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## Near-infrared spectroscopy using indocyanine green dye for minimally invasive measurement of respiratory and leg muscle blood flow in patients with COPD

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Running Head: Validation of NIRS-ICG derived BFI in COPD

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#### Abstract

Near-infrared spectroscopy (NIRS), measuring indocyanine green (ICG) after its peripheral venous injection, has been validated for minimally invasive measurement of relative muscle blood flow in fit young individuals, but not in COPD. Here we ask whether it could be used to evaluate respiratory and locomotor muscle perfusion in COPD patients. Vastus lateralis muscle blood flow (MBF, the reference method calculated from arterial and muscle ICG concentration curves) and a blood flow index (BFI, calculated using only the (same) muscle ICG concentration curves) were compared in 10 patients (FEV<sub>1</sub>:51±6% predicted) at rest and during cycling at 25%, 50%, 75% and 100% of WRpeak. Intercostal muscle MBF and BFI were also compared during isocapnic hyperpnea at rest, reproducing ventilation levels up to those at WRpeak. Intercostal and vastus lateralis BFI increased with increasing ventilation during hyperpnea (from  $2.5\pm0.3$  to  $4.5\pm0.7$  nM/s) and cycling load (from  $1.0\pm0.2$  to  $12.8\pm1.9$  nM/s), respectively. There were strong correlations between BFI and MBF for both intercostal (r=0.993 group mean data, r=0.872 individual data) and vastus lateralis (r=0.994 group mean data, r=0.895 individual data). Fold changes from rest in BFI and MBF did not differ for either the intercostal muscles or the vastus lateralis. Group mean BFI data showed strong interrelationships with respiratory and cycling workload, and whole body metabolic demand (r ranged from 0.913 to 0.989) simultaneously recorded during exercise. We conclude that BFI is a valid and minimally invasive tool for evaluating relative changes in respiratory and locomotor muscle perfusion from rest to peak exercise in COPD patient groups. 

34	News and Noteworthy
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36	We show that non-invasive near-infrared spectroscopic (NIRS) detection of indocyanine green
37	dye (ICG) after peripheral venous injection adequately reflects respiratory and locomotor
38	muscle perfusion during exercise and hyperpnea in COPD patients. Mean, individual, and fold-
39	change responses from rest to exercise or hyperpnea correlated closely with the reference
40	method, which requires arterial sampling. NIRS-ICG is a valid, robust and essentially non-
41	invasive tool for assessing relative changes in respiratory and locomotor muscle perfusion in
42	COPD patient groups.
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79	Keywords:	NIRS,	Indocyanine	Green d	lye,	muscle	perfusion,	respiratory	muscles,	exercise,
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- 80 COPD, validation, BFI

Introduction 120 121 In patients with Chronic Obstructive Pulmonary Disease (COPD) restrictions in 122 central hemodynamics owing to abnormal lung mechanics (18) can limit perfusion to the 123 working muscles thereby exacerbating the competition between respiratory and locomotor 124 125 muscles for the available blood flow (1, 5, 13). Measurement of muscle blood flow (MBF) in patients with COPD therefore provides an important tool for investigating the 126 pathophysiological mechanisms involved in exercise intolerance and for assessing the 127 effectiveness of pharmacological or non-pharmacological interventions. 128

129 Methods for measuring MBF have been developed over the past 120 years (2). 130 However, the traditional techniques for assessing MBF are highly invasive exposing the individuals to risks and posing difficulties to differentiate working and non working muscles 131 (3). For example, by using NIRS after an intravenous injection of the light-absorbing tracer 132 133 indocyanine green (ICG), it is possible to measure absolute values of MBF by applying the law of conservation of mass (3, 6). Specifically, the rate of accumulation of a tracer in a 134 given tissue is equal to its rate of inflow minus its rate of outflow. If a tracer is introduced 135 136 rapidly and its rate of accumulation is measured over time, blood flow can be measured as a ratio of the tracer accumulated to the quantity of tracer introduced over a given time. 137 However, to measure the quantity introduced requires continuous arterial blood sampling 138 over the time of tracer introduction, and thus the need for an indwelling arterial cannula. 139

Recent studies indicate that in young, fit subjects, relative values of muscle blood flow can be determined by NIRS after intravenous ICG injection, measuring only the accumulation rate of tracer without arterial sampling (7, 8). The result, termed the blood flow index (BFI), was found to reliably reflect local (i.e., within the sampling volume of each NIRS optode) changes in both respiratory and locomotor MBF from rest to maximal ventilation (i.e., ~120 liters/minute) during (isocapnic) hyperpnea or during graded cycling exercise up to maximal levels (as high as 360 watts) (7, 8). However, the validity of this method has not been investigated in a clinical population. Indeed, due to hemodynamic and
muscular differences between healthy subjects and COPD patients (11, 18), the results
obtained in healthy, fit subjects may not be transferrable to this patient population.

To this end, we retrospectively analyzed data obtained in a representative population of COPD patients across a wide range of exercise intensities and rates of minute ventilation during cycling and subsequently during isocapnic hyperpnea (16). Accordingly, the purpose of the present study was to examine in COPD patients whether NIRS-ICG derived BFI reflects changes in respiratory and locomotor muscle perfusions as reliably as simultaneously determined MBF calculated by Fick principle based on the law of conservation of mass.

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#### Materials and methods

158 Subjects

159 We retrospectively analyzed data from ten clinically stable patients with COPD (FEV<sub>1</sub>:51±6% predicted) classified by the Global Initiative for Chronic Obstructive Lung 160 Disease (GOLD, 2016) as stages II (n=4), III (n=3), and IV (n=3) from our previously 161 published work (16). In the original study (16), vastus lateralis and intercostal muscle blood 162 flow (MBF) were simultaneously measured in patients with COPD with the aim to 163 investigate whether during exercise, intercostal MBF competes with locomotor muscles for 164 165 the available blood flow (16). In the original study, only MBF data were analyzed and presented. In the present analysis, intercostal and vastus lateralis MBF are compared with 166 simultaneously measured blood flow index (BFI) in the same locations. Measurements were 167 made under two separate conditions: from rest to near-peak cycle exercise, and from resting 168 to peak (isocapnic) hyperpnea. 169

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#### 172 Study Design

The exercise protocols and isocapnic hyperpnea tests during which MBF and BFI were compared, as well as the methods used for assessing the physiologically relevant variables presented in this study, have been previously described (16), and are thus not repeated here.

176 Intercostal and quadriceps MBF and BFI

Absolute values of intercostal and vastus lateralis MBF during exercise were measured based on the principle of conservation of mass using an equation developed by (3). In brief, for both intercostal and vastus lateralis muscle, ICG concentration difference (measured by NIRS) at time t ([ICG]<sub>m</sub>) during dye accumulation in muscle tissue (between 10 and 70% of peak concentration) was divided by the integral under the arterial ICG concentration ([ICG]<sub>a</sub>) curve (measured by linear photodensitometer) until time t and multiplied by t and a factor k for unit conversion as previously described (16):

184 Blood flow (ml/100ml/min<sup>-1</sup>) = 
$$\frac{k \cdot [ICG]_m \cdot t}{\int_{a}^{t} [ICG]_a dt}$$

Relative perfusion (BFI) of intercostal and vastus lateralis muscles during exercise 185 186 was also obtained. This was done by dividing the NIRS-derived muscle ICG concentration difference (peak height of the ICG concentration curve) by the rise time from 10 to 90% of 187 peak as previously described (7-9). NIRS-derived muscle ICG concentration was expressed 188 189 in nanomoles per liter (nM). After dividing by the rise time (in seconds) the final unit of measurements of BFI was nM/s (7-9). Representative examples for calculating BFI for 190 intercostal and vastus lateralis muscle from one individual are shown in Figure 1a and 191 192 Figure 1b, respectively. ICG concentration curves were analyzed by using Chart5 v.5.4.2 (ADInstruments) program (8). NIRS data were sampled at 6 Hz and time was synchronized 193 to the metabolic, ventilatory, mechanical work and respiratory pressure data (16). 194

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#### 197 Statistical analysis

Data are presented as means  $\pm$  SEM. Pearson's correlation coefficient (r) was used to 198 199 establish associations between intercostal and vastus lateralis muscle BFI with MBF and 200 other physiologically related variables. ANOVA with repeated measures and post-hoc 201 comparisons (Tukey's test) were used to identify statistically significant differences across 202 the cycling exercise and isocapnic hyperpnea trials. Analysis of agreement between BFI and MBF measurements was performed by using Bland-Altman analysis. A satisfactory 203 204 agreement between BFI and MBF was considered when the difference between BFI and MBF measurements did not significantly vary from 0 (zero). For this purpose one sample t-205 test with testing variable the difference between BFI and MBF measurements and testing 206 207 value 0 (zero) was performed. For Bland-Altman analysis purposes, both BFI in nM/s and MBF in ml/min/100g were expressed as fold changes from rest in order to allow a 208 comparison between the two methods. The level of statistical significance was set at P < P209 210 0.05. All statistical analyses were performed using the SPSS statistical software (v. 20 IBM 211 SPSS Statistics, Chicago, IL, USA).

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#### Results

### 214 **Patient characteristics**

Detailed subject characteristics in terms of demographics, pulmonary function, and peak exercise performance data have been published and discussed previously (16). Not reported previously, patients exhibited impaired functional capacity in terms of the six minute walking distance test (389±31meters, 59±8% predicted), quadriceps muscle strength (28±5kg, 61±6% predicted), and quadriceps muscle endurance, (43±8 seconds 54±5% predicted, respectively).

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#### Perfusion responses to exercise

During the resting isocapnic hyperpnea trial, both intercostal muscle blood flow 224 225 (MBF) and blood flow index (BFI) increased from: 7.0±0.6 to 11.4±1.3 ml/min/100gr 226 (MBF) and from 2.5±0.3 to 4.5±0.7 nM/s, (BFI). Moreover, both MBF and BFI reached a 227 plateau at 75% of peak minute ventilation (Figure 2a). During the cycling trial, both vastus lateralis MBF and BFI increased progressively with increasing power output: from 2.8±0.4 228 to 39.7±7.0 ml/min/100gr (MBF) and from 1.0±0.2 to 12.8±1.9 nM/s, (BFI) up to 100% of 229 230 WRpeak (Figure 2c). When MBF and BFI data were expressed as changes relative to flows at rest, the pattern of increase in muscle perfusion with cycle exercise or with resting 231 hyperpnea did not significantly differ (between MBF and BFI) either for the intercostal or 232 the vastus lateralis muscle groups (Figures 2b and 2d) 233

#### 234 Comparison of BFI and MBF

During isocapnic hyperpnea, there was a highly significant and close correlation (r=0.993, P=0.001) between mean values of intercostal muscle BFI and MBF (Figure 3a). Furthermore, the individual values of intercostal muscle BFI and MBF showed a close linear relationship (r=0.872, P<0.001, Figure 3b). Normalization of BFI for body weight (BW), body mass index (BMI) and fat free mass index (FFMI) did not meaningfully change the correlation with MBF (r= 0.829, 0.856 and 0.837, P<0.001, respectively).

241 During cycling exercise, the correlation between mean vastus lateralis muscle BFI and MBF was also close and highly significant (r=0.994, P=0.001, Figure 3c). Again, correlation 242 analysis of individual patient vastus lateralis muscle BFI and MBF values showed close 243 244 association between the two methods (r=0.895, P<0.001, Figure 3d). Normalization of BFI 245 for BW, BMI and FFMI again did not meaningfully change the individual correlations with MBF (r=0.798, 0.843 and 0.809, P<0.001, respectively). Individual relationships between 246 intercostal and vastus lateralis muscle BFI with intercostal and vastus lateralis MBF are 247 shown in Figures 4 and 5, respectively. In addition, individual regression line slopes of the 248

individual relationships between intercostal and vastus lateralis muscle BFI and MBF areshown in Figure 6.

#### 251 Agreement between BFI and MBF

252 Figure 7 shows the results of Bland-Altman analysis between MBF and BFI data 253 expressed as fold change from rest for intercostal (Figure 7a) and vastus lateralis muscle groups (Figure 7b). Specifically, for both muscle groups there was a satisfactory agreement 254 255 between MBF and BFI as the difference between the two methods in fold change from rest 256 in muscle perfusion did not vary statistically significantly from 0 (zero) (intercostal muscles: mean difference: -0.05 fold change from rest, 95%CI: 0.07 to -0.17, p=0.424 and vastus 257 lateralis: mean difference: -1.51 fold change from rest, 95%CI: -4.2 to 1.2, p=0.268, 258 259 respectively).

#### 260 Associations between BFI and physiologically relevant variables

During the isocapnic hyperpnea trial the mean change from rest in intercostal muscle 261 BFI was linear with respect to the mean changes in whole body oxygen uptake (r=0.913, 262 263 P=0.030), minute ventilation (r=0.954, P=0.012), the power of breathing (r=0.922, P=0.026), 264 and tidal excursion in transdiaphragmatic pressure (r=0.962, P=0.009) (Figure 8). Similarly, the mean change from rest in vastus lateralis muscle BFI was linear with respect to mean 265 change in whole body oxygen uptake (r=0.989, P=0.001, and mean cycling load (r=0.984, 266 267 P=0.002) during graded cycling exercise (Figure 9). Individual r values for intercostal and vastus lateralis muscle between BFI and the aforementioned physiologically variables are 268 shown in Table 1. Specifically, for the majority of patients, BFI increased linearly with 269 270 physiological relevant variables, but the slope of the regression line varied considerably 271 between subjects.

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#### Discussion

#### 276 Main findings

277 This study examined the validity of the essentially non-invasive NIRS-derived blood 278 flow index (BFI) in reflecting respiratory and quadriceps muscle blood flow (MBF, the 279 reference method) in a representative COPD population over a wide range of ventilation and exercise workloads. Under all conditions examined, individual, mean and fold change 280 responses of BFI tracked MBF closely (Figures 2-5 and 7). Furthermore, while the slope of 281 282 individual patient relationships between BFI and MBF may differ (Figure 6) individual correlations between the two methods for both intercostal and vastus lateralis muscle groups 283 remained highly significant (Figure 3b and d and Figures 4 and 5). In addition, relative mean 284 and individual changes in intercostal and vastus lateralis muscle BFI from rest to peak 285 exercise or peak minute ventilation were strongly associated with measures of respiratory 286 load, mechanical workload, and whole body metabolic demand across a wide range of 287 physiologically relevant rates of minute ventilation and exercise intensities (Figures 8-9 and 288 Table 1). 289

The findings of the present study extend previous observations in young, fit individuals (6, 7) by demonstrating that BFI is able to detect changes in respiratory and leg MBF during exercise over a range of ventilation rates and exercise workloads typically experienced by patients with COPD. However, the BFI method does not allow absolute blood flow to be determined – it provides relative flow rates between conditions, but with the major advantage of avoiding arterial catheterization.

296 Comparison of MBF to BFI

This study provides strong support for the use of BFI in place of MBF as the data show. A difficulty in their comparison however is their very different units as explained in the Methods section. To compare side by side, we chose to calculate fold changes (from rest) in MBF and BFI, and these also indicated close similarity as shown (Figure 2b and

Figure 2d). However, fold changes are very sensitive to small variations in the resting values 301 (which form the denominator for fold changes). This explains the behavior noted in the 302 303 Bland-Altman plots (Figure 7) of some points with huge differences between the two 304 methods - the points identified with asterisks towards the right side of Figure 7a and 7b. 305 Specifically, for intercostal muscles (Figure 7a) one point lies at  $\sim +1$ -fold difference and two points at ~ -1 -fold difference whist for vastus lateralis muscle (Figure 7b) two points lie 306 307 at ~ +20-fold difference and three at ~ -20 -fold difference, which seem to indicate very high 308 variance. In Figures 4 and 5 we identify the same eight points, and it is apparent that the flows on exercise fit very well with the regression line in each case, reinforcing the 309 sensitivity of fold-change to the resting data. The Bland-Altman plot therefore distorts the 310 311 comparison of the MBF and BFI data because of the reliance on fold changes as the 312 comparison variable. However, that plot remains useful as an indicator that mean values 313 were not different from zero over the range encountered.

### 314 Advantages of BFI for measuring muscle blood flow in COPD.

BFI measures muscle blood flow by dividing the muscle ICG peak concentration 315 316 (assessed noninvasively by NIRS-ICG curve) by the rise time from 10 to 90% of peak (Figure 1). The use of these two cutoff points eliminates the need for exact temporal 317 definition of the start and end of the ICG washin, which are somewhat observer-dependent 318 319 (7). The only invasive component of this technique - compared to traditional methods for assessing MBF - is peripheral venous bolus injection of the ICG tracer. BFI requires only 320 the determination of the muscle ICG peak concentration and the times at 10% and 90% of 321 322 the rise in ICG concentration (Figure 1) thus reducing the possibility of observer errors. 323 Indeed, the study by Habazettl et al., (8) in normal subjects reported that the reproducibility of BFI is high as there is less inter-observer variability for analyses of intercostal and 324 quadriceps muscle BFI compared to the variability observed during MBF calculation 325 measured by NIRS-ICG relying on the Fick principle. Taking into consideration the 326

327 practical and methodological advantages of BFI as well as the findings of the present study 328 showing that mean and individual values of BFI and MBF were strongly associated over a 329 wide range of ventilatory rates and exercise intensities (Figures 2-5) we suggest that BFI can 330 be considered a reliable measure of relative blood flow changes from rest to exercise in 331 patients with COPD.

#### 332 Prior studies of the validation of the NIRS-ICG derived BFI

In clinical populations NIRS-ICG derived BFI has been shown to be sensitive and 333 334 reproducible in detecting relative perfusion differences in cerebral blood flow during bedside assessment in patients with acute ischemic stroke (9, 14, 19). Furthermore, a 335 validation study in young, fit subjects by Habazettl et al. (8) compared BFI values within the 336 vastus lateralis and 7<sup>th</sup> intercostal space against NIRS-ICG derived absolute MBF during 337 cycling exercise up to a maximal level (~360 watt). The results indicated a strong group 338 mean and individual agreement between BFI and MBF measured across both respiratory and 339 quadriceps muscles. In addition, the study by Guenette et al. (7) extends the observation by 340 Habazettl et al. (8) by investigating the sensitivity of BFI in intercostal and 341 342 sternocleidomastoid muscles across a wide range of ventilatory rates up to maximal levels (i.e., ~120 liters/min) during an isocapnic hyperpnea trial with simultaneous measurements 343 of electromyography and the work of breathing produced by these muscles. The results of 344 345 that study showed a strong correlation with the work of breathing and electromyography (EMG) data for both aforementioned respiratory muscles (7). 346

#### 347 Relationship between BFI and other physiological variables

In addition to the data comparing BFI to MBF, BFI was related to a number of physiologically relevant variables that were simultaneously recorded during the two protocols (i.e., cycling exercise and isocapnic hyperpnea trials). The strong group mean and individual associations that we found between BFI and those variables (Figures 8-9, Table 1) further support the physiological value of BFI as an index proportional to exercise-related

variables. In addition, these associations are also useful because considering the 353 comparisons between BFI and MBF, one could argue that a bias may occur owing to the 354 355 mathematical coupling between BFI and MBF variables (12, 20). This is because calculation 356 of both BFI and MBF use the same muscle ICG-concentration curve. Therefore, relating BFI 357 to independent relevant variables provides additional support for its validity. There were strong linear relationships between intercostal and vastus lateralis BFI and metabolic 358 requirement (VO<sub>2</sub>), ventilation, and mechanical loading (Figures 8-9 and Table 1) -359 360 relationships which avoid any mathematical coupling.

#### 361 **BFI** application to individual patients and to groups of patients

The data show excellent reliability when average values for groups are to be compared 362 (Figures 2, 3a and b). However it is more difficult to know whether BFI can be used to 363 estimate MBF on a single patient basis. This is because while BFI is an index proportional to 364 MBF, the slope of the BFI/MBF relationship varies across individuals (Figure 6). Therefore, 365 one cannot convert BFI to actual MBF in all patients by a single assumed average 366 proportionality constant (Figure 6). Figure 4 (Intercostals, hyperpnea trial), Figure 5 367 368 (Ouadriceps, cycling trial), and Figure 6 address this point by plotting for each of the 10 patients the individual relationships (Figures 4-5) and the regression line slopes (Figure 6) 369 between BFI and MBF. While in each case the relationships are closely linear for the 370 371 intercostal muscles, for two patients (i.e., subjects 6 and 7, Figure 4) the slopes between BFI and MBF relationship differ considerably from the other 8 subjects (Figure 6a). It is of 372 interest that for the same two subjects, the correlations between BFI and other 373 374 physiologically relevant variables are also found to be weak (Table 1), suggesting a physiological rather than technical explanation. For the quadriceps (Figure 6b) the 375 regression line slopes varied less as compared to intercostal muscles. In summary, 376 intercostal and quadriceps BFI cannot be used for conversion to MBF in all individual 377

subjects under different conditions, although use of an average proportionality constantwould be satisfactory for the majority of patients.

#### 380 Methodological considerations

381 The present study assessed respiratory muscle BFI in COPD patients at the level of the 7<sup>th</sup> intercostal space as this has been previously proposed for measuring absolute MBF by 382 studies using the NIRS-ICG approach (6, 7, 10, 15-17). This site of measurement (i.e., 7th 383 intercostal space), while easily accessible by NIRS, reflects both internal and external 384 385 intercostals muscles, thus providing an overall assessment of respiratory MBF. However, it does not reflect diaphragm blood flow. Nevertheless, in COPD patients, as the degree of 386 dynamic lung hyperinflation gradually increases during exercise, the pressure generated by 387 the diaphragm decreases and the act of inspiration is more dependent on the rib cage 388 389 inspiratory muscles such as the intercostal muscles (4). Indeed, in our study we did not find any impact of dynamic lung hyperinflation on intercostal muscle BFI responses as both 390 progressively increased, leveling off at 75% of WRpeak during isocapnic hyperpnea (actual 391 392 increase in end-expiratory lung volume at 25, 50, 75, 100% of WRpeak was 60±90, 240±50,  $550\pm110$ ,  $480\pm180$  ml, respectively). This further supports the use of 7<sup>th</sup> intercostal space for 393 assessing respiratory MBF in clinical populations irrespective of the occurrence of exercise-394 395 induced dynamic hyperinflation.

396 We found stronger associations between individual values of BFI and MBF (Figures 3b and d) in both respiratory (r=0.872 and locomotor muscles (r=0.895) compared to those 397 reported in young, fit subjects by Habazettl et al. (8) (i.e., r=0.730 and r=0.720, 398 399 respectively). In fact, the study by Habazettl et al. (8) reported substantial scattering of 400 individual data of BFI and MBF mostly exhibited during high levels of exercise. In our study, scattering of individual data of BFI and MBF appears much less with increasing 401 exercise intensity (Figures 3b and d). This may be attributed to the lower peak ICG values 402 that COPD patients exhibited compared to healthy subjects (both in muscle and arterial 403

blood) - as response to lower exercise intensity - and thus making easier to calculate the
peak ICG concentrations. Indeed, determination of peak ICG concentrations may be a source
of random errors that may contribute to the variability of individual BFI vs. MBF values (9),
especially when oscillations of the ICG concentration curve due to strong muscle contraction
dynamics during forced ventilation or high workloads require smoothing of the original ICG
concentration curve (for example see figure 1).

410 Another methodological consideration is the normalization of the BFI values for body 411 mass/composition when major variability in body mass (BM) and/or composition exists (Habazettl et al., 2010). In our study we did not normalize as we were most interested in 412 BFI/MBF relationships within each patient. However, we did analyze normalization of both 413 intercostal and vastus lateralis muscle BFI to BM, BMI and fat-free mass index (FFMI). 414 Although we found that BM, BMI and FFMI varied considerably among patients (i.e., BM: 415 from 52 kg to 110 kg, BMI: from 18.4 to 34.7 kg/m<sup>2</sup> and FFMI: from 17.5 to 23.1 kg/m<sup>2</sup>), 416 this did not affect the level of association between the BFI and MBF in both muscle groups. 417

#### 418 Conclusions

419 In a small population of patients with COPD, the essentially non-invasively derived BFI showed close agreement with MBF (determined from the same signals combined with 420 arterial ICG concentration/time curves) when applied to group data. This was found for 421 422 changes in intercostal and locomotor MBF across a wide range of ventilatory rates and exercise workloads up to peak levels. The findings support the quantitative validity of this 423 method, thereby allowing for minimally invasive assessment of relative muscle blood flow 424 425 in this population. While individual patient values of BFI and MBF were also generally 426 similar in both muscle groups, occasional patients displayed variances which give rise to caution in application to individual patients. 427

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	Intercostal muscles (hyperpnea trial)				Quadriceps muscle (cycling trial)		
Patients	BFI vs.VO <sub>2</sub>	BFI vs.V <sub>E</sub>	BFI vs. PoB	BFI vs. ΔPdi	BFI vs.VO <sub>2</sub>	BFI vs. Work	
1	0.909	0.953	0.962	0.961	0.937	0.958	
2	0.900	0.973	0.955	0.953	0.969	0.912	
3	0.915	0.949	0.891	0.893	0.895	0.937	
4	0.893	0.895	0.906	0.915	0.921	0.938	
5	0.710	0.713	0.568	0.559	0.698	0.541	
6	0.260	0.273	0.330	0.313	0.829	0.729	
7	0.686	0.493	0.289	0.328	0.903	0.927	
8	0.907	0.901	0.904	0.931	0.890	0.990	
9	0.925	0.942	0.958	0.959	0.989	0.970	
10	0.852	0.789	0.796	0.836	0.875	0.938	
565					·		

Table 1. Regression analysis for individual subjects between BFI and physiologically relevant variables

Individual Pearson r values relating intercostal muscles blood flow index (BFI) oxygen uptake (VO<sub>2</sub>), minute ventilation (V<sub>E</sub>), power of breathing (PoB) and transdiaphragmatic pressure ( $\Delta$ Pdi) recorded from rest to peak minute ventilation during the isocapnic hyperpnea trial and quadriceps muscle blood flow index (BFI) with quadriceps muscle blood flow (MBF), oxygen uptake (VO<sub>2</sub>) and work (watts) recorded from rest to WRpeak during cycling exercise.

591	Figure legends							
592 593	Figure 1. Representative examples of an intercostal (a) and quadriceps (b) muscle							
594	indocyanine green (ICG) concentration curve recorded by near-infrared spectroscopy							
595	(NIRS) during isocapnic hyperpnea and cycling trials at 75% of WRpeak in an individual							
596	subject. Isocapnic hyperpnea was sustained at the level of minute ventilation (i.e., 40 l/min)							
597	similar to that recorded during cycling exercise at 75% of WRpeak (i.e., 58 watts). The							
598	original tracing (gray line) appears with marked oscillations owing to muscle contraction							
599	and relaxation during exercise. Low-pass filtering with a cutoff frequency of 0.5 Hz							
600	produced the smoothed curve (black line) that was used for blood flow index (BFI)							
601	calculation. ICG concentrations expressed in nanomoles/liter (nM) and the rise time							
602	expressed in seconds between 10% and 90% of ICG concentration peak are indicated, and							
603	intercostal and quadriceps muscle BFI calculations and results are shown.							
604								

**Figure 2.** Group mean responses of intercostal (a) and quadriceps (c) muscle blood flow (MBF) and blood flow index (BFI) during isocapnic hyperpnea and cycling trials respectively. Relative changes from rest of intercostal (b) and quadriceps (d) muscle blood flow (MBF) and blood flow index (BFI) are also shown. Data are presented as mean  $\pm$  SEM. Asterisks denote significant differences from 100% of WRpeak, P<0.05. (MBF data were reproduced from reference 16).

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**Figure 3.** Comparison of mean and individual blood flow index (BFI) and muscle blood flow (MBF) values for intercostal (a) and (b) and quadriceps (c) and (d) muscles during isocapnic hyperpnea and cycling trials, respectively. Linear regression equations, regression coefficients, and significance levels are presented in each figure. (MBF data were reproduced from reference 16).

Figure 4. Individual patient correlations between intercostal muscle blood flow index (BFI)
and intercostal muscle blood flow (MBF). Linear regression equations and regression
coefficients are presented in each figure. (MBF data were reproduced from reference 16).

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Figure 5. Individual correlations between vastus lateralis muscle blood flow index (BFI)
with vastus lateralis muscle blood flow (MBF). Linear regression equations and regression
coefficients are presented in each figure. (MBF data were reproduced from reference 16).

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**Figure 6.** Individual regression line slopes between (a) intercostal muscle blood flow index (BFI) and intercostal muscle blood flow (MBF) and (b) vastus lateralis muscle blood flow index (BFI) with vastus lateralis muscle blood flow (MBF). Mean regression line slope and equation is presented in each figure (MBF data were reproduced from reference 16).

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Figure 7. Bland-Altman plots comparing fold changes form rest of intercostal (a) and
quadriceps (b) muscle blood flow (MBF) and blood flow index (BFI) during isocapnic
hyperpnea and cycling trials, respectively (MBF data were reproduced from reference 16).

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**Figure 8.** Regression analyses of mean intercostal muscle blood flow index (BFI) and (a) oxygen uptake, (b) minute ventilation, (c) power of breathing, and (d) tidal excursion in transdiaphragmatic pressure during the isocapnic hyperpnea trial. Regression coefficients and significance levels are presented in each figure. (physiologically relevant data were reproduced from reference 16).

640

Figure 9. Regression analyses of mean quadriceps muscle blood flow index (BFI) and (a)oxygen uptake and (b) work rate recorded at rest and during the cycling trial. Regression

- 643 coefficients and significance levels are presented in each figure. (physiologically relevant
- 644 data were reproduced from reference 16).















