

This is a repository copy of *The effect of aging and cardiorespiratory fitness on the lung diffusing capacity response to exercise in healthy humans*.

White Rose Research Online URL for this paper: http://eprints.whiterose.ac.uk/114363/

Version: Accepted Version

Article:

Coffman, KE, Carlson, AR, Miller, AD et al. (2 more authors) (2017) The effect of aging and cardiorespiratory fitness on the lung diffusing capacity response to exercise in healthy humans. Journal of Applied Physiology, 122 (6). pp. 1425-1434. ISSN 8750-7587

https://doi.org/10.1152/japplphysiol.00694.2016

© 2016, Journal of Applied Physiology. This is an author produced version of a paper published in Journal of Applied Physiology. Uploaded in accordance with the publisher's self-archiving policy.

Reuse

Unless indicated otherwise, fulltext items are protected by copyright with all rights reserved. The copyright exception in section 29 of the Copyright, Designs and Patents Act 1988 allows the making of a single copy solely for the purpose of non-commercial research or private study within the limits of fair dealing. The publisher or other rights-holder may allow further reproduction and re-use of this version - refer to the White Rose Research Online record for this item. Where records identify the publisher as the copyright holder, users can verify any specific terms of use on the publisher's website.

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



The effect of aging and cardiorespiratory fitness on the lung diffusing capacity response to exercise in healthy humans Authors Kirsten E. Coffman¹, Alex R. Carlson², Andrew D. Miller², Bruce D. Johnson², Bryan J. Taylor^{2,3} **Author Affiliations** ¹Mayo Graduate School, Mayo Clinic, Rochester, MN, USA ²Division of Cardiovascular Diseases, Department of Internal Medicine, Mayo Clinic, Rochester, MN, USA ³School of Biomedical Sciences, Faculty of Biological Sciences, University of Leeds, Leeds, UK **Contact Information for Corresponding Author** Bryan J. Taylor, PhD University Academic Fellow School of Biomedical Sciences Faculty of Biological Sciences University of Leeds Garstang 5.68 Leeds, UK LS2 9JT Office: +44(0)113 343 0482 E-mail: b.j.taylor@leeds.ac.uk

39 ABSTRACT

Aging is associated with deterioration in the structure and function of the pulmonary circulation. 40 We characterized the lung diffusing capacity for carbon monoxide (DLCO), alveolar-capillary 41 membrane conductance (Dm_{CO}), and pulmonary-capillary blood volume (V_C) response to 42 discontinuous incremental exercise at 25, 50, 75, and 90% of peak work (W_{peak}) in four groups: 43 1) Young [27 \pm 3 y, maximal oxygen consumption ($\dot{V}O_{2max}$) 110 \pm 18% age-predicted]; 2) 44 Young Highly-Fit (27 \pm 3 y, \dot{VO}_{2max} 147 \pm 8% age-predicted); 3) Old (69 \pm 5 y, \dot{VO}_{2max} 116 \pm 45 13% age-predicted); and 4) Old Highly-Fit (65 \pm 5 y, $\dot{V}O_{2max}$ 162 \pm 18% age-predicted). At rest 46 47 and at 90% W_{peak}, DLCO, Dm_{CO}, and V_C were decreased with age. At 90% W_{peak}, DLCO, Dm_{CO} and $V_{\rm C}$ were greater in Old Highly-Fit vs. Old adults. The slope of the DLCO-cardiac output (Q) 48 relationship from rest to end-exercise at 90% W_{peak} was not different between Young, Young 49 Highly-Fit, Old and Old Highly-Fit (1.35 vs. 1.44 vs. 1.10 vs. 1.35 ml_{CO}·mmHg⁻¹·Lblood⁻¹, P =50 0.388), with no evidence of a plateau in this relationship during exercise; this was also true for 51 Dm_{CO} - \dot{Q} and V_{C} - \dot{Q} . $\dot{V}O_{2max}$ was positively correlated with: 1) DLCO, Dm_{CO} , and V_{C} at rest. 2) 52 the rest to end-exercise change in DLCO, Dm_{CO}, and V_C. In conclusion, these data suggest that 53 despite the age-associated deterioration in the structure and function of the pulmonary 54 circulation, expansion of the pulmonary capillary network does not become limited during 55 exercise in healthy individuals regardless of age or cardiorespiratory fitness level. 56

57

58 **KEY WORDS:** Maximal aerobic capacity, lung diffusing capacity, pulmonary circulation,

59 alveolar-capillary membrane conductance, pulmonary-capillary blood volume

- 60
- 61

62 NEW & NOTEWORTHY

Healthy aging is a crucial area of research. This manuscript details how differences in age and cardiorespiratory fitness level affect lung diffusing capacity, particularly during heavy exercise. We conclude that highly fit older adults do not experience a limit in lung diffusing capacity during heavy exercise. Interestingly, however, we found that highly fit older individuals demonstrate greater values of lung diffusing capacity during heavy exercise than their less fit age-matched counterparts.

85 INTRODUCTION

Maximal aerobic capacity (\dot{VO}_{2max}) has been shown to, at least in part, determined by the structure and function of the pulmonary vasculature in health and chronic disease (1, 10, 22, 23, 25). For example, it has been shown that \dot{VO}_{2max} is positively correlated with resting pulmonary capillary blood volume (V_C) and pulmonary vasculature distensibility, and inversely related to pulmonary vascular resistance at maximal exercise in healthy individuals (23). This suggests that a larger, more distensible pulmonary vascular network is associated with greater aerobic exercise capacity in humans.

93

Measures of lung diffusing capacity for carbon monoxide (DLCO) and nitric oxide (DLNO), 94 alveolar-capillary membrane conductance (Dm_{co}) and pulmonary capillary blood volume (V_c) 95 are considered to reflect the pulmonary vascular response to whole-body exercise. Indeed, 96 increased cardiac output and pulmonary perfusion pressure during exercise cause a marked 97 expansion of the highly compliant pulmonary capillary network that is associated with an 98 increase DLCO, DLNO, Dmco and Vc (17, 27, 34). Additionally, it is thought that the 99 DLNO/DLCO ratio provides insight into the mechanism by which expansion of the pulmonary 100 capillary network during exercise occurs, with an increase in the ratio indicating a thinning of the 101 pulmonary capillary sheet (i.e. predominant vessel recruitment) and a decrease in the ratio 102 indicating a thickening of the blood sheet (i.e. predominant vessel distension) (13, 23). 103

104

Heathy aging is associated with a progressive deterioration in the structure and function of the
pulmonary circulation that is characterized by an increase in pulmonary vascular stiffness,
pulmonary vascular pressures and pulmonary vascular resistance (20, 24, 28). Additionally, from

maturity to senescence there is a decrease in resting V_C and Dm_{CO} that is consistent with a 108 reduction in alveolar-capillary surface area (1, 15). These age-related changes in the pulmonary 109 vasculature may impair recruitment and/or distension of the pulmonary capillaries during 110 exercise in healthy older adults, subsequently impairing the increase in alveolar-capillary surface 111 area needed for effective gas exchange and resulting in an excessive rise in pulmonary vascular 112 113 pressures relative to the metabolic demand of exercise. However, it has been shown that DLCO and V_C increase linearly relative to exercise intensity in old as well as young healthy adults (34), 114 indicating that expansion of the pulmonary capillaries does not become limited during exercise 115 116 in these individuals. This finding implies that the changes in the pulmonary circulation that occur with healthy aging are somewhat mild and not sufficient to affect pulmonary vascular expansion 117 and the recruitment of effective alveolar-capillary surface area for gas exchange during exercise 118 119 in healthy older adults.

120

The pulmonary vascular response to exercise in aged adults who have maintained a high 121 cardiorespiratory fitness is, however, currently less well characterized. Theoretically, better 122 maintenance of $\dot{V}O_{2max}$ through conditioning may cause the demand for \dot{Q} and pulmonary blood 123 124 flow during exercise to remain elevated in endurance trained highly fit older subjects compared to their younger counterparts. This, in the face of age-related alterations in the structure and 125 function of the pulmonary circulation, may predispose highly fit older adults to impairments in 126 127 pulmonary vascular expansion and pulmonary gas exchange relative to metabolic demand during exercise. Accordingly, the aim of the present study was to characterize the DLCO, Dm_{CO}, and V_C 128 response to incremental exhaustive exercise in healthy, aerobically-trained older adults relative 129 130 to their age-matched less aerobically fit counterparts, as well as younger adults of various

131 cardiorespiratory fitness levels. We hypothesized that those older individuals who had 132 maintained an elevated cardiorespiratory fitness would encroach upon their maximal ability to 133 expand the pulmonary vascular network during severe exercise, as evidenced by a plateau and/or 134 decrease in the rate of rise in one or more of DLCO, Dm_{CO} or V_{C} .

135

136 METHODS

137 Subjects

Sixteen young adults $(27 \pm 3 \text{ y})$ and 15 older adults $(67 \pm 6 \text{ y})$ who had pulmonary function 138 within normal limits participated in the study (Table 1). The subjects were sub-divided into four 139 groups according to age (≤ 30 y = "young"; ≥ 60 y = "old") and cardiorespiratory fitness (\dot{VO}_{2max} 140 \geq 140% of age predicted = "highly-fit"). Group 1) Young: age 27 ± 3 y (range 22 - 29), $\dot{V}O_{2max}$ 141 $110 \pm 18\%$ predicted (range 85 – 133) (n = 9); Group 2) Young Highly-Fit: age 27 ± 3 y (range 142 23 - 30), \dot{VO}_{2max} 147 ± 8% predicted (range 140 - 163) (n = 7); Group 3) Old: age 69 ± 5 y 143 (range 60 – 76), \dot{VO}_{2max} 116 ± 13% predicted (range 100 – 132) (*n* = 7, 1 female); Group 4) Old 144 Highly-Fit: age 65 ± 5 y (range 60 - 74), \dot{VO}_{2max} $162 \pm 18\%$ predicted (range 140 - 198) (n = 8, 145 1 female) (Table 1). All subjects were healthy and had no history of respiratory, cardiovascular, 146 147 or metabolic disease. Each participant gave written informed consent after being provided a detailed description of the study requirements. The experimental procedures were approved by 148 the Mayo Clinic Institutional Review Board and were performed in accordance with the ethical 149 150 standards of the Declaration of Helsinki.

151

152

153

154 Experimental Procedures

The experimental procedures were conducted during two laboratory visits separated by at least 2 155 but no more than 14 days. The subjects abstained from caffeine for 12 h and exercise for 24 h 156 prior to each visit. During visit 1, pulmonary function was assessed via full body 157 plethysmography (MedGraphics Elite Series Plethysmograph, Medical Graphics Corporation, St. 158 159 Paul, MN, USA) according to standard procedures (26). Next, subjects performed a maximal incremental exercise test on an electromagnetically braked cycle ergometer (Lode Corival, Lode 160 B.V. Medical Technology, Groningen, The Netherlands) for determination of peak work rate 161 (W_{peak}) and maximal oxygen consumption (VO_{2max}). Exercise was initiated at 60, 80, or 100 162 Watts (W), depending on self-reported fitness, and work rate was increased by 20 W every 2 min 163 until volitional exhaustion. W_{peak} was calculated as the sum of the final work rate completed plus 164 the fraction of the partially completed work rate before exhaustion. $\dot{V}O_{2max}$ was taken to be the 165 highest mean value within the final 20 seconds of exercise. 166

167

During visit 2, subjects performed discontinuous graded cycle exercise. Following 5 minutes of 168 quiet rest, participants cycled for 6 minutes at 25, 50, and 75% of W_{peak}, before cycling at 90% of 169 W_{peak} to volitional exhaustion. Between exercise bouts, subjects recovered quietly until heart rate 170 returned to within 10 bpm of resting values (at least 4 minutes). Additionally, a 1 min 'warm-up' 171 at 40% of the workload about to be completed was allowed before each exercise bout. Lung 172 173 diffusing capacity for carbon monoxide (DLCO) and nitric oxide (DLNO) and cardiac output (Q) were measured in duplicate at rest and during the final 90 s of each exercise bout via a rebreathe 174 technique. 175

176

177 Lung diffusing capacity and cardiac output

Lung diffusing capacity for carbon monoxide (DLCO), lung diffusing capacity for nitric oxide 178 (DLNO), and cardiac output (O) were assessed using a rebreathe technique as we have described 179 previously (5, 36). Using this technique, DLCO, DLNO, and Q are determined via the rate of 180 disappearance of CO, NO, and acetylene (C₂H₂), respectively. Briefly, subjects sat upright on a 181 cycle ergometer and breathed through a two-way switching valve (Hans Rudolph 4285 series, 182 Hans Rudolph, Kansas City, MO, USA) connected to a pneumotachometer (MedGraphics 183 PreVent Pneumotach, Medical Graphics Corporation, St. Paul, MN,USA), mass spectrometer 184 (Marquette 1100 Medical Gas Analyser, Perkin-Elmer, St. Louis, MO, USA) and NO analyzer 185 (Sievers 280i NOA, Sievers, Boulder, CO, USA). The inspiratory port of the switching valve was 186 open to room air or a 6-L anesthesia bag filled with 0.3% CO (C¹⁸O), 40 ppm NO, 9% He, 0.6% 187 C₂H₂, 35% O₂ and N₂ balance. The total volume of gas added to the rebreathe bag was 188 determined as the average tidal volume of the subject during the 20-30 s immediately prior to 189 each measurement. To ensure the volume of the test gas was consistent across multiple rebreathe 190 maneuvers the bag was filled using a timed switching circuit that, given a constant flow rate 191 from the gas tank, resulted in the desired volume. The test gas volume given by the switching 192 circuit was verified before exercise using a 3 L syringe. Following a normal expiration, subjects 193 were switched into the rebreathe bag and instructed to nearly empty the bag with each breath for 194 8-10 consecutive breaths. A respiratory frequency of 32 breaths per min was maintained by 195 following a metronome with inspiratory and expiratory tones; this respiratory rate was necessary 196 in order to collect enough data to correctly trace NO decay. If the subject's respiratory frequency 197 was above 32 breaths per min during exercise, the subject was allowed to breathe at a higher 198

rate. This maneuver was performed in duplicate at rest and during the final 90 s of each exercisebout.

201

From the measurements of DLCO and DLNO, alveolar-capillary membrane conductance (Dm_{CO}) and pulmonary capillary blood volume (V_C) were calculated (5, 32). The coefficient relating DLNO to Dm_{CO} (α -ratio) was set at 2.26 (5) such that Dm_{CO} was calculated as DLNO/ α -ratio. Next, the reaction rate of CO with hemoglobin (θ_{CO}) was calculated using the equation derived by Reeves and Park (5, 29) in which θ_{CO} is dependent on the capillary partial pressure of oxygen (P_{cap}O₂):

208
$$\frac{1}{\theta_{co}} = 0.008 * P_{cap}O_2 + 0.0156 \qquad \text{Equation (1)}$$

where $P_{cap}O_2$ is estimated as alveolar $PO_2 - VO_2/(DLCO \times 1.23)$ with partial pressures in mmHg and VO_2 in ml/min. Finally, the values of Dm_{CO} and θ_{CO} were used to solve for V_C according to the following equation derived by Roughton and Forster (31):

212
$$\frac{1}{DLCO} = \frac{1}{DmCO} + \frac{1}{\theta_{CO}*Vc}$$
 Equation (2)

Capillary blood was sampled from an earlobe and measured for hemoglobin (Hb) concentration via centrifugal hematology (QBC Autoreader, Bector Dickinson, Port Matilda, PA). V_C was then corrected for standard concentrations of Hb in men (14.6 g/dl) and women (13.4 g/dl) as calculated $V_C \times$ (standard Hb concentration/measured Hb concentration).

217

218 Lung Diffusing Capacity-Cardiac Output Slope

We calculated the slope of the DLCO- \dot{Q} , Dm_{CO}- \dot{Q} , and V_C- \dot{Q} relationship from rest to endexercise in each experimental group by plotting DLCO, Dm_{CO} or V_C as a function of \dot{Q} . These slopes represent unit DLCO, Dm_{CO}, and V_C changes per unit change in pulmonary vascular blood flow (Q) and provide an indirect measure of the hemodynamic response of the pulmonary
circulation to exercise.

224

225 Statistical Analysis

One-way ANOVA with Tukey-Kramer post-hoc analysis (two-tailed) was used to compare 1) 226 subject characteristics, 2) pulmonary function, and 3) measures of DLCO, Dm_{CO}, and V_C at rest 227 and at 90% of W_{peak}. The effect of exercise on DLCO, Dm_{CO}, V_C, and DLNO/DLCO was tested 228 using a linear mixed effects model. In this model, the dependent variable was DLCO, Dm_{CO}, V_C 229 230 or DLNO/DLCO, and the five exercise levels (rest, 25, 50, 75 and 90% W_{peak}) were treated as repeated measures with Q as the continuous variable. Age (young or old) and cardiorespiratory 231 fitness (normal or highly-fit) were included as independent predictor variables. That is, using this 232 model the response of a given variable to exercise as well as the offset in baseline values are 233 assessed as a function of age and/or cardiorespiratory fitness. Coefficient of determination (r^2) 234 was computed to assess the proportion of \dot{VO}_{2max} that was predicted by 1) resting DLCO, resting 235 Dm_{CO} , and resting V_C ; 2) the change (Δ) in DLCO, Dm_{CO} , and V_C from rest to end-exercise; and 236 3) the DLCO- \dot{Q} , Dm_{CO}- \dot{Q} , and V_C- \dot{Q} slope in response to exercise. In all analyses, the acceptable 237 type I error was set at P < 0.05. Results are expressed as mean \pm SD. The linear mixed effects 238 model was performed in Matlab (version R2016a, MathWorks, Natick, MA); all other statistical 239 analyses were performed using IBM SPSS Statistics 20 for Windows (IBM, Armonk, NY). 240

241

242 **RESULTS**

243 Subjects

Subject characteristics and pulmonary function are shown in Table 1. Group mean age was not
different in Young vs. Young Highly-Fit, or in Old vs. Old Highly-Fit. In addition, all subject

groups were well matched for height, body mass, and BMI. Absolute and relative (*to body mass*) \dot{VO}_{2max} and W_{peak} were greater in Young Highly-Fit compared to Young, Old and Old Highly-Fit (all P < 0.01, Table 1). In addition, \dot{VO}_{2max} (absolute and relative) and W_{peak} were lower in Old versus Young, Young Highly-Fit and Old Highly-Fit (all P < 0.01, Table 1). Interestingly, however, neither \dot{VO}_{2max} nor W_{peak} was different in Young vs. Old Highly-Fit (Table 1).

251

252 Lung Diffusing Capacity at Rest and at 90% W_{peak}

At rest, group mean DLCO, Dm_{CO} , and V_C were lower in Old vs. Young ($P \le 0.003$) and Young Highly-Fit (P < 0.001) (Table 2 and Figure 1). In addition, group mean resting DLCO, Dm_{CO} , and V_C were lower in Old Highly-Fit compared to Young Highly-Fit (P < 0.001, 0.016, and 0.021, respectively) (Table 2 and Figure 1). No other differences in resting measures of DLCO, Dm_{CO}, and V_C were observed between the four experimental groups (Table 2).

258

At 90% of W_{peak} , group mean DLCO, Dm_{CO} , and V_{C} were lower in Old vs. Young (P < 0.001), 259 Young Highly-Fit (P < 0.001), and Old Highly-Fit ($P \le 0.050$) (Table 2 and Figure 1). Also, 260 group mean DLCO and Dm_{CO} at 90% of W_{peak} were lower in Old Highly-Fit compared to Young 261 $(P \le 0.019)$ and Young Highly-Fit $(P \le 0.001)$. Moreover, V_C was lower in Old Highly-Fit vs. 262 Young Highly-Fit at 90% of Wpeak (P = 0.016) (Table 2 and Figure 1). These data suggest that, 263 at rest and during exercise, lung diffusing capacity and its component parts (i.e. DLCO, Dm_{CO}, 264 265 and V_C) are decreased with age. Additionally, lung diffusing capacity and its component parts are greater during near-maximal exercise highly-fit older adults compared to their age matched 266 267 less fit counterparts.

268

269

270

Lung Diffusing Capacity Response to Exercise: effect of age and cardiorespiratory fitness 271 DLCO, Dm_{CO}, and V_C rose steadily with increasing Q during exercise, with no evidence of a 272 plateau and/or decrease in the rate of rise in these variables from rest to end-exercise in any of 273 the experimental groups (Figure 1). These data suggest that no group encroached upon their 274 maximal ability to expand the pulmonary vascular network and increase lung surface area for gas 275 exchange during exercise. Throughout exercise, DLCO was significantly lower with greater age 276 (P < 0.001) but significantly higher with greater fitness (P = 0.016) (Figure 2). Dm_{CO} and V_C 277 were significantly lower with greater age (P < 0.001) throughout exercise; the effect of fitness 278 was not significant (Figure 2). The relationships between lung diffusing capacity and its 279 280 component variables (i.e. DLCO, Dm_{CO} , and V_{C}) and \dot{Q} from rest to end-exercise are shown in Figure 2. The rate of rise in DLCO, Dm_{CO} , and V_{C} relative to \dot{Q} during exercise was remarkably 281 similar between the four experimental groups. Indeed, there was no significant effect of age or 282 fitness on the slope of the DLCO- \dot{Q} , Dm_{CO}- \dot{Q} and V_C- \dot{Q} response to exercise (Figure 2). These 283 data suggest that all groups experienced a similar pulmonary vascular response to exercise. 284

285

The DLNO/DLCO ratio decreased from rest to throughout exercise in Young (P = 0.033), Old (P = 0.028), and Old Highly-Fit (P = 0.004), indicating that pulmonary capillary expansion during exercise was at least partially achieved via vessel distension; this fall was not significant in the Young Highly-Fit individuals (P = 0.051) (Figure 6). Furthermore, there was no significant effect of age (P = 0.055) or fitness on the rate of fall in DLNO/DLCO from rest to end-exercise (Figure 6). 292

293

294 Relationship of Lung Diffusing Capacity and VO_{2max}

The relationships between $\dot{V}O_{2max}$ and 1) resting measures of DLCO, Dm_{CO} , and V_{C} , 2) the 295 change (Δ) in DLCO, Dm_{CO}, and V_C from rest to end-exercise, and 3) the DLCO- \dot{Q} , Dm_{CO}- \dot{Q} , 296 and V_C-Q slope in response to exercise for all 31 subjects are shown in Figure 3, Figure 4 and 297 Figure 5, respectively. A significant positive correlation was found between $\dot{V}O_{2max}$ and resting 298 measures of DLCO ($r^2 = 0.587$, P < 0.001), Dm_{CO} ($r^2 = 0.402$, P < 0.001), and V_C ($r^2 = 0.584$, P 299 < 0.001) (Figure 3). Similarly, there was a significant positive relationship between $\dot{V}O_{2max}$ and 300 the rest to end-exercise change (Δ) in DLCO ($r^2 = 0.502$, P < 0.001), Dm_{CO} ($r^2 = 0.412$, P < 0.001) 301 0.001), and V_c ($r^2 = 0.273$, P = 0.003) (Figure 4). Finally, there was a positive relationship 302 between $\dot{V}O_{2max}$ and the DLCO- \dot{Q} slope in response to exercise ($r^2 = 0.152$, P = 0.030); the 303 relationship between $\dot{V}O_{2max}$ and the Dm_{CO} - \dot{Q} and V_C - \dot{Q} slope in response to exercise was not 304 statistically significant (Figure 5). Together, these data suggest that a higher baseline DLCO, 305 Dm_{CO}, and V_C, as well as a larger increase these values in response to exercise are associated 306 with greater maximal aerobic capacity in young and older adults regardless of cardiopulmonary 307 fitness level. 308

309

310 DISCUSSION

311 Major Findings: comparison to previous findings

In the present study, we characterized the lung diffusing capacity for carbon monoxide (DLCO), alveolar-capillary membrane conductance (Dm_{CO}), and pulmonary-capillary blood volume (V_C) response to discontinuous incremental exercise in healthy, aerobically-trained older adults relative to their age-matched less fit counterparts as well as younger adults. We hypothesized

that healthy older adults (~65 years old) with a high cardiorespiratory fitness level ($\dot{V}O_{2max}$ 316 ~162% of age-predicted) would encroach upon their maximal ability to expand the pulmonary 317 capillary network during severe cycle exercise, as evidenced by a limit in DLCO, Dm_{CO}, and V_C 318 near maximal end-exercise. The main findings were: 1) healthy aging was associated with a 319 decrease in DLCO, Dm_{CO}, and V_C at rest and during near maximal exercise (Table 2); 2) better 320 maintained cardiorespiratory fitness was associated with greater DLCO, Dm_{CO,} and V_C during near-321 maximal exercise in older adults (Table 2); 3) there was no plateau (i.e. a limitation) in the DLCO, 322 Vc and Dm_{CO} response to exercise in any subject group regardless of age or cardiorespiratory fitness 323 level (Figure 1); 4) throughout exercise, DLCO, Dm_{CO}, and V_C were systematically lower in older 324 325 individuals regardless of fitness, and DLCO was systematically higher with maintained cardiorespiratory fitness regardless of age; and 5) the slope, or rate of rise, of the DLCO- \dot{Q} , Dm_{CO}- \dot{Q} , 326 and V_C-Q relationship from rest to end-exercise was not different between subjects regardless of age 327 or cardiorespiratory fitness level (Figure 2). 328

329

Our findings are confirmatory that healthy aging is associated with a progressive decline in 330 resting DLCO (1, 6, 9, 11, 15, 33, 34). For example, Guénard and Marthan demonstrated that 331 332 both DLCO and DLCO relative to minute ventilation (DLCO/ \dot{V}_E) are negatively correlated with age according to the equations DLCO = $126 - 0.90 \times \text{age}$ and DLCO/ $\dot{V}_E = 13.5 - 0.85 \times \text{age}$, 333 334 respectively (15). Likely contributors to this age-related reduction in DLCO are a decrease in the number of capillaries perfusing the lungs with a reduction in Vc (6, 7, 12), as well as a decrease 335 in alveolar surface area with a consequent reduction in membrane diffusing capacity (6, 12). Our 336 337 findings are also in agreement with previous reports that exercise is associated with a marked, mostly linear, increase in DLCO, Vc and Dm_{CO} in healthy adults (16, 18, 27, 34). This increase, 338

Downloaded from http://jap.physiology.org/ by 10.220.33.5 on March 30, 2017

at least in part, reflects expansion of the highly compliant pulmonary capillary network
secondary to the elevation in Q and pulmonary perfusion pressure that occurs with exercise.

341

To date, however, there is a relative paucity of data regarding the effect of healthy aging on the 342 lung diffusing capacity response to exercise. In a limited number of subjects (n = 12) of a broad 343 age range (23 to 79 years), Tamhane et al. reported that DLCO, Dm_{CO} and lung diffusing 344 capacity for nitric oxide (DLNO) increased linearly with Q from rest to exercise regardless of 345 age (34). However, while the authors found that age was a significant determinant of resting 346 347 DLNO, no such analysis was done examining the influence of age on the lung diffusing capacity response to *exercise*. Additionally, we are unaware of any previous study that has examined the 348 influence of healthy aging *plus* maintained cardiorespiratory fitness on lung diffusing capacity 349 during exercise. In combination, the present findings suggest that despite the age-associated 350 deterioration in the structure and function of the pulmonary circulation, expansion of the 351 pulmonary capillary network does not become limited during severe exercise (i.e. there is still a 352 reserve to recruit the pulmonary vasculature) in healthy individuals regardless of age or 353 cardiorespiratory fitness level. In addition, we suggest that maximal oxygen consumption 354 (\dot{VO}_{2max}) is positively related DLCO, Dm_{CO} , and V_{C} , both at rest and in response to exercise, across 355 all ages and cardiorespiratory fitness levels. 356

357

358 Lung Diffusing Capacity Response to Exercise: effect of age and cardiorespiratory fitness

Exercise is associated with an increase in cardiac output and pulmonary perfusion pressure that causes both recruitment of under-perfused pulmonary capillaries and distension of already perfused pulmonary blood vessels, as evidenced by an increase in DLCO, Dm_{CO} , and V_C (16, 18, 22, 23, 36). The resulting increase in pulmonary blood flow and marked expansion of the highly 363

compliant pulmonary vasculature acts to increase the alveolar-capillary surface area available for effective gas exchange. 364

365

Healthy aging is associated with a deterioration in the structure and function of the pulmonary 366 vasculature that is characterized by an increase in pulmonary vascular stiffness, pulmonary 367 vascular pressures and pulmonary vascular resistance (20, 24, 28), as well as reductions in 368 resting V_C and Dm_{CO} (1, 9, 12, 15). It has been shown, however, that despite these age-related 369 changes DLCO does not become limited during heavy to maximal exercise in the older adult of 370 371 average cardiorespiratory fitness (34). This is likely because the age-associated decline in the maximal metabolic demand of exercise occurs at rate equal to or greater than the deleterious 372 changes in the pulmonary circulation (14, 19, 37). However, it is conceivable that healthy older 373 individuals who have maintained cardiorespiratory fitness, and thus metabolic demand, at an 374 exceedingly high level may experience a limit in the capacity of the pulmonary vasculature to 375 expand relative to the demand for pulmonary blood flow during exercise. 376

377

In the present study, we found that DLCO, Dm_{CO}, and V_C increased steadily with increasing 378 cardiac output (Q), with no evidence of a plateau in these variables in both young and old 379 subjects, regardless of cardiorespiratory fitness level (Figure 1). Moreover, the slope of the 380 DLCO-Q, Dm_{CO}-Q, and V_C-Q relationship from rest to end-exercise was not different between 381 382 Young, Young Highly-Fit, Old and Old Highly-Fit subjects (Figure 2). This suggests that the recruitment and/or distension of the pulmonary capillaries and thus expansion of alveolar-383 capillary surface area remain adequate for the metabolic demand of exercise regardless of age 384 385 and cardiorespiratory fitness level. In agreement with previous findings, DLCO, Dm_{CO}, and V_C

at rest and during exercise were decreased with advanced age (1, 6, 11, 12, 15, 33, 34). Interestingly, however, we also found that DLCO, Dm_{CO} , and V_C were greater during nearmaximal exercise in highly fit older individuals compared to their less fit counterparts (Figure 1). Additionally, regardless of age, maintained cardiorespiratory fitness was associated with a significantly greater DLCO from rest through to maximal exercise.

391

It has been suggested previously that exercise training has no effect on DLCO and its 392 components parts (8, 30). For example, Reuschlein et al. reported no change in DLCO or Vc at 393 394 rest and during submaximal exercise from before to after 5 months of combined strength and endurance training (30). By contrast, it has been shown that cardiac and great vessel function is 395 better in older, habitually active, fit adults relative to their more sedentary counterparts. Indeed, 396 Arbab-Zadeh et al. reported that prolonged, sustained endurance training improves stroke 397 volume for a giving filling pressure and preserves left ventricular compliance in aged adults such 398 that the capillary wedge pressure-LV end diastolic volume curve in Masters athletes was 399 indistinguishable from that of young, sedentary control subjects (2). In addition, central arterial 400 compliance is 20-35% greater in endurance-trained middle-aged and older men compared to their 401 402 less active age matched counterparts (35). Furthermore, 3 months of aerobic exercise training increases central arterial compliance (~25%) in middle-aged men (~53 years) to a level similar to 403 that observed in older endurance-trained men (35), possibly due to modified cross-linking of 404 405 "stretched" collagen fibres and/or a reduction in the chronic suppressive influence exerted by sympathetic adrenergic tone (4, 35). Theoretically, it is entirely possible that habitual physical 406 activity may also better preserve the function of the pulmonary circulation, attenuating the age-407 408 related decline in pulmonary vascular distensibility (28) and compliance, allowing for greater

409 expansion of the pulmonary capillary network with increasing cardiac output, and thus 410 facilitating a greater DLCO, Dm_{CO} , and V_C response to exercise in highly fit older individuals 411 compared to their less fit counterparts. However, whether preservation of cardiorespiratory 412 fitness does indeed 'protect' against the age-related deterioration of the pulmonary circulation 413 cannot be deduced from the present findings, and as such remains purely speculative.

414

415 Expansion of the Pulmonary Capillary Network during Exercise: Recruitment vs. 416 Distension

The mechanism by which the pulmonary vasculature expands to accept increased blood flow and 417 increase effective alveolar-capillary surface area during exercise is also of interest. In the healthy 418 pulmonary circulation, vessel distensibility is largely independent of vessel size, location and 419 animal species (21), but appears to be lower in older adults relative to their younger counterparts 420 (28). Assessing the change in the ratio between DLNO and DLCO during exercise may allow 421 insight into whether pulmonary vascular volume increases due to recruitment or distension of the 422 pulmonary capillaries (13, 23, 27). Specifically, a fall in the DLNO/DLCO ratio with exercise is 423 thought to indicate a disproportionate rise in V_C relative to Dm_{CO}, which in turn suggests 424 predominant pulmonary capillary distension over recruitment (13). In the present study, all 425 groups with the exception of the Young Highly-Fit (P = 0.051) demonstrated a significant 426 reduction in the DLNO/DLCO ratio from rest to end-exercise (Figure 3). This finding suggests 427 that despite the previously reported decay in pulmonary vascular distensibility associated with 428 healthy aging, increased blood flow through the pulmonary circulation during exercise is at least 429 partially achieved via distension of the pulmonary capillaries in old as well as young individuals. 430

431

Downloaded from http://jap.physiology.org/ by 10.220.33.5 on March 30, 2017

432 Relationship between Lung Diffusing Capacity Response to Exercise and VO_{2max}

The importance of pulmonary vascular hemodynamics in determining \dot{VO}_{2max} in both health and disease is well known (10, 23). Fujii et al. reported that slope of the mean pulmonary artery pressure-to-cardiac output relationship was negatively correlated with \dot{VO}_{2max} in COPD patients (10). That is, a more compliant pulmonary vasculature, or one that experiences lower vascular pressures, allows for greater maximal aerobic capacity in healthy individuals as well as diseased patients.

439

In this study, we did not examine pulmonary vascular pressures in response to exercise. 440 Conceptually, however, a greater increase lung diffusing capacity relative to cardiac output in 441 response to exercise could be indicative of a greater ability of the pulmonary vasculature to 442 expand and accept the increase in pulmonary blood flow whilst minimizing the increase in 443 pulmonary vascular pressure. That is, it is conceivable that a steeper DLCO-Q slope during 444 exercise is reflective of a more compliant pulmonary vascular network. In support of this notion, 445 we found that DLCO- \dot{Q} was positively correlated to $\dot{V}O_{2max}$ across all subjects (Figure 6). In 446 addition, resting values of DLCO, Dm_{CO}, and V_C, as well as the absolute change in these 447 variables from rest to maximal exercise, are positively correlated with VO_{2max} across all subjects 448 (Figures 3 and 4). That expansion of the pulmonary vasculature appears not to reach a maximum 449 during exercise (Figure 1) likely serves to allow an increase in the alveolar-capillary surface area 450 451 for effective gas exchange whilst minimizing the exercise-induced increase in pulmonary arterial pressure, pulmonary vascular resistance and right-ventricular afterload, and thus allowing a 452 greater maximum cardiac output and maximum oxygen consumption in healthy adults regardless 453 454 of age and cardiorespiratory fitness (22, 23, 27).

455

456 **Technical Considerations**

457 *Effect of acetylene solubility upon calculation of cardiac output*

A concern with the use of acetylene uptake in the noninvasive determination of Q is that an 458 assumption, rather than direct assessment, of acetylene solubility (λ) in individual subjects can 459 result in considerable error in the measurement of Q. For example, Barker et al. suggested that 460 failure to account for inter-subject variability in λ can lead to substantial underestimation (up to 461 27%) or overestimation (up to 13%) of \dot{Q} in young healthy adults (3). In the present study, we 462 463 did not measure, and subsequently account for, between-subject differences in λ upon the calculation of Q. However, while any such error may have negatively impacted the accuracy of 464 our measure of Q, we are able to demonstrate good within-session within-subject reliability of 465 our acetylene derived \dot{Q} measure at rest (CV; coefficient of variation = 8.5%). This is 466 comparable with the CV reported previously by our group for open-circuit acetylene wash-in 467 estimated Q (17). In addition, although not assessed in the present study, it is likely that the 468 variability in our measure of Q improved to ~4% during exercise (17). As such, based on the 469 relatively low CV (i.e. good reproducibility) along with the repeated measures design of our 470 study, we are confident that our findings are not greatly affected by any underlying variability in 471 λ between individual subjects. 472

473

474 CONCLUSION

475 In conclusion, DLCO, Dm_{CO} , and V_C are decreased with age and increased with greater 476 cardiorespiratory fitness in older individuals near maximal exercise. Interestingly, there is a 477 systematic increase in DLCO throughout exercise with maintained cardiorespiratory fitness, regardless of age. Older highly fit individuals do not appear to encroach upon a pulmonary vascular limit, and expansion of the pulmonary capillary network is able to adequately increase DLCO, Dm_{CO} , and V_{C} , even during heavy exercise. Furthermore, the response (i.e., rate of increase) of DLCO, Dm_{CO} , and V_{C} to exercise is not altered by age and/or cardiorespiratory fitness level. Future studies should incorporate measures of pulmonary vascular pressures in order to elucidate the relationship between increases in lung diffusing capacity and the pulmonary vascular response to exercise.

485

486 ACKNOWLEDGEMENTS

The authors would like to acknowledge Josh O'Malley for his work as a research technologist during these studies. KEC is supported by Mayo Graduate School and National Heart, Lung, And Blood Institute of the National Institutes of Health (NIH) grant F31HL131076. BJT is supported by American Heart Association grant AHA12POST12070084. This study was funded by NIH grant HL71478.

492 493	REFERENCES
493 494	KEFERENCES
495	1. Al Dandachi G, Londner C, Caumont-Prim A, Plantier L, Chevalier-Bidaud B,
496	Toussaint JF, Desgorces FD, and Delclaux C. Ageing and endurance training effects on
497	quantity and quality of pulmonary vascular bed in healthy men. Respir Res 15: 8, 2014.
498	2. Arbab-Zadeh A, Dijk E, Prasad A, Fu Q, Torres P, Zhang R, Thomas JD, Palmer
499	D, and Levine BD. Effect of aging and physical activity on left ventricular compliance.
500	Circulation 110: 1799-1805, 2004.
501	3. Barker RC, Hopkins SR, Kellogg N, Olfert IM, Brutsaert TD, Gavin TP, Entin PL,
502	Rice AJ, and Wagner PD. Measurement of cardiac output during exercise by open-circuit
503	acetylene uptake. J Appl Physiol (1985) 87: 1506-1512, 1999.
504	4. Bruel A, Ortoft G, and Oxlund H. Inhibition of cross-links in collagen is associated
505	with reduced stiffness of the aorta in young rats. Atherosclerosis 140: 135-145, 1998.
506	5. Ceridon ML, Beck KC, Olson TP, Bilezikian JA, and Johnson BD. Calculating
507	alveolar capillary conductance and pulmonary capillary blood volume: comparing the multiple-
508	and single-inspired oxygen tension methods. J Appl Physiol (1985) 109: 643-653, 2010.
509	6. Chang SC, Chang HI, Liu SY, Shiao GM, and Perng RP. Effects of body position and
510	age on membrane diffusing capacity and pulmonary capillary blood volume. Chest 102: 139-142,
511	1992.
512	7. Crapo RO, Morris AH, and Gardner RM. Reference values for pulmonary tissue
513	volume, membrane diffusing capacity, and pulmonary capillary blood volume. Bull Eur
514	Physiopathol Respir 18: 893-899, 1982.
515	8. Dempsey JA, Gledhill N, Reddan WG, Forster HV, Hanson PG, and Claremont AD.
516	Pulmonary adaptation to exercise: effects of exercise type and duration, chronic hypoxia and
517	physical training. Ann N Y Acad Sci 301: 243-261, 1977.

518 9. Ehrsam RE, Perruchoud A, Oberholzer M, Burkart F, and Herzog H. Influence of
519 age on pulmonary haemodynamics at rest and during supine exercise. *Clin Sci (Lond)* 65: 653520 660, 1983.

Fujii T, Kurihara N, Fujimoto S, Hirata K, and Yoshikawa J. Role of pulmonary
vascular disorder in determining exercise capacity in patients with severe chronic obstructive
pulmonary disease. *Clin Physiol* 16: 521-533, 1996.

524 11. Garcia-Rio F, Dorgham A, Galera R, Casitas R, Martinez E, Alvarez-Sala R, and
525 Pino JM. Prediction equations for single-breath diffusing capacity in subjects aged 65 to 85
526 years. *Chest* 142: 175-184, 2012.

527 12. Georges R, Saumon G, and Loiseau A. The relationship of age to pulmonary membrane
528 conductance and capillary blood volume. *Am Rev Respir Dis* 117: 1069-1078, 1978.

529 13. Glenet SN, De Bisschop C, Vargas F, and Guenard HJ. Deciphering the nitric oxide to
530 carbon monoxide lung transfer ratio: physiological implications. *J Physiol* 582: 767-775, 2007.

531 14. Goldspink DF. Ageing and activity: their effects on the functional reserve capacities of
532 the heart and vascular smooth and skeletal muscles. *Ergonomics* 48: 1334-1351, 2005.

533 15. Guenard H, and Marthan R. Pulmonary gas exchange in elderly subjects. *Eur Respir J*534 9: 2573-2577, 1996.

Hsia CC, McBrayer DG, and Ramanathan M. Reference values of pulmonary
diffusing capacity during exercise by a rebreathing technique. *Am J Respir Crit Care Med* 152:
658-665, 1995.

Johnson BD, Beck KC, Proctor DN, Miller J, Dietz NM, and Joyner MJ. Cardiac
output during exercise by the open circuit acetylene washin method: comparison with direct
Fick. *J Appl Physiol (1985)* 88: 1650-1658, 2000.

Johnson RL, Jr., Spicer WS, Bishop JM, and Forster RE. Pulmonary capillary blood
volume, flow and diffusing capacity during exercise. *J Appl Physiol* 15: 893-902, 1960.

543 19. Kasch FW, Boyer JL, Van Camp S, Nettl F, Verity LS, and Wallace JP.
544 Cardiovascular changes with age and exercise. A 28-year longitudinal study. *Scand J Med Sci*545 *Sports* 5: 147-151, 1995.

546 20. Kovacs G, Berghold A, Scheidl S, and Olschewski H. Pulmonary arterial pressure
547 during rest and exercise in healthy subjects: a systematic review. *Eur Respir J* 34: 888-894,
548 2009.

549 21. Krenz GS, and Dawson CA. Flow and pressure distributions in vascular networks
550 consisting of distensible vessels. *Am J Physiol Heart Circ Physiol* 284: H2192-2203, 2003.

La Gerche A, MacIsaac AI, Burns AT, Mooney DJ, Inder WJ, Voigt JU, Heidbuchel
H, and Prior DL. Pulmonary transit of agitated contrast is associated with enhanced pulmonary
vascular reserve and right ventricular function during exercise. *J Appl Physiol (1985)* 109: 13071317, 2010.

Lalande S, Yerly P, Faoro V, and Naeije R. Pulmonary vascular distensibility predicts
aerobic capacity in healthy individuals. *J Physiol* 590: 4279-4288, 2012.

Lam CS, Borlaug BA, Kane GC, Enders FT, Rodeheffer RJ, and Redfield MM.
Age-associated increases in pulmonary artery systolic pressure in the general population. *Circulation* 119: 2663-2670, 2009.

Lewis GD, Bossone E, Naeije R, Grunig E, Saggar R, Lancellotti P, Ghio S, Varga J,
Rajagopalan S, Oudiz R, and Rubenfire M. Pulmonary vascular hemodynamic response to
exercise in cardiopulmonary diseases. *Circulation* 128: 1470-1479, 2013.

563 26. Miller MR, Hankinson J, Brusasco V, Burgos F, Casaburi R, Coates A, Crapo R,

564 Enright P, van der Grinten CP, Gustafsson P, Jensen R, Johnson DC, MacIntyre N,

565 McKay R, Navajas D, Pedersen OF, Pellegrino R, Viegi G, Wanger J, and Force AET.

566 Standardisation of spirometry. *Eur Respir J* 26: 319-338, 2005.

567 27. Pavelescu A, Faoro V, Guenard H, de Bisschop C, Martinot JB, Melot C, and Naeije
568 R. Pulmonary vascular reserve and exercise capacity at sea level and at high altitude. *High Alt*569 *Med Biol* 14: 19-26, 2013.

570 28. Reeves JT, Linehan JH, and Stenmark KR. Distensibility of the normal human lung
571 circulation during exercise. *Am J Physiol Lung Cell Mol Physiol* 288: L419-425, 2005.

572 29. Reeves RB, and Park HK. CO uptake kinetics of red cells and CO diffusing capacity.
573 *Respir Physiol* 88: 1-21, 1992.

30. Reuschlein PS, Reddan WG, Burpee J, Gee JB, and Rankin J. Effect of physical
training on the pulmonary diffusing capacity during submaximal work. *J Appl Physiol* 24: 152158, 1968.

31. Roughton FJ, and Forster RE. 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* 11: 290-302, 1957.

Sackner MA, Greeneltch D, Heiman MS, Epstein S, and Atkins N. Diffusing
capacity, membrane diffusing capacity, capillary blood volume, pulmonary tissue volume, and
cardiac output measured by a rebreathing technique. *Am Rev Respir Dis* 111: 157-165, 1975.

Stam H, Hrachovina V, Stijnen T, and Versprille A. Diffusing capacity dependent on
lung volume and age in normal subjects. *J Appl Physiol (1985)* 76: 2356-2363, 1994.

Tamhane RM, Johnson RL, Jr., and Hsia CC. Pulmonary membrane diffusing
capacity and capillary blood volume measured during exercise from nitric oxide uptake. *Chest*120: 1850-1856, 2001.

Tanaka H, Dinenno FA, Monahan KD, Clevenger CM, DeSouza CA, and Seals DR.
Aging, habitual exercise, and dynamic arterial compliance. *Circulation* 102: 1270-1275, 2000.

36. Taylor BJ, Carlson AR, Miller AD, and Johnson BD. Exercise-induced interstitial
pulmonary edema at sea-level in young and old healthy humans. *Respir Physiol Neurobiol* 191:
17-25, 2014.

Trappe SW, Costill DL, Vukovich MD, Jones J, and Melham T. Aging among elite
distance runners: a 22-yr longitudinal study. *J Appl Physiol (1985)* 80: 285-290, 1996.

597

Fig. 1 Group mean values of lung diffusing capacity for carbon monoxide (DLCO), alveolar-598 capillary membrane conductance (Dm_{CO}), and pulmonary-capillary blood volume (V_C) as a 599 function of cardiac output (Q) during exercise. DLCO, Dm_{CO}, and V_C increase relatively linearly 600 from rest to exercise at 90% W_{peak} in all groups. Error bars denote standard deviation from the 601 mean. Closed circles = Young Highly-Fit (HF); Open circles = Young; Closed squares = Old 602 Highly-Fit (HF); Open squares = Old. 603

604

Fig. 2 Individual values of lung diffusing capacity for carbon monoxide (DLCO), alveolar-605 capillary membrane conductance (Dm_{CO}), and pulmonary-capillary blood volume (V_C) as a 606 607 function of cardiac output (Q) during exercise. The mean and standard deviation of the slope of the lung diffusing capacity-cardiac output relationship is given for each group; there was no 608 significant difference in slope between groups for any measure. Closed circles = Young Highly-609 Fit (HF); Open circles = Young; Closed squares = Old Highly-Fit (HF); Open squares = Old. 610

611

Fig. 3 Individual values of \dot{VO}_{2max} as a function of resting values of lung diffusing capacity for 612 carbon monoxide (DLCO), alveolar-capillary membrane conductance (Dm_{CO}), and pulmonary-613 capillary blood volume (V_C). A linear regression was fit to all data points for each measure. 614 $\dot{V}O_{2max}$ is positively correlated with resting values of DLCO, Dm_{CO} , and V_{C} . Closed circles = 615 Young Highly-Fit (HF); Open circles = Young; Closed squares = Old Highly-Fit (HF); Open 616 squares = Old. 617

618

exercise at 90% of W_{peak} . A _x is positively correlated with to exercise at 90% of W_{peak} . Closed squares = Old Highlyardiac output slope for each data points for each measure. is not significantly correlated for carbon monoxide; Dm_{CO} , llary blood volume. Closed puares = Old Highly-Fit (HF);

Fig. 4 Individual values of $\dot{V}O_{2max}$ as a function of the absolute change (Δ) in values of lung diffusing capacity for carbon monoxide (DLCO), alveolar-capillary membrane conductance (Dm_{CO}), and pulmonary-capillary blood volume (V_C) from rest to exercise at 90% of W_{peak}. A linear regression was fit to all data points for each measure. $\dot{V}O_{2max}$ is positively correlated with the absolute change in values of DLCO, Dm_{CO}, and V_C from rest to exercise at 90% of W_{peak}. Closed circles = Young Highly-Fit (HF); Open circles = Young; Closed squares = Old Highly-Fit (HF); Open squares = Old.

626

Fig. 5 $\dot{V}O_{2max}$ as a function of the lung diffusing capacity-cardiac output slope for each individual subject (from Figure 2). A linear regression was fit to all data points for each measure. $\dot{V}O_{2max}$ is positively correlated with the DLCO- \dot{Q} slope; $\dot{V}O_{2max}$ is not significantly correlated with the Dm_{CO}- \dot{Q} or V_C- \dot{Q} slopes. DLCO, lung diffusing capacity for carbon monoxide; Dm_{CO}, alveolar-capillary membrane conductance; V_C, pulmonary-capillary blood volume. Closed circles = Young Highly-Fit (HF); Open circles = Young; Closed squares = Old Highly-Fit (HF); Open squares = Old.

634

Fig. 6 Group mean values of the ratio of lung diffusing capacity for nitric oxide to carbon monoxide (DLNO/DLCO) as a function of cardiac output (\dot{Q}) during exercise. DLNO/DLCO falls from rest to exercise at 90% W_{peak} in YNGm, OLDm, and OLDh. Error bars denote standard deviation from the mean. Closed circles = Young Highly-Fit (HF); Open circles = Young; Closed squares = Old Highly-Fit (HF); Open squares = Old.

640

641

642 **Table 1** Subject characteristics

	Young	Young Highly-Fit	Old	Old Highly-Fit	
Age, y	27 ± 3	27 ± 3	$69 \pm 5^{*}$ †	$65 \pm 5^{*}$ †	
N (female)	9 (0)	7 (0)	7 (1)	8 (1)	
Height, cm	178 ± 4	178 ± 4	173 ± 4	176 ± 7	
Mass, kg	78.0 ± 5.8	$72.5 \hspace{0.2cm} \pm \hspace{0.2cm} 6.8$	$73.6 \hspace{0.2cm} \pm \hspace{0.2cm} 8.8$	$75.4 \hspace{0.1in} \pm \hspace{0.1in} 9.0$	
BMI, kg/m ²	$24.7 \hspace{0.2cm} \pm \hspace{0.2cm} 2.0$	22.8 ± 2.1	$24.8 \hspace{0.2cm} \pm \hspace{0.2cm} 3.4$	$24.5 \hspace{0.2cm} \pm \hspace{0.2cm} 2.2$	
W _{peak} , W	$269 \ \pm \ 43$	$338 \pm 28*$	157 ± 26*†	239 ± 29†#	
VO₂max, ml/min	$3519 \ \pm \ 449$	$4577 \pm 419^*$	$2050 \pm 381*$ †	3140 ± 368 †#	
VO_{2max}, ml/kg/min	$45.3 \hspace{0.2cm} \pm \hspace{0.2cm} 6.5$	$63.4 \pm 6.0*$	$28.2 \pm 6.4*$ †	41.8 ± 3.8 †#	
% Pred. VO _{2max}	110 ± 18	$147 \pm 8*$	116 ± 13†	$162 \pm 18*\#$	
FEV ₁ /FVC	82 ± 6	$80~\pm~4$	78 ± 5	$74 \pm 5*$	
% pred.	99 ± 8	96 ± 5	105 ± 7	98 ± 5	
FEV ₁ , L	$4.8 \hspace{0.2cm} \pm \hspace{0.2cm} 0.4$	5.0 ± 0.8	$3.2 \pm 0.3*^{+}$	$3.7 \pm 0.9*$ †	
% pred.	106 ± 9	110 ± 13	110 ± 12	112 ± 21	
FVC, L	5.9 ± 0.7	6.3 ± 0.9	$4.2 \pm 0.4*^{+}$	5.0 ± 1.2 †	
% pred.	106 ± 10	114 ± 12	104 ± 11	114 ± 21	
FEF ₂₅₋₇₅ , L	$4.6 \hspace{0.2cm} \pm \hspace{0.2cm} 0.7$	4.5 ± 1.1	$2.9 \pm 0.7*$ †	$2.8 \pm 1.0^{*}$	
% pred.	100 ± 19	97 ± 21	126 ± 32	104 ± 30	
TLC, L	7.7 ± 0.9	8.0 ± 1.3	$6.9 \pm 1.0^{*}$ †	$8.2 \pm 1.2*$ †	
% pred.	112 ± 10	116 ± 16	106 ± 12	121 ± 11	

644 Values are reported as mean \pm SD. BMI, body mass index; W_{peak} , peak power output during maximal 645 exercise test; \dot{VO}_{2max} , maximal oxygen consumption; FEV_1 , forced expiratory volume in 1 s; FVC, forced 646 vital capacity; FEF_{25-75} , average forced expiratory volume during middle portion of FVC; TLC, total lung 647 capacity. Significance set at P < 0.05; *significantly different vs. Young, [†]significantly different vs. 648 Young Highly-Fit, [#]significantly different vs. Old.

649**Table 2** Lung diffusing capacity and cardiac output

650

	Young	Young Highly-Fit	Old	Old Highly-Fit	Young	Young Highly-Fit	Old	Old Highly-Fit
	Rest				<i>Exercise at 90% of</i> W_{peak}			
DLCO, ml/min/mmHg	28 ± 4	33 ± 4	$17 \pm 3*;$	$23 \pm 5^{\dagger}$	$47 \hspace{0.1in} \pm \hspace{0.1in} 6$	53 ± 7	25 ± 3*†	$37 \pm 10^{*}$ †#
DLNO, ml/min/mmHg	101 ± 22	112 ± 16	$67 \pm 10^{*}$	$85 \pm 18^{\dagger}$	153 ± 16	171 ± 18	87 ± 7*†	$122 \pm 30^{*}$ †#
DLNO/DLCO, unitless	3.6 ± 0.4	3.4 ± 0.4	$4.0 \hspace{0.1in} \pm \hspace{0.1in} 0.6$	3.8 ± 0.4	3.3 ± 0.3	3.2 ± 0.2	3.6 ± 0.2	3.3 ± 0.2
Dm _{CO} , ml/min/mmHg	$45 \hspace{0.1in} \pm \hspace{0.1in} 10$	50 ± 7	$30 \pm 4^{*}$ †	$37 \pm 8^{\dagger}$	71 ± 12	$77 \hspace{.1in} \pm \hspace{.1in} 10$	$39 \pm 3^{*}$ †	54 ± 13*†#
V _C , ml	82 ± 20	$95~\pm~14$	44 ± 9*†	68 ± 20 †	127 ± 30	$149 \hspace{.1in} \pm \hspace{.1in} 15$	$66 \pm 10^{*}$ †	110 ± 29†#
Q, L/min	5.6 ± 2.0	6.2 ± 1.1	$3.5 \pm 0.6*$ †	5.1 ± 1.0	17.3 ± 3.1	$20.3 \hspace{0.2cm} \pm \hspace{0.2cm} 2.2$	$10.3 \pm 1.6*$ †	14.7 ± 3.5†#

651Values are reported as mean ± SD. DLCO, lung diffusing capacity for carbon monoxide; DLNO, lung diffusing capacity for nitric oxide; 652Dm_{CO}, alveolar-capillary membrane conductance; V_C, pulmonary-capillary blood volume; Q, cardiac output. Significance set at P < 0.05; 653^{*}significantly different vs. Young, [†]significantly different vs. Young Highly-Fit, [#]significantly different vs. Old. Downloaded from http://jap.physiology.org/ by 10.220.33.5 on March 30, 2017











