JCI The Journal of Clinical Investigation

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J Clin Invest. 1967;46(3):463-474. https://doi.org/10.1172/JCI105548.

Research Article

A method for appraising the distribution of diffusing capacity of the lungs (\mathbb{Q}) in relationship to pulmonary capillary blood flow ([unk]Q_C) in normal human subjects was derived from measurements of oxygen diffusing capacity (\mathbb{P}_{O_2}) and carbon monoxide diffusing capacity ($\mathbb{D}_{L_{CO}}$) performed during breath holding. This method utilizes the fact that the observed \mathbb{P}_{O_2} is considerably reduced in value if uneven distribution of \mathbb{D}_L with respect to [unk]Q_C (uneven \mathbb{D}_L /[unk]Q_C) is present. In contrast, $\mathbb{D}_{L_{CO}}$ is barely affected by uneven \mathbb{D}_L /[unk]Q_C, and from its measured value one can calculate the value \mathbb{Q}_{O_2} would have if no uneven \mathbb{D}_L /[unk]Q_C were present (true $\mathbb{D}_{L_{O_2}}$). Once observed $\mathbb{D}_{L_{O_2}}$ and true $\mathbb{D}_{L_{O_2}}$ are known, the degree of uneven \mathbb{D}_L /[unk]Q_C in the lung can be calculated.

In five normal, resting, sitting subjects average values for true P_{D_2} were 57 ml per (minute × mm Hg), and the directly measured $D_{L_{D_2}}$ was 33 ml per (minute × mm Hg). These values could be explained if one-half of total [unk]@ were distributed to approximately 15% of total D_L .

These measurements did not permit the determination of the alveolar to end capillary Q gradient, but calculations demonstrate that an important factor in determining its size may be the pattern of uneven $D_L/[unk]Q_C$ present in the lungs. Estimations of the alveolar-end capillary O_2 gradient from measurements of $D_{L_{CO}}$ or $D_{L_{O_2}}[...]$



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Determination of Distribution of Diffusing Capacity in Relation to Blood Flow in the Human Lung *

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Summary. A method for appraising the distribution of diffusing capacity of the lungs (DL) in relationship to pulmonary capillary blood flow ($\dot{Q}c$) in normal human subjects was derived from measurements of oxygen diffusing capacity (DL_{O_2}) and carbon monoxide diffusing capacity (DL_{CO}) performed during breath holding. This method utilizes the fact that the observed DL_{O_2} is considerably reduced in value if uneven distribution of DL with respect to $\dot{Q}c$ (uneven $DL/\dot{Q}c$) is present. In contrast, DL_{CO} is barely affected by uneven $DL/\dot{Q}c$, and from its measured value one can calculate the value DL_{O_2} would have if no uneven $DL/\dot{Q}c$ were present (true DL_{O_2}). Once observed DL_{O_2} and true DL_{O_2} are known, the degree of uneven $DL/\dot{Q}c$ in the lung can be calculated.

In five normal, resting, sitting subjects average values for true DL_{O2} were 57 ml per (minute \times mm Hg), and the directly measured DL_{O2} was 33 ml per (minute \times mm Hg). These values could be explained if one-half of total \dot{Qc} were distributed to approximately 15% of total DL.

These measurements did not permit the determination of the alveolar to end capillary O₂ gradient, but calculations demonstrate that an important factor in determining its size may be the pattern of uneven D_L/\dot{Q}_C present in the lungs. Estimations of the alveolar-end capillary O₂ gradient from measurements of $D_{L_{CO}}$ or $D_{L_{O_2}}$ that do not take into account uneven D_L/\dot{Q}_C may underestimate its size.

Introduction

Although the distribution of ventilation with respect to perfusion in the lungs has been extensively investigated, little attention has been given to the distribution of diffusing capacity in

* Submitted for publication June 29, 1966; accepted December 1, 1966.

This work was supported in part by U. S. Public Health Service grant HE 10324-01 and by a grant from the Life Insurance Medical Research Fund.

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Present address: Institute of Diseases of the Chest, Brompton, London, S.W.3, England. relation to blood flow (1–4). Recently a breathholding method for the determination of DL_{O_2} has been developed that permits the simultaneous measurement of DL_{O_2} and DL_{CO} (5). The numerical value of DL_{O_2} was found on a theoretical basis to be quite sensitive to uneven distribution of diffusing capacity with respect to blood flow (uneven $DL/\dot{Q}c$), whereas DL_{CO} was relatively insensitive. By comparing the numerical values of DL_{O_2} and DL_{CO} , we can evaluate the degree of uneven $DL/\dot{Q}c$ in the lungs. The present work uses this approach to study the diffusion-blood flow relationships in the lungs of five normal resting subjects.

Methods

Theory

Both steady state and breath-holding methods for measuring O_2 diffusing capacity of the lungs require the assumptions of even distribution of diffusing capacity (DL),



FIG. 1. DIAGRAMS OF THE PULMONARY CAPILLARY BED. Diagram A illustrates the model of the lung and its pulmonary capillary bed that must be assumed for the determination of oxygen diffusing capacity (DLo,) by either the breath-holding method with ³⁴O₂ or by the steady state method. Diffusing capacity, capillary blood flow, Qc, lung volume, and ventilation are assumed to be distributed evenly. This assumption makes the capillary transit time (TL) from the pulmonary arterioles (labeled P.A.) to the pulmonary veins (labeled P.V.) constant and the end capillary oxygen pressure (Po₂ or P³⁴O₂) constant throughout the lung. Diagram B, adapted from Von Hayek (9), represents a more anatomically correct view of the pulmonary capillary bed, and since it is likely that some of the blood travels from P.A. to P.V. at different speeds, uneven distribution of diffusing capacity to capillary blood flow (uneven DL/Oc) is a distinct possibility. Diagram C is the lung model used in this paper. Each alveolus is considered to be perfused by two capillaries that may have different lengths and receive different amounts of Qc. Unlike diagram A, in this model uneven DL/Qc may be present depending on the values chosen for the length of the two capillaries and their respective Qc.

pulmonary capillary blood flow ($\dot{Q}c$), and pulmonary capillary blood volume (Vc) along the capillaries. These assumptions are made in order to have a constant alveolarend capillary gradient for O₂ (6) or O₂ isotope throughout the lungs (5). A number of anatomical observations *in vivo* and *in vitro* indicate that the pulmonary capillary bed is in fact a meshwork of capillaries along the wall of each alveolus (7, 8). It is therefore likely that as blood moves from the pulmonary arterioles to the pulmonary veins, some capillary pathways will be longer than others. In comparison to the shorter ones, these long capillaries have a greater surface area and thereby a larger DL. If blood flow is not distributed to the different capillary pathways in proportion to their respective lengths (7), uneven distribution of DL with respect to $\dot{Q}c$ will result (see Figure 1). Since each alveolus has many capillary pathways of varying length, there is the possibility of a large number of different $DL/\dot{Q}c$ compartments within each gas exchange unit.

An analysis that takes into consideration many DL/Qc compartments, although probably more representative of the lungs in vivo, makes calculations extremely tedious. We, therefore, assumed a two compartment system representing two capillaries of different length coursing along the alveolar wall (see Figure 1C). In addition, we assumed that all the capillaries have the same cross-sectional area and capillary wall thickness (even distribution of diffusing capacity with respect to pulmonary capillary blood volume). The DL/Qc ratio of different capillary pathways can then be expressed in terms of capillary transit time (T_L) .¹ For instance, pathways with a low DL/Qc ratio will have a relatively short TL, and capillaries with a high DL/Qc ratio will have a relatively long TL. It should be noted that this model places uneven DL/Qc within each gas exchange unit rather than in gross regions of the lungs such as lobes or segments.

Effect of uneven $DL/\dot{Q}c$ on DL_{O_2} . The measurement of DLO2 with 34O2 requires breath holding after the inspiration of a gas mixture enriched with the stable O₂ isotope of mass 34 (${}^{34}O_2$). The partial pressure of ${}^{34}O_2$ (P ${}^{34}O_2$) is very low in the blood at the start of the capillary and then progressively increases until at the end of the capillary its mean value determined experimentally is approximately 70% of the alveolar $P^{34}O_2$ (5). However, if uneven DL/Qc is present, in those capillaries with a long TL (or high $DL/\dot{Q}c$), the alveolar P³⁴O₂ and capillary P³⁴O₂ will have sufficient time to come almost into equilibrium before reaching the end of the capillary. Once equilibrium is approached, minimal ³⁴O₂ uptake takes place along that portion of the capillary. Because ³⁴O₂ uptake is slight at the end of those capillaries with a long TL, the ³⁴O₂ uptake for any given alveolar P³⁴O₂ is reduced. This reduction results in a decrease in calculated DL_{02} . Therefore if uneven DL/Qc is present, the observed diffusing capacity will be lower than if $DL/\dot{Q}c$ is evenly distributed (see Figure 2).

Effect of uneven $DL/\dot{Q}c$ on DL_{CO} . In the determination of DL_{CO} alveolar CO is so low (approximately 1.5 mm Hg) that only a small amount of CO diffuses into the blood in the pulmonary capillaries. Because the CO entering the capillary blood is such a small fraction of its total capacity for CO, unlike the case for ${}^{34}O_2$ described above, there is little change in capillary Pco during breath holding. Even if in some of the capillaries TL is considerably longer than the mean TL, there is only a slight increase in capillary

¹ The assumption of even distribution of diffusing capacity with respect to Vc is not essential, but seems reasonable on the basis of both anatomical and physiological considerations (8, 10). If this assumption is granted, Vc is proportional to DL, and the TL for any capillary or group of capillaries can be calculated from the following relationship: TL = Vc/Qc. This equation permits the analysis of uneven DL/Qc in terms of TL, which is synonymous with the terms alveolar-capillary contact time (11) or time along the capillary (12) used by others.

Pco as blood moves along the pulmonary capillaries. Because there is little increase in capillary Pco, total CO uptake and calculated DL_{CO} are relatively insensitive to uneven $DL/\dot{Q}c$ compared to DL_{O2} . For instance, if in 80% of the capillary bed the transit time increased from a normal value of about 0.8 second to 5 seconds, the end capillary Pco would only rise to about 0.14 mm Hg, which is less than 10% of the alveolar Pco.² Such an increase in end



FIG. 2. Change in measured DL_{O_2} (observed DL_{O_2}) PRODUCED BY DIFFERENT AMOUNTS OF QC GOING THROUGH 25% OF THE LUNG'S DIFFUSING CAPACITY. A hypothetical lung is considered to have two compartments containing 25% and 75% of the total diffusing capacity. Total or true DL_{0_2} is given a value of 57 ml per (minute \times mm Hg), Qc a value of 7,200 ml per minute, and pulmonary capillary blood volume (Vc) a value of 100 ml. Observed DL₀₂, the value of DLO2 that would be determined by using the breath-holding isotope method (5), according to the mathematical methods derived in the text will only equal true DLO2 when the ratio of diffusing capacity to Qc of each compartment is 57 to 7,200 (even DL/Qc and indicated by 1 arbitrary unit on the upper horizontal axis). If DL/Qc is uneven (25% of the diffusing capacity receives either more or less than 25% of total $\dot{Q}c$), observed DL_{O_2} will be less than its maximal value of 57 ml per (minute \times mm Hg). Note that the greater the abnormality in DL/Qc, the greater is the fall in observed DLO2. Capillary transit time for the small compartment was calculated by the method described in the first footnote.



FIG. 3. GRAPHICAL SOLUTION OF EQUATION 4 IN TEXT. This chart permits the determination of the dimensions of DL and Qc in a lung with two compartments that might have different values for DL and Qc. Numerical constants used are representative values for our subjects, namely true $DL_{02} = 57$ ml per (minute \times mm Hg), total Qc = 7,200 ml per minute, the Bunsen solubility coefficient for total O_2 in blood $(\alpha b_{O_2}) = 2.5$, and Vc = 100 ml. The diagonal solid slightly curved lines are isopleths of observed DL_{O_2} as a percentage of true DL_{O_2} . The broken lines are isopleths of TL in seconds. As an example, assume that one compartment contains 20% of DL and observed DL_{O_2} is 60% of true DL_{O_2} . One then reads across from the 20% point on the upright axis to the diagonal solid line labeled 60. At this point the corresponding point on the horizontal axis is 56%, which is the per cent of the total Qc passing through the compartment. Since the interrupted line labeled 0.3 passes through the same point, TL for this compartment is 0.3 second. The other compartment must then contain 80% of DL and 44% of Qc, and its TL determined from the chart would be 1.5 seconds. At the present time we have no method of determining which of the solutions lying along the isopleths of observed DL_{O_2} as a percentage of true DL_{O_2} represents the correct answer (or answers), and since the lung is most likely composed of a large number of DL-Qc compartments, a two compartment model can only be a rough approximation of the situation present in vivo.

capillary Pco would result in a fall of DL_{CO} of less than 5%.³ However, with the same increase in TL, DL_{02} would

³ Even though the end capillary Pco is 0.14 mm Hg in the capillaries with a TL of 5 seconds, from multiple deter-

² The end capillary Pco of 0.14 mm Hg was obtained by the following calculation: In a subject with a DL_{CO} of 35 ml per (minute \times mm Hg), at an average alveolar Pco of 1.5 mm Hg the amount of CO entering 80% of the capillary bed in 5 seconds would be 0.8 \times 35 \times 1.5 \times (5/60) or 3.5 ml. If the Vc were 100 ml and the hemoglobin concentration 15g per 100 ml, then the total CO capacity of the blood in 80% of Vc would be 0.8 \times 100 \times 1.34 \times (15/ 100) or 16 ml. The per cent carboxyhemoglobin saturation at the end of these capillaries would be (3.5/16) \times 100

or 22%. From the Haldane relationship (13) with a value of 240 for the relative affinity of hemoglobin for CO as compared with O_2 (M) and 120 mm Hg for intracapillary Po_2 , end capillary $Pco = (120 \times 22)/[(240)(99 - 22)]$ or 0.14 mm Hg.

fall to less than one-third of the value it would have if $DL/\dot{Q}c$ were evenly distributed.⁴

Determination of value of DL_{02} if DL and $\dot{Q}c$ are evenly distributed (true DLO2). Since uneven DL/Qc may decrease the numerical value of DLO2 measured with 34O2, DLO2 measured by this technique was called the "observed DLOs." The degree of unevenness of DL/Qc was calculated by determining the difference between the observed DL_{O_2} and the true DL_{O_2} . By "true DL_{O_2} " we mean the value of DLO2 that would be present if diffusing capacity and Qc were evenly distributed. Since DLco is barely affected by uneven DL/Qc, true DL_{02} was determined indirectly from measurements of the pulmonary capillary blood volume (Vc) and diffusing capacity of the alveolarcapillary membrane for CO (DM_{CO}) calculated by measuring DLCO at alveolar O2 tensions of approximately 150 mm Hg and 600 mm Hg (14). The following formula was then used to calculate true DL_{0_2} :

$$\frac{1}{\text{true } DL_{O_2}} = \frac{1}{DM_{34O_2}} + \frac{1}{(\text{Vc})(\theta O_2)}$$
$$= \frac{1}{(1.19)(DM_{CO})} + \frac{1}{(\text{Vc})(\theta O_2)}.$$
 [1]

The two right-hand terms of the above equation equal, respectively, the resistance to diffusion from the alveolus through the alveolar-capillary membrane into the plasma in the capillaries $(1/DM_{340_2})$ and the resistance to diffusion from the plasma into the interior of the red blood cells $[1/(Vc \times \theta O_2)]$. Their sum, which is the left-hand term $(1/\text{true } DL_{O_2})$, equals the total resistance to diffusion. θO_2 is the diffusing capacity of the red blood cell for O_2

expressed in milliliters per (minute \times millimeters Hg \times milliliters), and the values reported by Staub, Bishop, and Forster were used (15). DM₀₂, the diffusing capacity of the alveolar-capillary membrane for ³⁴O₂, expressed in milliliters per (minute \times millimeters Hg), was considered to be equal to DM_{CO} multiplied by 1.19 on the basis of Graham's law of diffusivity of gases.⁵ The derivation of an expression similar to Equation 1 is described by Roughton and Forster (14).

Quantitative analysis of uneven $DL/\dot{Q}c$. Observed DL_{02} is calculated by the following formula, whose derivation is given in detail elsewhere (5):

observed
$$DL_{0_2} = \frac{\dot{Q}c(\alpha b_{0_2})}{760} \ln\left(\frac{1}{K}\right),$$
 [2]

where Qc is the pulmonary capillary blood flow in milliliters per minute measured by the acetylene breath-holding method (16), and αb_{0_2} is the Bunsen solubility coefficient for total O₂ in the blood in milliliters per milliliter per standard atmosphere calculated from the capillary Po₂ present during breath holding and the O₂ capacity of the subject's blood. K is a constant and is defined in the following manner:

$$K = \frac{\text{alveolar } P^{34}O_2 - \text{end capillary } P^{34}O_2}{\text{alveolar } P^{34}O_2 - \text{mixed venous } P^{34}O_2}.$$
 [3]

K is determined from the rate of disappearance of ${}^{34}O_2$ during breath holding and has a numerical value of about 0.3 in normal subjects. If the lung is divided into two compartments with different DL/Qc ratios such as illustrated in Figure 1C, the relationship between observed DL₀₂ and true DL₀₂ will be the following:

observed
$$DL_{0_2} = \frac{\dot{Q}c(\alpha b_{0_2})}{760} \ln \left[\frac{\dot{Q}c}{(\dot{Q}c_1)e^{[-DL_1(760)/\dot{Q}c_1(\alpha b_{0_2})]} + (\dot{Q}c_2)e^{[-DL_2(760)/\dot{Q}c_1(\alpha b_{0_2})]}} \right].$$
 [4]

 \dot{Q}_{c_1} and \dot{Q}_{c_2} are the respective blood flows through the two compartments, and their sum equals \dot{Q}_c . DL₁ and DL₂ are the respective diffusing capacities of the two compartments, and their sum equals true DL₀₂. The detailed

minations of the capillary Pco over the 5-second interval, it is apparent that the mean capillary Pco would be less than half this value. Since CO uptake and DL_{CO} are determined by the alveolar-mean capillary CO gradient, the reduction in calculated DL_{CO} in this part of the capillary bed would be less than 5%. In addition, in the 20% of the capillary bed with a short TL, the mean capillary Pco would be negligible, so that calculated DL_{CO} in these capillaries would be unaffected by uneven $DL/\dot{Q}c$.

⁴ See Figure 3 for dimensions of DL_{02} , Vc, and Qc used. Since 80% of Vc had a TL of 5 seconds, Qc of this compartment could be calculated from the relationship Qc = Vc/ TL and was found to be 14% of total Qc. Since it is assumed that Vc and DL are evenly distributed, a compartment with 80% of total Vc contains 80% of total DL₀₂. From Figure 3 it is apparent that if one compartment has 80% of total DL₀₂ and 14% of Qc, the observed DL₀₂ is 30% of the true DL₀₂ (*vide infra*). derivation of Equation 4 is given in the Appendix. Figure 3 is a graphical solution of Equation 4 that permits the determination of the dimensions of the two diffusing capacity-blood flow compartments for any given value of observed DL_{0_2} expressed as a percentage of true DL_{0_2} . Numerical constants used are given in the legend to the Figure.

Procedures

Equation 4 was applied in the following manner: Observed DL_{0_2} and $\dot{Q}c$ were measured from the rate of disappearance of ${}^{34}O_2$ and acetylene (C_2H_2) during breath holding (5, 16). The sum of DL_1 plus DL_2 , which by definition equals true DL_{0_2} , was determined by the following steps: First the DL_{CO} was measured at alveolar PO_2 of approximately 150 and 600 mm Hg, and then the values for Vc

⁵ According to Graham's law, the diffusivity of two gases through a liquid is proportional to the ratio of their solubilities and inversely proportional to the ratio of the square root of their molecular weights. If the alveolarcapillary membrane is considered to be essentially water, $DM_{34O_9} = DM_{CO}$ (0.244/0.0185)(28/34) = (1.19)(DM_{CO}).

and DM_{CO} were calculated (14). These values were then substituted into Equation 1 in order to obtain the value of true DLo₂ for each subject. The term DL₂ in Equation 4 can then be eliminated by replacing it with true DL_{0_2} -DL₁. Likewise since $\dot{Q}c_1 + \dot{Q}c_2 = \dot{Q}c$, $\dot{Q}c_2$ can be replaced by the term $\dot{Q}c - \dot{Q}c_1$. Because two unknown terms remain in Equation 4, namely DL_1 and Qc_1 , these values can be expressed as a locus of possible solutions on a graph. For example, Figure 3 is the graphical solution for Equation 4 in a hypothetical subject with a true DLO2 of 57 ml per (minute \times mm Hg), Qc of 7,200 ml per minute, and αb_{O_2} of 2.5 ml per ml per standard atmosphere. Note that any point on one of the unbroken curved lines of the Figure represents a possible pair of values for DL_1 and Qc_1 for a particular observed DLO2 expressed as a percentage of true DLO2. Because Equation 4 must be solved by a method of trial and error or with a computer, its graphical solution presented in Figure 3 is helpful in estimating the values of DL_1 and $\dot{Q}c_1$ that would satisfy the values of observed DL_{0_2} and true DLO2 obtained in our subjects. With the assistance of this Figure, Equation 4 was solved for each subject, and the solutions were then plotted in Figure 4.

Subjects

Measurements for the determination of uneven $DL/\dot{Q}c$ were made in five male laboratory personnel, skilled at respiratory maneuvers, in the sitting position. Values for observed DL_{02} previously reported were used (5). At the time of the measurements no symptoms of respiratory disease were present. Chest X rays were read as showing no active disease, and vital capacities as well as maximal midexpiratory flow rates were within the normal limits recently published by Bates and Christie (17). Physical characteristics of the subjects are listed in Table I.



FIG. 4. EXPERIMENTALLY DETERMINED DISTRIBUTION OF DL_{02} WITH RESPECT TO QC. The curved lines represent the patterns of uneven DL/Qc that would explain the experimental findings for each subject. The two older subjects (JC and REF) have the more uneven DL/Qc.

	TABI	ĿΕ	I	
Physical	characteristics	of	experimental	subjects

Subject	ubject Age		Weight	Surface area	
	years	inches	pounds	m^2	
RR	24	72	175	2.01	
GGP	28	73	175	2.03	
ŔŴĦ	34	71	160	1.94	
REF	44	74	170	2.02	
JC	46	64	132	1.65	
•					

Results

The data for determining true DL_{0_2} and observed DL_{0_2} are listed in Table II. For the five subjects the average value for DL_{CO} was 44 ml per (minute \times mm Hg), and observed DL_{0_2} measured simultaneously was 33 ml per (minute \times mm Hg). The value for Vc calculated from measurements of DL_{CO} at different alveolar Po_2 was 104 ml, and DM_{CO} was 61 ml per (minute \times mm Hg). From these figures the value for true DL_{0_2} for the five subjects was calculated. The average was 57 ml per (minute \times mm Hg), which is considerably greater than the observed DL_{0_2} of 33 ml per (minute \times mm Hg).

The pattern of uneven DL/Qc for each subject calculated from Equation 4 is plotted in Figure 4. Unfortunately at the present time we have no method of determining which one of the solutions shown in this Figure represents the correct two compartment model, but it is evident that any of the possible solutions shown represents a fairly marked degree of uneven DL/Qc. For instance, if the lung is divided into two compartments with equal amounts of DL, the amount of blood flowing through the compartment with the lesser amount of Qc varied from a low value of 8% of total Qc in subject JC to a high value of 28% in subject RR. The median value, represented by subject GGP, was 13%. (If no uneven DL/Oc were present, Qc through each compartment would be 50% of total Qc.)

Uneven DL/ $\dot{Q}c$ showed a tendency to be more marked with increasing age, for in the two subjects over 40 years old 50% of DL received on the average 9% of the total $\dot{Q}c$, whereas in the three subjects under 40, 50% of DL received 20% of total $\dot{Q}c$. Measurements in more subjects over a wider age range would be needed to confirm this suggestion of progressively severe uneven DL/ $\dot{Q}c$ with advancing age.

Subject	Qc	Vc	Dlcot	Dmco‡	Dм ₃₄₀₂ §	Alveolar Po2	αb₀₂¶	θο₂¶	True DL02	Observed DL02	Observed DLO2 as per cent of true DLO2
	L/min	ml	ml	ml	ml	mm Hg	ml/ml	ml	ml	ml	%
			min ×mm Hg	min ×mm Hg	min ×mm Hg			min ×mm Hg ×ml	min ×mm Hg	min ×mm Hg	
RR	8.2	98	44	59	70	36.0	2.49	2.8	56	46	82
GGP	8.0	106	51	72	86	41.5	2.71	2.8 '	62	34	55
RWH	7.1	112	44	57	68	42.0	2.62	2.7	55	35	64
REF	6.2	80	40	53	63	49.0	2.60	2.5	48	23	48
JC	6.3	123	43	66	79	43.5	2.46	2.7	64	26	41
Average	72	104	44	61	73	42.0	2.58	27	57	33	58

TABLE II									
Data for determination	of uneven	DL/Qc in	ı five subjects*						

* $\dot{Q}c$ = pulmonary capillary blood flow; Vc = pulmonary capillary blood volume; DL_{C0} and DL_{Q2} = carbon monoxide and oxygen diffusing capacity of the lungs; DM_{CO} and DM_{HO_2} = carbon monoxide and oxygen diffusing capacity of the pulmonary membrane; PO_2 = oxygen tension; αb_{O_2} = Bunsen solubility coefficient for total O_2 in blood; θo_2 = diffusing capacity of the pulmonary membrane PO_2 = oxygen tension; αb_{O_2} = Bunsen solubility coefficient for total O_2 in blood; θo_2 = diffusing capacity of the pulmonary membrane PO_2 = oxygen tension; αb_{O_2} = Bunsen solubility coefficient for total O_2 in blood; θo_2 = diffusing capacity of the pulmonary membrane PO_2 = oxygen tension; αb_{O_2} = Bunsen solubility coefficient for total O_2 in blood; θo_2 = diffusing capacity of the pulmonary membrane PO_2 = oxygen tension; αb_{O_2} = Bunsen solubility coefficient for total O_2 in blood; θo_2 = diffusing capacity of the pulmonary membrane PO_2 = d ing capacity of red blood cells for oxygen.

DLco measured at alveolar Po2 listed in the sixth column of figures.

DMco calculated from measurements of DLco performed at alveolar Po2 of approximately 15 mm Hg and 600 mm Hg

 $\frac{5}{5}$ D_{MM02} calculated by multiplying DMc0 by 1.19 on basis of Graham's law (see text).

Alveolar Po₂ present during the determination of observed DL_{0_2} . ¶ αb_{0_2} and θo_2 for alveolar Po₂ present during determination of observed DL_{0_2} .

Discussion

The above results indicate that a considerable degree of uneven distribution of diffusing capacity (DL) with respect to pulmonary capillary blood flow (Qc) is present in normal resting human subjects. Among the alternative explanations for the data is the possibility that the DL/Qc patterns determined are an artifact due to inaccuracies in the measured quantities of either observed DLo, or true DLo,.

Accuracy of observed DLO2. Sources of error in the measurement of observed DL_{0_2} have been described in detail elsewhere (5). It was concluded that the error in the measurement is less than $\pm 15\%$, which is not sufficient to alter dramatically the degree of uneven DL/Qc calculated for our subjects.

Accuracy of true DL_{0_2} . Appraisal of the accuracy of true DL_{0_2} is more complex because its value depends on the terms from which it is calculated in Equation 1, namely Vc, DM_{CO} , θO_2 , and the factor 1.19 used to convert DM_{CO} into $D_{M_{340_{2}}}$. Errors in the determination of Vc and $D_{M_{CO}}$ by performing $D_{L_{CO}}$ at different alveolar Po₂ have been discussed previously (14, 18). It should be noted that for the subjects in this report errors in Vc and θ_{O_2} produce only onethird the error in true DL₀₂. Therefore it seems unlikely that those measurements would be sufficiently inaccurate to alter markedly true DL₀₂. Errors in DM_{CO} , however, produce almost the same per cent change in true DLO2, but fortunately a lower limit for the value of DM_{CO} can be calculated in the following manner: DLco was measured simultaneously with observed DL_{0_2} . The average value of DL_{CO} for the five subjects was 44 ml per (minute \times mm Hg), which theoretically represents the minimal value possible for DM_{CO} under the conditions of the experiment. Even if DM_{CO} were this minimal value, the average value for true DLo2 calculated by Equation 1 would be 43 ml per (minute X mm Hg), which is still considerably greater than 33 ml per (minute \times mm Hg), the average value of observed DL_{O_2} . The factor 1.19, used to convert DM_{CO} into DM₃₄₀, was calculated on the basis of Graham's law and the assumption that the solubilities of CO and ³⁴O₂ in the alveolar-capillary membrane are similar to their respective solubilities in water. Though the alveolar-capillary membrane is predominantly water, it contains proteins and lipid membranes that might alter the solubilities of these two gases. Experimentally it has been shown that the relative solubility of O₂ of mass 32 compared to CO is changed from 1.25 to 0.78 if the measurement is made in a solu-

tion containing equal quantities of glycerol and buffer solution instead of buffer alone (19). If in these experiments the conversion factor used should have been 0.78 instead of 1.19, the average value of true DLO2 would fall from 57 ml per (minute \times mm Hg) to 40 ml per (minute \times mm Hg). This value is still greater than 33 ml per (minute \times mm Hg), the average value of observed DL₀, so that even if there is an appreciable error in the factor used to convert DM_{CO} into $D_{M_{0_2}}$, there will still be a considerable degree of uneven DL/Qc in the lungs.

In addition to errors in observed DL_{0_2} and true DLO2 several other factors might influence the determination of uneven DL/Qc, namely the artificiality of constructing a two compartment lung, pulsatile pulmonary capillary blood flow, and the upright posture of the subjects.

Errors arising from the assumption of a two compartment lung. Equation 4 is listed in a form that is applicable to a lung with only two DL/Qc compartments, but by adding to the denominator inside the large bracket terms of the form, $\dot{Q}c_n e^{[-DL_n(760)]/[\dot{Q}c_n(\alpha bO_2)]}$, the equation can be

used to construct a lung with many compartments. Although at the present time we have no measurements available that justify the complexity of constructing more than two compartments, it seems likely that the lung may have many capillary pathways with different patterns of DL/Qc distribution. Although a knowledge of the dimensions of a greater number of compartments would be expected to give a more exact picture, a multicompartment model must still take into account the experimental finding of a large difference between true DLo2 and observed DLO2. A multicompartment model of the lungs would therefore not invalidate the existence of uneven DL/Qc. It would, however, demonstrate the pattern of distribution of uneven DL/Qc more clearly.

Effect of pulsatile Qc on uneven DL/Qc. Measurements in man of instantaneous Qc indicate that Qc is pulsatile (20). If some of the red blood cells rapidly traversed the capillaries during systole, whereas others had a prolonged exposure to the alveolar gas during diastole, variation in DL/Qc during the cardiac cycle would

TABLE III

Effect of uneven distribution of alveolar volume (VA), pulmonary capillary blood flow (Qc), and pulmonary diffusing capacity on measured values of CO diffusing capacity (DL_{CO}), O₂ diffusing capacity (DL_{O2}), and the alveolar to end capillary O₂ gradient (A-c O₂ gradient)*

		Com	Compartment dimensions								
		part- ment no.	True DL ₀₂	Qc	VA	Capillary transit time	A-c O2 gradient†	Total A-c O2 gradient‡	Dlco	Observed DL02	
			ml (min× mm Hg)	ml/min	ml	sec	mm Hg	mm Hg	ml (min× mm Hg)	$\frac{ml}{(min \times mm Hg)}$	
А.	Upright posture	1	9.7	980	1,800	1.04	0				
	(rer. 1, 21)	2	18.8	2,380	1,800	0.83	0	0	(-7%)	50	
		3	28.5	3,840	1,800	0.78	0			(-12%)	
в.	Anatomical and	1	10.0	3,600	T	0.29	0.5				
	(ref. 7, 9, 22)	2	30.0	3,600	Ţ	0.88	0	0.25	44 (0%)§	27.2	
		3	17.0	0	T		0			(-52%)	
c.	Pulmonary capil- lary transit time of 0.1 sec in 5% of capillary channels (ref. 23)	1	1.46	1,530	T	0.10	16.5				
		2	55.54	5,670	¶	1.03	0	5	44 (0%)§	38.1 (−33%)∥	
D.	Measurements in anesthetized dogs (ref. 3)	1	1.14	1,200	T	0.12	12				
		2	55.86	6,000	¶	0.86	0	2	44 (0%)§	40.1 (-30%)	

* Representative measurements in our subjects were used, namely, DL_{O_2} calculated from the CO data (true DL_{O_2}) = 57 ml per (minute \times mm Hg), total O_C = 7,200 ml per minute, total V_A = 5,400 ml, DL_{O_0} = 44 ml per (minute \times mm Hg), and capillary blood volume (VC) = 100 ml. † A-c O_2 gradients were taken from Figure 5. ‡ The numbers in this column were calculated from the A-c O_2 gradient of each compartment. Alveolar Po₂ was assumed to be 100 mm Hg, and capillary O_2 content for each compartment was obtained from a standard O_2 dissociation curve at pH of 7.40. § Per cent reduction from true DL_{O_2} of 57 ml per (minute \times mm Hg).

¶ Uneven $DL/\dot{Q}c$ was considered to be within each gas exchange unit so that all alveoli had the same rate of change of CO and $4O_2$ during breath holding.

result (2). Because this form of uneven DL/Qcmight explain the difference between observed DL_{O_2} and true DL_{O_2} seen in our subjects, we evaluated this possibility by obtaining representative values of pulsatile flow from the data published by Linderholm, Kimbel, Lewis, and DuBois (20) and then calculated an observed DL_{O_2} . We used the values of total Qc, Vc, and true DL_{0_2} listed in Table III, a pulse rate of 100 per minute, a systole of 0.12 second with a Qc of 300 ml per second, and diastole of 0.48 second with a Qc of 75 ml per second. No significant change in DL₀₂ was produced by this pattern of Qc, but if pulsatile blood flow were combined with uneven DL/Qc such as illustrated in Figures 1B and 1C, rapid pulsatile flow through capillaries of short length would permit some blood to pass through them during systole and thereby produce a very brief exposure to alveolar gas. Such a condition would decrease observed DL₀, without changing true DLo2. At the present time we have no method of separating uneven DL/Qc due to pulsatile flow from uneven DL/Qcfrom other causes, so that pulsatile flow may be a significant contributing factor to the uneven DL/Qc observed in our subjects.

Effect of uneven perfusion and diffusion per unit of lung volume secondary to the upright position. Evidence has been presented that in the upright posture the upper zones of the lungs have considerably less Qc and DL per unit of lung volume (VA) than is found in the lower zones (21, 24). To establish whether this type of uneven distribution might explain our data, we calculated the changes in DL_{0_2} and DL_{CO} that would result in a subject whose lungs had three zones of equal VA, but proportionately less amounts of DL and Qc in the upper zones. Numerical values for total VA, DL, and Oc are representative for our subjects, and the dimensions of the three zones were chosen on the basis of observations made by others using regional scanning of the lungs after the inspiration of radioactive CO and CO_2 (1, 21) (see Table III). This model of uneven distribution produced a decrease in DL_{0_2} of 12% and a decrease in DL_{CO} of 7%. Since these changes in part cancel out during the determination of uneven $DL/\dot{Q}c$ and also are too small to account for the 18% to 64% difference between observed DL_{0_2} and true DL_{0_2} seen in our subjects, we do not think the sitting position alone explains the uneven $DL/\dot{Q}c$ we calculated.

In addition, the effect of uneven distribution of alveolar volume (VA) with respect to DL (uneven VA/DL) resulting from the upright position was evaluated with compartmental dimensions based on the data reported by Burrows and co-workers (25). These calculations showed that there would be changes in DLO2 and DLCO but insufficient in magnitude or direction to produce significant changes in the calculated values of uneven DL/Qc. We conclude that uneven distribution of DL with respect to Qc within each gas exchange unit of the lung is the best explanation for our experimental findings. Although it is likely that there is uneven distribution of DL/Qc and DL/VA in gross anatomical zones of the lung, the value of observed DL₀, would be only slightly reduced. The comparison of observed DL₀₂ to true DL₀₂ is, therefore, not a good method for investigating this particular type of uneven distribution.

Comparison to other measurements of uneven DL/Qc. Piiper, Haab, and Rahn calculated the degree of uneven DL/Qc in anesthetized dogs from measurements of the alveolar-arterial O₂ gradient (3). They concluded that diffusion of O₂ into the blood takes place from two functional compartments, namely one small compartment containing 2% of the diffusing capacity and receiving 14% of total Qc and one large compartment containing 98% of the diffusing capacity and receiving 86% of Qc.6 If there were a similar pattern of uneven DL/Qc in the lungs of our human subjects, it would have no influence on the value of DL_{CO}, but DL_{O2} would decrease from its average "true" value of 57 ml per (minute \times mm Hg) to an observed DL₀₂ of 40.1 ml per (minute \times mm Hg), which is slightly greater then the experimentally determined average value of 33 ml per (minute X mm Hg). Preliminary studies in resting man using breath holding with 5% carbon monoxide (26) and the steady state O_2 diffusing capacity method (27)

⁶ These authors reported that a small portion of the perfusion (1.5%) probably behaves like a true shunt $(DL/\dot{Q}c = 0)$. This compartment was omitted in our analysis because the method of measuring uneven $DL/\dot{Q}c$ described in this paper is only influenced by perfusion that comes in contact with alveolar CO, acetylene, and ${}^{34}O_{2}$ during the breath-holding maneuver.

indicate that two-thirds of $\dot{Q}c$ is delivered to 20% of the diffusing capacity. In the subjects of the present study this pattern of uneven DL/ $\dot{Q}c$ would result in an observed DL₀₂ of approximately 30 ml per (minute \times mm Hg) (see Figure 3), which is in good agreement with the experimentally determined average value of 33 ml per (minute \times mm Hg).

Influence of uneven DL/Qc on the pulmonary alveolar to end capillary O_2 gradient (A-c O_2 gradient). The presence of uneven $DL/\dot{Q}c$ within the alveoli of the lungs has considerable bearing on the estimation of the size of the A-c O₂ gradient. Previous reports do not agree about the magnitude of this gradient in normal individuals or even in diseased subjects. On the basis of measurements of distribution of ventilation to perfusion, right to left shunting around the alveoli, and the single breath DL_{CO}, some authors have stated that even in the so-called alveolar-capillary block syndrome an A-c O₂ gradient does not exist (28, 29). On the other hand, more recently Johnson, Taylor, and De-Graff have cited evidence showing that the arterial O2 desaturation seen in certain diffuse restrictive diseases of the lung tissues may in part be explained by an A-c O₂ gradient produced by the presence of red blood cell transit times of different duration in the pulmonary capillary bed (30). After a careful analysis of the influence of both uneven DL/Oc and uneven ventilation-perfusion ratios upon the A-c O₂ gradient, Piiper and co-workers concluded that the O₂ gradient they measured in anesthetized dogs was principally due to uneven DL/Qc rather than uneven ventilation-perfusion ratios (3).

Because we have no data that permit the determination of the precise pattern of uneven $DL/\dot{Q}c$ present in the lungs of our subjects, its exact contribution to the A-c O₂ gradient cannot be calculated at the present time. As an alternative we chose four patterns of uneven $DL/\dot{Q}c$ fashioned after reports in the literature and calculated the A-c O₂ gradient that would result in a resting subject with values for true DL_{O_2} , $\dot{Q}c$, and Vc approximately the same as found in our subjects (see Table III). Using these values together with a mixed venous PO₂ of 42.5 mm Hg and an alveolar PO₂ of 100 mm Hg, one can calculate the rate at which the PO₂ in the red blood



FIG. 5. RATE OF RISE OF PO₂ IN A RED BLOOD CELL AS IT TRAVERSES A CAPILLARY IN THE LUNGS OF A SUBJECT WITH A MIXED VENOUS PO₂ OF 42.5 MM HG AND AN ALVEO-LAR PO₂ OF 100 MM HG. Oxygen diffusing capacity of the membrane was considered to be 73 ml per (minute \times mm Hg) and Vc 100 ml, which are representative values for our subjects. [For details of the calculation see (12)]. PO_{2,RBC} is the PO₂ in the capillary blood for any particular transit time. Note that when the capillary transit time falls below 0.2 second, a significant alveolar to end capillary O₂ gradient develops.

cells increases as they traverse the capillaries (12). This calculation is presented graphically in Figure 5, and from it we determined the end capillary Po₂ for a DL/ $\dot{Q}c$ compartment whose capillary transit time (TL) is known.⁷

A-c O_2 gradient secondary to the upright posture. Table IIIA shows the effect of uneven $DL/\dot{Q}c$ and uneven $VA/\dot{Q}c$ secondary to the upright posture on the A-c O_2 gradient. Because the shortest TL for this model is 0.78 second, essentially no A-c O_2 gradient is present.

A-c O_2 gradient secondary to the DL/Qc distribution observed in microscopic studies of the pulmonary capillary bed. Von Hayek (9) and Weibel (22) have pointed out that the capillary pathway from the pulmonary arterioles to the pulmonary veins is a meshwork of vessels whose length may

⁷ Calculation of end capillary Po₂ from Figure 5 requires the assumption of even distribution of Vc with respect to membrane diffusing capacity (*vide supra*). Once this assumption is accepted, the per cent of the membrane diffusing capacity of a compartment equals the same percentage of total Vc. TL of the compartment can then be calculated by the following relationship: TL = Vc of compartment/ $\dot{Q}c$ of compartment. If Vc is not distributed evenly with respect to membrane diffusing capacity, the end capillary Po₂ can still be calculated, but the procedure becomes more laborious.

vary from about 60 μ to 250 μ or longer. It has been postulated from in vivo observations that shorter routes stay open whereas longer ones may be intermittently closed off. Blood flow in some capillaries has been seen to stop and even reverse direction (7). On the basis of the above information we constructed a lung model in which, in each gas exchange unit, 50% of the blood flow goes through capillaries 80 μ long, and the other 50% traverses capillaries 240 μ long. In addition, 30% of the capillary blood volume contains blood that is not flowing (see Table IIIB). Although this pattern of uneven DL/Qc according to Equation 4 would produce an observed DL_{0_2} of 27 ml per (minute \times mm Hg) compared to the true DL_{O_2} of 57 ml per (minute \times mm Hg) (values similar to those seen in our subjects, the A-c O₂ gradient is barely perceptible (0.25 mm Hg).

A-c O_2 gradient secondary to the uneven DL/Qcsuggested by cinematographic observations of the pulmonary capillary bed in vivo. Schlosser, Heyse, and Bartels recently developed a method that permits the visualization of the subpleural pulmonary capillaries of rabbit lungs. They reported that the average transit time for a red blood cell through these capillaries is only 0.1 second (23). On the basis of their observations we calculated the observed DL_{O_2} and A-c O_2 gradient that would result if 5% of the capillary channels carrying blood from the pulmonary arteries to the pulmonary veins were 60 μ long and had a transit time of 0.1 second. The remaining 95% of the channels were assumed to be 120 μ long and would receive 79% of the total blood flow (see Table IIIC). According to Equation 4 this pattern of uneven DL/Qc results in an observed DL₀₂ of 38 ml per (minute \times mm Hg), a value 33% less than the true DL_{0} of 57 ml per (minute \times mm Hg). In the fast compartment the A-c O_2 gradient would be 16.5 mm Hg, and after mixing with the blood from the slow compartment, the total pulmonary A-c O_2 gradient would be 5 mm Hg. These data suggest that a significant amount of the difference between alveolar and arterial Po₂ could be due to uneven DL/Qc.

A-c gradient secondary to the pattern of uneven $DL/\dot{Q}c$ observed in anesthetized dogs. Piiper and co-workers concluded that in anesthetized dogs diffusion of O_2 in the pulmonary capillary blood

takes place from two functional compartments, a smaller one receiving 14% of Qc, but containing only 2% of the diffusing capacity, and a larger compartment with 86% of $\dot{Q}c$ and 98% of the diffusing capacity (vide supra). For our human subjects the A-c O₂ gradient for the smaller compartment according to Figure 5 would be 12 mm Hg and after mixing with the blood flowing through the larger compartment would decrease to 2 mm Hg. The comparable figure for the A-c O₂ gradient from both compartments calculated by Piiper and co-workers was 10 mm Hg. This discrepancy can be accounted for by a number of factors, including the larger value of DL_{0_2} in the human subjects [0.8 ml per (minute \times mm Hg) per kg vs. 0.3 ml per (minute \times mm Hg) per kg in dogs]; the difference in the O_2 dissociation curves used, which is quite critical in the calculation of the A-c O_2 gradient (31); and the fact that the fall in DL_{0_2} with rising intracapillary O2 recently demonstrated in vivo (5) was taken into account in the humans but not in the dogs.

Relationship between DL and the size of the A-c O₂ gradient. The data in Table III demonstrate an apparent paradox: A lung model such as IIIB may have a lower observed DL_{0_2} than another model, yet its A-c O₂ gradient may be smaller. Therefore the observed values of DL_{0_2} and DL_{CO} in the presence of uneven DL/Qcdo not necessarily give information relevant to the estimation of the size of the A-c O₂ gradient. The precise pattern of distribution of uneven DL/Qc may be of greater importance in determining this gradient than the absolute values of diffusing capacity. For example, in the four hypothetical lungs listed in Table III, the A-c O_2 gradient varied from essentially 0 to 5 mm Hg depending more on the pattern of uneven $DL/\dot{Q}c$ chosen than on the measured values of DL_{CO} or DL_{O_2} .

It has been stated that in normal subjects breathing air at rest or even during heavy exercise, as well as in patients with chronic airway obstruction, no A-c O₂ gradient will be present because of the large size of the diffusing capacity (12, 28, 32). All of these calculations have assumed even distribution of diffusing capacity to blood flow. The data presented in this report indicate that fairly marked degrees of uneven DL/ $\dot{Q}c$ may be present in resting man, and it is possible for this uneven $DL/\dot{Q}c$ to be distributed in a manner that produces an appreciable A-c O₂ gradient. In the face of this information we believe it is hazardous to judge the size of the A-c O₂ gradient on the basis of CO diffusing capacities alone. The distribution of diffusing capacity with respect to $\dot{Q}c$ must be taken into account.

Although our results do not permit the calculation of the A-c O_2 gradient, they do suggest that the development of more precise methods for the measurement of the pattern of uneven $DL/\dot{Q}c$ may permit an accurate estimation of its size.

Appendix

Derivation of Equation 4 in the text, which expresses observed DL_{O_2} as a function of the distribution of diffusing capacity (DL) with respect to pulmonary capillary blood flow (Qc) in the lung.

The total ${}^{34}O_2$ leaving the lungs per minute during breath holding equals the amount leaving via the pulmonary capillaries less the amount arriving in the mixed venous blood, or:

total ³⁴O₂ leaving lungs per minute in milliliters per

minute =
$$\frac{\dot{Q}c(\alpha b_{O_2})(Pc_{34_{O_2}} - Pmv_{34_{O_2}})}{760}$$
, [5]

where $\dot{Q}c$ is expressed in milliliters per minute, Pc_{34O_2} equals the end capillary $P^{34}O_2$ in millimeters Hg, Pmv_{34O_2} is the mixed venous $P^{34}O_2$ in millimeters Hg, and αb_{O_2} is the Bunsen solubility coefficient for total O_2 in the blood in milliliters per milliliter per standard atmosphere calculated from the capillary PO₂ present during breath holding and the O₂ capacity of the subject's blood.

If a lung with uneven $DL/\dot{Q}c$ could be divided into compartments within which diffusing capacity and $\dot{Q}c$ were distributed evenly, then as in Equation 2:

$$DL_1 = \frac{\dot{Q}c_1(\alpha b_{0_2})}{760} \ln\left(\frac{1}{K_1}\right),$$

$$DL_2 = \frac{\dot{Q}c_2(\alpha b_{0_2})}{760} \ln\left(\frac{1}{K_2}\right), \quad DL_3 = \dots \text{ etc.}, \quad [6]$$

where DL_1 , DL_2 , DL_3 , ..., $\dot{Q}c_1$, $\dot{Q}c_2$, ..., and K_1 , K_2 , ..., are, respectively, the true DL_{02} , $\dot{Q}c$, and K for each compartment. The amount of ${}^{34}O_2$ removed from each compartment per minute can be calculated by applying Equation 5, and for example for compartment 1 would equal:

$$\frac{Qc_1(\alpha b_{O_2})({}^{1}Pc_{34_{O_2}} - Pmv_{34_{O_2}})}{760},$$
 [7]

where ${}^{1}\text{Pc}_{34_{O_2}}$ equals the end capillary $P^{34}\text{O}_2$ in compartment 1.

For a two compartment lung the total milliliters of ${}^{34}O_2$ leaving the lung per minute during breath holding must equal the sum of the amount leaving the two compartments in the blood less the amount entering, or:

$$\frac{\dot{Q}c(\alpha b_{02})(Pc_{34_{02}} - Pmv_{34_{02}})}{760} = \frac{\dot{Q}c_1(\alpha b_{02})({}^{1}Pc_{34_{02}} - Pmv_{34_{02}})}{760} + \frac{\dot{Q}c_2(\alpha b_{02})({}^{2}Pc_{34_{02}} - Pmv_{34_{02}})}{760}.$$
 [8]

K for a compartment of the lung can be defined as in Equation 3 and for compartment 1 would be:

$$K_{1} = \frac{(\text{alveolar } P^{34}O_{2} - {}^{1}Pc_{34}O_{2})}{(\text{alveolar } P^{34}O_{2} - Pmv_{34}O_{2})}.$$
 [9]

The terms $Pc_{34_{0_2}}$, ${}^{1}Pc_{34_{0_2}}$, and ${}^{2}Pc_{34_{0_2}}$ can be eliminated from Equation 8 by substituting in their respective values obtained from Equations 3 and 9. The resultant equation simplifies to the following form:

$$\frac{1}{K} = \frac{\dot{Q}c}{\dot{Q}c_1(K_1) + \dot{Q}c_2(K_2)}.$$
 [10]

Equation 6 can be solved for K_1 and K_2 and then substituted into Equation 10, giving:

$$\frac{1}{K} = \frac{Qc}{\dot{Q}c_1e^{[-DL_1(760)/\dot{Q}c_1(\alpha bo_2)]}} + \dot{Q}c_2e^{[-DL_2(760)/\dot{Q}c_2(\alpha bo_2)]}}$$

Equation 9 can be substituted into Equation 2 so as to eliminate 1/K giving:

observed
$$DL_{0_2} = \frac{\dot{Q}c(\alpha b_{0_2})}{760}$$

 $\times \ln \left[\frac{\dot{Q}c}{\dot{Q}c_1 e^{[-DL_1(760)/\dot{Q}c_1(\alpha b_{0_2})]}} + \dot{Q}c_1 e^{[-DL_2(760)/\dot{Q}c_2(\alpha b_{0_2})]} \right].$ [12]

Acknowledgment

The authors wish to express their appreciation to Dr. Robert E. Forster for his interest and advice.

Addendum

While this paper was in press, we obtained an article by Thews and Witte (33), in which they included a theoretical analysis of the influence of nonuniform distribution of O_2 diffusion on the alveolar to end capillary O_2 gradient. Their calculations showed that if distribution of O_2 diffusing capacity is nonhomogeneous, the "apparent" O_2 diffusing capacity will always be less than the "true" O_2 diffusing capacity, and the end capillary O_2 tension will be lower than if diffusing capacity were uniformly distributed. Their findings, which are in agreement with ours, indicate that the steady state O_2 diffusing capacity is affected by uneven DL/Qc in a manner similar to that described in this report for the single breath O_2 diffusing capacity.

References

 Dollery, C. T., N. A. Dyson, and J. D. Sinclair. Regional variations in uptake of radioactive CO in the normal lung. J. appl. Physiol. 1960, 15, 411.

- Piiper, J. Unequal distribution of pulmonary diffusing capacity and the alveolar-arterial Po₂ differences: theory. J. appl. Physiol. 1961, 16, 493.
- Piiper, J., P. Haab, and H. Rahn. Unequal distribution of pulmonary diffusing capacity in the anesthetized dog. J. appl. Physiol. 1961, 16, 499.
- Piiper, J. Variations of ventilation and diffusing capacity to perfusion determining the alveolararterial O₂ difference: theory. J. appl. Physiol. 1961, 16, 507.
- Hyde, R. W., R. E. Forster, G. G. Power, J. Nairn, and R. Rynes. Measurement of O₂ diffusing capacity of the lungs with a stable O₂ isotope. J. clin. Invest. 1966, 45, 1178.
- Lilienthal, J. L., Jr., R. L. Riley, D. D. Proemmel, and R. E. Franke. An experimental analysis in man of the oxygen pressure gradient from alveolar air to arterial blood during rest and exercise at sea level and at altitude. Amer. J. Physiol. 1946, 147, 199.
- Wearn, J. T., A. C. Ernstene, A. W. Bromer, J. S. Barr, W. J. German, and L. J. Zschiesche. The normal behavior of the pulmonary blood vessels with observations on the intermittence of the flow of blood in the arterioles and capillaries. Amer. J. Physiol. 1934, 109, 236.
- Miller, W. S. The Lung, 2nd ed. Springfield, Ill., Charles C Thomas, 1947, pp. 74–77.
- 9. Von Hayek, H. The Human Lung, V. E. Krahl, Trans. New York, Hafner, 1960, p. 253.
- Sackner, M. A., K. A. Feisal, and D. N. Karsch. Size of gas exchange vessels in the lung. J. clin. Invest. 1964, 43, 1847.
- Rossier, P. H., A. A. Buhlmann, and K. Wiesinger. Respiration: Physiologic Principles and Their Clinical Applications, P. C. Luchsinger and K. M. Moser, Trans. St. Louis, C. V. Mosby, 1960, pp. 89–93.
- Staub, N. C. Alveolar-arterial oxygen tension gradient due to diffusion. J. appl. Physiol. 1963, 18, 673.
- Roughton, F. J. W. Transport of oxygen and carbon dioxide *in* Handbook of Physiology, Section 3, Respiration, W. O. Fenn and H. Rahn, Eds. Washington, D. C., American Physiological Society, 1964, vol. 1, p. 781.
- 14. Roughton, F. J. W., and R. E. Forster. Relative importance of diffusion and chemical reaction rates in determining rate of exchange of gases in the human lung, with special reference to true diffusing capacity of pulmonary membrane and volume of blood in the lung capillaries. J. appl. Physiol. 1957, 11, 290.
- Staub, N. C., J. M. Bishop, and R. E. Forster. Importance of diffusion and chemical reaction rates in O₂ uptake in the lung. J. appl. Physiol. 1962, 17, 21.
- Cander, L., and R. E. Forster. Determination of pulmonary parenchymal tissue volume and pulmonary capillary blood flow in man. J. appl. Physiol. 1959, 14, 541.
- 17. Bates, D. V., and R. V. Christie. Respiratory Func-

tion in Disease. Philadelphia, W. B. Saunders, 1964, p. 92.

- Krumholz, Richard A. Pulmonary membrane diffusing capacity and pulmonary capillary blood volume: an appraisal of their clinical usefulness. Amer. Rev. resp. Dis. 1966, 94, 195.
- Ackerman, E., and R. L. Berger. Reaction of oxyhemoglobin with carbon monoxide. Biophys. J. 1963, 3, 493.
- Linderholm, H., P. Kimbel, D. H. Lewis, and A. B. DuBois. Pulmonary capillary blood flow during cardiac catheterization. J. appl. Physiol. 1962, 17, 135.
- West, J. B. Regional differences in gas exchange in the lung of erect man. J. appl. Physiol. 1962, 17, 893.
- Weibel, E. R. Morphometrics of the Lung *in* Handbook of Physiology, Section 3, Respiration, W. O. Fenn and H. Rahn, Eds. Washington, D. C., American Physiological Society, 1964, vol. 1, p. 285.
- Schlosser, D., E. Heyse, and H. Bartels. Flow rate of erythrocytes in the capillaries of the lung. J. appl. Physiol. 1965, 20, 110.
- Bryan, A. C., L. G. Bentivoglio, F. Beerel, H. Mac-Leish, A. Zidulka, and D. Bates. Factors affecting regional distribution of ventilation and perfusion in the lung. J. appl. Physiol. 1964, 19, 395.
- Burrows, B., A. H. Niden, C. Mittman, R. C. Talley, and W. R. Barclay. Non-uniform pulmonary diffusion as demonstrated by the carbon monoxide equilibration technique: experimental results in man. J. clin. Invest. 1960, 39, 943.
- Hyde, R. W., R. E. Forster, J. Nairn, G. G. Power, and R. Rynes. Distribution of pulmonary diffusing capacity (DL) in relation to pulmonary capillary blood flow (Qc) (abstract). Physiologist 1965, 8, 199.
- Miller, J. M., and R. L. Johnson, Jr. Effect of nonuniform blood flow on oxygen diffusing capacity (abstract). Physiologist 1965, 8, 235.
- Finley, T. N., E. W. Swenson, and J. H. Comroe, Jr. The cause of arterial hypoxemia at rest in patients with "alveolar-capillary block syndrome." J. clin. Invest. 1962, 41, 618.
- 29. Hamer, J. Cause of low arterial oxygen saturation in pulmonary fibrosis. Thorax 1964, 19, 507.
- Johnson, R. L., Jr., H. F. Taylor, and A. C. DeGraff, Jr. Functional significance of a low pulmonary diffusing capacity for carbon monoxide. J. clin. Invest. 1965, 44, 789.
- Haab, P. E., J. Piiper, and H. Rahn. Simple method for rapid determination of an O₂ dissociation curve of the blood. J. appl. Physiol. 1960, 15, 1148.
- Ayres, S. M., and S. Giannelli, Jr. Causes of arterial hypoxemia in patients with obstructive pulmonary emphysema. Amer. J. Med. 1965, 39, 422.
- Thews, von G., and K. Witte. Der Einfluss statistisch verteilter ungleicher O₂-Diffusionswiderstände der Lungenmembran auf die O₂-Diffusionskapazität. Beitr. Silikose-Forsch. 1963, 5, 329.