO2}$. This result is comparable with a model study by Hardman and Aitkenhead.

Our results show that the model simulates realistically the gas exchange of a human lung both dynamically and in steady state.

**Appendix C. Calculation of ventilation and perfusion to the three perfused and ventilated compartments in the model**

In Part 3 of this study the ventilation and perfusion of the three ventilated and perfused alveolar compartments, $V_1$, $V_2$, and $V_3$, and $Q_1$, $Q_2$, and $Q_3$, litre min$^{-1}$ respectively, were varied to simulate lungs with various degrees of $V/Q$ inhomogeneity. The $V/Q$ ratios of the ventilated and perfused compartments were fixed as follows:

$$V_1/Q_1 = 0.1$$

$$V_2/Q_2 = 1$$

$$V_3/Q_3 = 10$$

The total ventilation $V$, total perfusion $Q$, shunt blood flow ($Q_s$) and alveolar dead-space ventilation ($V_{D_{alv}}$) were known, therefore:

$$V_1 + V_2 + V_3 + V_{D_{alv}} = \dot{V}$$

$$Q_1 + Q_2 + Q_3 + Q_s = \dot{Q}$$

There were six unknown parameters in five equations (Equations 5–9). We assigned values to $Q_3$ and calculated the remaining five unknown parameters ($V_1$, $V_2$, $V_3$, $Q_1$, and $Q_2$) by solving Equations 5–9 simultaneously. We used software written in Matlab (Mathworks, Natick, MA, USA) to solve the equations and hence determine the fractions of ventilation and perfusion directed to each alveolar compartment.

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