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**Bailey, TG, Perissiou, M, Windsor, M, Russell, FD, Golledge, J, Green, DJ and Askew, CD**

**Cardiorespiratory Fitness Modulates The Acute Flow-Mediated Dilation Response Following High-Intensity But Not Moderate-Intensity Exercise In Elderly Men.**

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### Article

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2  
3 **CARDIORESPIRATORY FITNESS MODULATES THE ACUTE**  
4 **FLOW-MEDIATED DILATION RESPONSE FOLLOWING**  
5 **HIGH-INTENSITY BUT NOT MODERATE-INTENSITY**  
6 **EXERCISE IN THE ELDERLY**

7  
8 TOM G. BAILEY<sup>1</sup>  
9 MARIA PERISSIOU<sup>1</sup>  
10 MARK WINDSOR<sup>1</sup>  
11 FRASER RUSSELL<sup>1,4</sup>  
12 JONATHAN GOLLEDGE<sup>2</sup>  
13 DANIEL J. GREEN<sup>3,4</sup>  
14 CHRISTOPHER D. ASKEW<sup>1</sup>

15  
16 <sup>1</sup>*VasoActive Research Group, School of Health and Sport Sciences,*  
17 *University of the Sunshine Coast, Queensland, Australia.*

18 <sup>2</sup>*Queensland Research Centre for Peripheral Vascular Disease,*  
19 *James Cook University and the Townsville Hospital*

20 <sup>3</sup>*School of Sport Science, Exercise and Health, The University of Western Australia*

21 <sup>4</sup>*Research Institute for Sport and Exercise Sciences, Liverpool John Moores University*  
22  
23

24 **SHORT TITLE:** Exercise intensity and FMD in elderly males

25  
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28  
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34 **AUTHOR FOR CORRESPONDENCE:**

35 \*Tom Bailey, PhD. *VasoActive Research Group* – School of Health and Sport Sciences,  
36 University of the Sunshine Coast, Locked bag 4, Maroochydore DC, Australia. Email  
37 [tbailey@usc.edu.au](mailto:tbailey@usc.edu.au)  
38

39 **ABSTRACT**

40 Impaired endothelial function is observed with ageing and with low cardiorespiratory fitness  
41 ( $VO_{2peak}$ ) whilst improvements in both are suggested to be reliant on higher-intensity exercise in  
42 the elderly. This may be due to the flow-mediated dilation (FMD) response to acute exercise of  
43 varying intensity. We examined the hypothesis that exercise-intensity alters the FMD response in  
44 healthy elderly adults, and would be modulated by  $VO_{2peak}$ . Forty-seven elderly men were  
45 stratified into lower- ( $VO_{2peak} = 24.3 \pm 2.9 \text{ ml.kg}^{-1}.\text{min}^{-1}$ ,  $n=27$ ) and higher-fit groups ( $VO_{2peak} =$   
46  $35.4 \pm 5.5 \text{ ml.kg}^{-1}.\text{min}^{-1}$ ,  $n=20$ ) after a test of cycling peak power output (PPO). In randomised  
47 order, participants undertook 27 min moderate-intensity continuous (MICE; 40% PPO) or high-  
48 intensity interval cycling exercise (HIIE; 70% PPO), or no-exercise control. Brachial FMD was  
49 assessed at rest, 10 and 60 min after exercise. In control, FMD reduced in both groups ( $P=0.05$ ).  
50 FMD increased after MICE in both groups [increase of 0.86 % (95% CI, 0.17 to 1.56),  $P=0.01$ ],  
51 and normalised after 60 min. In the lower-fit, FMD reduced after HIIE [reduction of 0.85 %  
52 (95% CI, 0.12 to 1.58),  $P=0.02$ ], and remained decreased at 60 min ( $P=0.05$ ). In the higher-fit  
53 FMD was unchanged immediately after HIIE and increased after 60 min [increase of 1.52 %  
54 (95% CI, 0.41 to 2.62),  $P<0.01$ ], which was correlated with  $VO_{2peak}$  ( $r = 0.41$ ;  $P<0.01$ ). Exercise-  
55 intensity alters the FMD response in elderly adults, and  $VO_{2peak}$  modulates the FMD response  
56 following HIIE, but not MICE. The sustained decrease in FMD in the lower-fit may represent a  
57 signal for vascular adaptation or endothelial fatigue.

58

59 **Key Words:** exercise, endothelial function, FMD, ageing, cardiorespiratory fitness

60

61 **New and noteworthy**

62

63 This study is the first to show that moderate-intensity continuous exercise increased FMD

64 transiently before normalisation of FMD after one hour, irrespective of cardiorespiratory fitness

65 level in the elderly. Interestingly, we show increased FMD after high-intensity exercise in

66 higher-fit, with a sustained reduction in FMD in lower-fit. The prolonged reduction in FMD after

67 high-intensity exercise may be associated to future vascular adaptation but may also reflect a

68 period of increased cardiovascular risk in lower-fit elderly.

69

70 **INTRODUCTION**

71 Ageing is associated with chronic low-grade inflammation, oxidative stress and impaired nitric-  
72 oxide (NO) bioavailability that contribute to endothelial dysfunction and large artery stiffness  
73 (54, 55). Endothelial dysfunction is considered an important prognostic factor and precursor to  
74 the development of atherosclerosis (22, 46), and is strongly associated with the risk of  
75 cardiovascular events (22, 57). In addition, endothelial dysfunction is suggested to contribute to  
76 other age-associated disorders including cognitive impairment and insulin resistance (60, 62, 72).  
77 As such, interventions that prevent or slow the detrimental changes in endothelial function are  
78 important in reducing cardiovascular risk and mortality associated with increasing age (56, 57).

79

80 Importantly, age-associated endothelial dysfunction, measured using flow-mediated dilation  
81 (FMD) of the brachial artery (59), can be attenuated with both regular physical activity (71) and  
82 exercise training (16, 23). Results of cross-sectional studies indicate that exercise-trained older  
83 adults have preserved endothelial function (17, 40, 45, 49), and reduced cardiovascular disease  
84 risk (63), compared with those who are not habitually active. This adaptive response is  
85 commonly attributed to the repeated episodes of elevated blood flow, and consequently shear  
86 stress, during exercise that induces vascular adaptation (21).

87

88 While the positive impact of chronic aerobic exercise on endothelial function is well described,  
89 the significance of the transient changes observed in endothelial function with an acute bout of  
90 exercise is less clear (15). To elucidate which forms of exercise are most likely to benefit  
91 cardiovascular health and function, recent studies have focussed on the acute FMD response and  
92 how it is modulated by factors such as exercise intensity. Some evidence suggests that the FMD

93 response to acute exercise may be *biphasic*, involving an immediate decrease, followed by a  
94 transient increase in FMD before returning to baseline levels (15). This may represent the  
95 initiation of an adaptive response, and be linked to the long-term benefit provided by exercise  
96 training on endothelial function at rest (23). This response is suggested to be exaggerated  
97 following higher-intensity exercise e.g. a larger immediate reduction followed by transient  
98 improvement in FMD (4, 11, 15, 32), and may contribute to recent observations of larger  
99 improvements in FMD following high-intensity interval exercise (HIIE) compared to moderate-  
100 intensity continuous exercise (MICE) training (47, 52). We hypothesize that the *bi-phasic* FMD  
101 response would be further exaggerated in individuals with endothelial dysfunction, a low  
102 cardiorespiratory fitness or no training history.

103

104 To date, there have been no comparisons of the FMD response to acute exercise between  
105 individuals of a higher and lower cardiorespiratory fitness. There is a strong association between  
106 a higher cardiorespiratory fitness and maintenance of FMD with aging (40). HIIE training  
107 improves cardiorespiratory fitness in healthy elderly adults to a greater extent than MICE  
108 training (28), suggesting that it may also modulate the FMD response to training. Despite this, no  
109 study has investigated the influence of a lower and higher cardiorespiratory fitness on the FMD  
110 response following acute exercise in the elderly. We therefore aimed to determine whether the  
111 effect of acute exercise on FMD differed between MICE and HIIE cycling in elderly males,  
112 when controlling for both exercise work and duration. In addition, we assessed the influence of  
113 cardiorespiratory fitness on the acute effect of exercise intensity on the FMD response between  
114 participants with higher and lower cardiorespiratory fitness. In line with previous findings in the  
115 young (4, 11), we hypothesised that acute HIIE would stimulate greater immediate reductions in

116 endothelial function compared to MICE, with subsequent elevation in FMD after 60-min. We  
117 also hypothesised that this overall response would be attenuated in those with a higher  
118 cardiorespiratory fitness.

119

## 120 **METHODS**

### 121 **Research Design**

122 Participants underwent four laboratory visits, each following an overnight fast, refraining from  
123 alcohol and exercise for 24h, and caffeine for 12h, before each visit. Participants consumed a  
124 standardised snack (4 oat breakfast biscuits, 20g carbohydrate, 8g fat) 3h prior to attending the  
125 laboratory, and the macronutrient content of this snack was unlikely to influence endothelial  
126 function (24, 70). Visit 1 consisted of baseline measurements of height, body mass and estimated  
127 body composition using bio-impedance scales (BC 545N, Tanita, Australia). After 10 min of  
128 supine rest, blood pressure was measured using a manual sphygmomanometer, which was  
129 followed by a maximal cycling test to determine cardiorespiratory fitness ( $VO_{2peak}$ ) and peak  
130 power output (PPO). Experimental visits (2-4) were randomised, counter-balanced and consisted  
131 of two separate acute cycling exercise conditions (moderate-intensity continuous vs. high-  
132 intensity interval) or a no-exercise control condition. Blood pressure and brachial FMD were  
133 assessed at baseline following 20 min of supine rest, and then repeated at 10- and 60-min  
134 following exercise/control. Laboratory conditions were standardised for each visit (room  
135 temperature:  $23 \pm 1^{\circ}C$ ) (63). To control for diurnal variation in blood pressure and vascular  
136 function, each visit was performed at the same time of day (33), and separated by 7 days.

137



138 **Participants**

139 Forty-seven healthy elderly males (mean  $\pm$  SD, aged  $70\pm 5$  y; BMI  $25.3\pm 3.4$  kg.m<sup>2</sup>) were  
140 recruited. Participants were included if they were able to exercise and were non-smokers (>12  
141 months no smoking history). Participants were excluded if they were aged >86 years, had a BMI  
142 >39, or a chronic cardiovascular or metabolic condition. During the study, participants were  
143 requested to continue to take all prescribed medication. Participants were informed of the  
144 methods and study design verbally and in writing before providing written informed consent.  
145 The study conformed to the Declaration of Helsinki and was approved by the institutional ethics  
146 committees.

147

148 **Maximal cardiorespiratory cycling test:** A maximal incremental cardiorespiratory fitness test  
149 was performed on an electro-magnetically braked cycle ergometer (Lode Corival, Groningen,  
150 Netherlands). Following a 3 min warm up at 0 W, the test began at 20 W and then increased by  
151 10 W each min until volitional cessation. Participants were required to self-select a pedal  
152 cadence (between 60 and 90 RPM) and maintain this throughout the test. Expired respiratory  
153 gases were collected throughout the test and data were averaged every 15 s (Parvo Medics, UT,  
154 USA) for the determination of oxygen consumption ( $VO_2$ ; mL·kg<sup>-1</sup>·min<sup>-1</sup>). Peak  $VO_2$  was  
155 determined as the highest 15 s average over the last 60 s of maximal exercise ( $VO_{2peak}$ ). Heart  
156 rate was measured continuously using 12-lead ECG (Mortara Inc., WI, USA) and recorded,  
157 along with perceived exertion (RPE) using the 0-10 Borg scale, during the final 10 s of each  
158 stage. All participants reached the criteria for maximum effort based upon attaining >2 of the  
159 following: a peak heart rate within 10 bpm of predicted age-related maximum; RPE (>9); a fall  
160 in pedal cadence (>10 RPM); a plateau in  $VO_2$  despite an increase in workload; and a respiratory

161 exchange ratio >1.15. Peak power output (W) was then used to establish the exercise intensity in  
162 the subsequent test visits.

163

164 **Acute exercise/control protocols:** Following pre-test measurements, participants performed 27  
165 min of continuous or interval cycling exercise, or no-exercise control (seated-rest). Both acute  
166 exercise protocols commenced with a 3-minute warm-up at 0 W, followed by either 24 min of:  
167 *i)* continuous moderate-intensity cycling at 40% PPO, or *ii)* high-intensity interval cycling  
168 involving 12 x 60 s bouts at 70% PPO, with each separated by 60 s at 10% PPO. Heart rate and  
169 RPE were recorded every 2 min. This design ensured the continuous and interval cycling  
170 exercise protocols were duration and work-matched. Immediately following exercise/control  
171 (<60 s), participants were moved to the supine position and asked to remain supine for post-test  
172 FMD measurements (at 10 and 60-min). Right brachial artery blood pressure was measured in  
173 triplicate using an automated device (Sphygmocor XCEL, AtCor Medical, NSW, Australia) 10-  
174 min before each FMD time-point to negate any effect of cuff inflation on FMD.

175

176 **Brachial artery flow-mediated dilation:** Brachial artery FMD was used as a measure of  
177 endothelial function (63). Measurements were performed in the supine position, on the right arm  
178 with the cuff placed distal to the olecranon process. High-resolution duplex ultrasound (T3000;  
179 Terason, Burlington, MA) with a 12-MHz multi-frequency linear array probe was used to image  
180 the brachial artery at the distal third of the upper arm and simultaneously record the longitudinal  
181 B-mode image and Doppler blood velocity trace. The angle of Doppler insonation was 60°.  
182 Images were optimised, and settings (depth, focus position and gain) were maintained between  
183 FMD assessments within each individual visit, and the location of the transducer was recorded

184 and marked on the skin using an indelible marker. Following a 60 s baseline recording period,  
185 the cuff was rapidly inflated to 220 mmHg and maintained for 5 min (D.E. Hokanson, Bellevue,  
186 WA). Ultrasound recordings resumed 30 s prior to rapid cuff deflation (<2 s) and continued for  
187 3 min thereafter, in accordance with recommendations (12, 63). All ultrasound scans were  
188 performed by the same trained sonographer.

189

190 Analysis of brachial artery diameter was performed using custom-designed edge-detection and  
191 wall-tracking software, which is largely independent of investigator bias. Recent papers describe  
192 the analysis approach in detail (12, 63). Briefly, from recordings of the synchronised artery  
193 diameter and blood velocity data, blood flow (the product of lumen cross-sectional area and  
194 Doppler velocity) was calculated at 30 Hz. Shear rate (an estimate of shear stress independent of  
195 viscosity) was calculated as 4 times mean blood velocity/vessel diameter. This semi-automated  
196 software possesses an intra-observer coefficient of variation (CV) of 6.7% and reduces error,  
197 with the reproducibility of diameter measurements significantly better than manual methods (64,  
198 73).

199

## 200 **Statistical analysis**

201 To differentiate the cohort on the basis of cardiorespiratory fitness, each participant was stratified  
202 into lower- ( $VO_{2\text{ peak}} < 27 \text{ ml.kg.min}^{-1}$ ) and higher ( $VO_{2\text{ peak}} > 31 \text{ ml.kg.min}^{-1}$ ) fitness (fit) group  
203 based on age- and sex-specific normative data (1). A three-way (fitness\*protocol\*time) linear  
204 mixed model (LMM) was employed to analyse changes in FMD parameters [brachial diameter,  
205 peak diameter and FMD (mm), FMD (%), time to peak, shear rate area-under-the-curve (SRAUC),  
206 blood flow,] and blood pressure between the two fitness groups (low and high fitness), across

207 “time” (baseline, 10- and 60-min post) during each protocol (control, moderate- and high-  
208 intensity exercise). As variability in the baseline artery diameter and shear rate may influence  
209 the magnitude of the FMD response (65), these parameters were included in the analysis as  
210 covariates (2, 10). In line with recent recommendations (5-7), we also performed an additional  
211 three-way LMM analysis of logarithmically transformed absolute diameter change (difference  
212 between peak and baseline diameter as the outcome, in mm), with logarithmically transformed  
213 baseline diameter and shear rate again included as covariates, specific to each FMD test. The  
214 logged absolute diameter change was then also interpreted in the conventional manner and is  
215 presented as “adjusted FMD%” for comparative purposes as suggested (9), in line with recent  
216 reports (4, 67). This allometric approach may be more accurate for scaling changes in diameter  
217 than percentage change alone, which makes implicit assumptions about the linearity of the  
218 relationship between baseline diameter and peak diameter (8). The strength of the relationships  
219 between cardiorespiratory fitness and changes in FMD after exercise and/or control were  
220 assessed using Pearson correlation coefficient.

221  
222 Similarly, a three-way LMM analysis was used to detect any differences in heart rate and  
223 perceived exertion in response to the acute protocols between the two fitness groups (low- and  
224 high-fit), across time (at 2 minute intervals) during each protocol (control, moderate- and high-  
225 intensity exercise). Statistically significant interactions were further investigated with multiple  
226 comparisons using the least significant difference approach (43, 51). Analyses were conducted  
227 using the Statistical Package for Social Sciences (Version 22; IBM SPSS Inc., Chicago, IL).  
228 Statistical significance was delimited at  $P \leq 0.05$  and exact  $P$  values are cited ( $P$  values of “0.00”

229 are reported as “<0.01”). Data are presented in the text as mean (95% confidence interval,  
230 95%CI) unless otherwise stated.

231

## 232 **Results**

### 233 **Baseline:**

#### 234 *Participant characteristics.*

235 Participant characteristics are presented in Table 1. Participant age was higher in the lower-fit  
236 compared to the higher-fit group [mean difference of 3 years (95% CI, -1 to 6),  $P=0.05$ ].  
237 Approximately one quarter of the participants were hypertensive (30% and 26% in the lower and  
238 higher fitness groups, respectively) and all hypertensive participants were taking blood-pressure  
239 controlling medication. Resting heart rate was lower in the higher-fit compared to lower-fit  
240 [mean difference 6  $\text{b}\cdot\text{min}^{-1}$  (95% CI, 2 to 10),  $P = 0.01$ ], but there were no differences in resting  
241 blood pressure or anthropometric variables between lower- and higher-fit groups.

#### 242 *Cardiorespiratory fitness.*

243 There was a mean difference of 11  $\text{ml}\cdot\text{kg}^{-1}\cdot\text{min}^{-1}$  (95% CI, 8 to 13,  $P<0.01$ ) in  $\text{VO}_2$  peak and 50  
244 Watts (95% CI, 30 to 70,  $P<0.01$ ) between higher and lower-fit groups.

245

### 246 **Heart rate and perceived exertion during the exercise protocols**

247 Heart rate responses were normalised for peak heart rate obtained during the cardiorespiratory  
248 fitness test. Heart rate was significantly higher during high-intensity exercise [mean 65 % $\text{HR}_{\text{peak}}$   
249 (95% CI, 62 to 68 %)] compared to moderate-intensity exercise [mean 58 % $\text{HR}_{\text{peak}}$  (95% CI, 55  
250 to 61%,  $P<0.01$ )], whilst both were elevated compared to control [mean 37 % $\text{HR}_{\text{peak}}$  (95% CI, 34  
251 to 40),  $P<0.01$ ]. There was no effect of fitness on the heart rate responses ( $P=0.24$ ). RPE was

252 higher during the HIIE [mean RPE 4 AU (95% CI, 3 to 5)] compared to moderate-intensity  
253 exercise [mean RPE 3 AU (95% CI, 2 to 4,  $P < 0.01$ )]. There was no effect of fitness on the RPE  
254 responses ( $P = 0.58$ ).

255

## 256 **Brachial artery flow-mediated dilation**

### 257 *Baseline flow-mediated dilation.*

258 The coefficient of variation for baseline FMD% across the three visits in this study was  $11.8 \pm 3.9$   
259 %, which is similar to those previously reported (10.1-14.7%) (66, 73). There were no  
260 differences in resting (pre-exercise/control) brachial diameter,  $FMD_{mm}$ , FMD%, or  $SR_{AUC}$  across  
261 the three separate testing days (Table 2;  $P > 0.05$ ).

### 262 *Effect of fitness on baseline flow-mediated dilation.*

263 There was no significant difference in resting FMD% between the lower- (Table 3a) and higher-  
264 fit groups (Table 3b) [mean difference of 0.2 % (95% CI, -0.8 to 0.9),  $P = 0.82$ ].  $SR_{AUC}$  was  
265 significantly higher in the lower-fit compared to the higher-fit group [mean difference of 3.2  
266  $10^3 \cdot s^{-1}$  (95% CI, 1.3 to 6.3),  $P = 0.04$ ], despite no differences in baseline diameter between fitness  
267 groups [mean difference of 0.2 mm (95% CI, -0.6 to 0.8),  $P = 0.13$ ]. Furthermore, time to peak  
268 diameter was significantly longer in the lower-fit compared to the higher-fit group [mean  
269 difference of 10 s (95% CI, 1 to 17),  $P = 0.02$ ].

270

### 271 **Effect of exercise intensity on the acute flow-mediated dilation response to exercise:**

272 Baseline and recovery (10 and 60 min post) brachial FMD% and associated variables are detailed  
273 in Tables 3a and 3b for the lower- and higher-fit groups, respectively. For clarity, post-hoc  $P$

274 values are reported only in the text. Delta FMD% data are summarised in Figure 1, which shows  
275 the change in FMD% from baseline during recovery (10 and 60 min post).

276

277 In both fitness groups, FMD decreased by 0.74 % (95% CI, -1.34 to -0.03) after 60-min of  
278 recovery in control compared to baseline ( $P=0.05$ ). There was no effect of fitness on this  
279 response. There was a significant fitness x condition x time interaction for FMD% ( $P=0.01$ ).

280 FMD% was significantly reduced compared to baseline following high-intensity exercise in the

281 lower-fit group at both 10 min [mean reduction of 0.85 % (95% CI, 0.12 to 1.58),  $P=0.02$ ] and

282 60 min post [mean reduction of 0.72 % (95% CI, 0.02 to 1.46),  $P=0.05$ ] (see Table 3a). In the

283 higher-fit group, a negligible change in FMD% was observed 10 min after high-intensity

284 exercise [mean difference of 0.13 % (95% CI, -0.73 to 0.98),  $P=0.77$ ], however there was a

285 significant increase in FMD % compared to baseline after 60-min of 0.84 % (95% CI, -0.12 to

286 1.69;  $P=0.05$ ) (see Figure 1). The improved FMD% response following HIIE elicited a mean

287 difference of 1.52 % (95% CI, 0.41 to 2.62) after 60 min in the higher-fit compared to the lower-

288 fit group ( $P=0.01$ ; Table 3a and 3b). In support of this difference between fitness groups, the

289 delta change in FMD% after high-intensity exercise at 60 min was significantly correlated with

290  $VO_{2peak}$  ( $r = 0.41$ ;  $P<0.01$ ). Furthermore, in the higher-fit group, FMD% was elevated after 60-

291 min compared to moderate-intensity and control protocols [mean difference of 0.92% (95% CI,

292 0.05 to 1.78,  $P=0.01$ ) and 1.54% (95% CI, 0.65 to 2.42,  $P=0.02$ ) (Table 3b). These changes in

293 FMD% were also observed for absolute FMD (mm), with an increase 60-min following high-

294 intensity exercise in the higher-, but not lower-fit group ( $P=0.04$ ; Table 3a and 3b).

295

296 FMD% increased significantly from baseline 10 min after moderate-intensity exercise [mean  
297 change of 0.86 % (95% CI, 0.17 to 1.56),  $P=0.02$ ; Figure 1], and returned to baseline levels after  
298 60 min [mean difference to baseline of 0.30 % (95% CI, -0.59 to 0.53),] with no effect of fitness  
299 on the response [mean between fitness group difference of 0.43 % (95% CI, -0.28 to 1.13),  
300  $P=0.23$ ;  $r = -0.13$ ,  $P=0.38$ ]. Furthermore, the FMD% response 10-min after moderate-intensity  
301 exercise was increased compared to the high-intensity response [mean difference of 1.15 % (95%  
302 CI, 0.58 to 1.72),  $P<0.001$ ] and control [mean difference of 1.23 % (95% CI, 0.72 to 1.88),  
303  $P<0.001$ ] in both fitness groups (Figure 1). In the lower-fit group, an increase in FMD% was  
304 observed 10 min after moderate-intensity exercise compared to the reduction observed after  
305 high-intensity exercise [mean difference of 1.34 % (95% CI, 0.60 to 2.09),  $P<0.001$ ] and control  
306 [mean difference of 0.99% (95% CI, 0.23 to 1.75),  $P = 0.01$ ] (Table 3a).

307  
308 We also present covariate “adjusted FMD%” values (Table 3a/b). This analysis was consistent  
309 with our initial observations in FMD%, with a significant interaction between condition, fitness  
310 and time ( $P=0.04$ ). Post-hoc analysis revealed significant differences between the lower- and  
311 higher-fit groups 60-min after HIIE ( $P<0.01$ ).

312

### 313 **Blood flow and shear rate responses**

314 Resting blood flow was significantly elevated 10 min following both exercise protocols  
315 compared to control ( $P<0.01$ ), and was higher following high-intensity exercise compared with  
316 moderate-intensity [mean difference of 0.36 mL.s<sup>-1</sup> (95% CI, -0.03 to 0.66),  $P=0.05$ ]. There was  
317 no effect of fitness on the blood flow responses to exercise ( $P=0.79$ ) (Table 3a and 3b). Shear  
318 rate demonstrated a similar pattern where it was elevated 10 min after both exercise protocols



319 compared with control ( $P=0.01$ ), and was higher immediately after high-intensity compared to  
320 moderate-intensity exercise [mean difference of  $17.38 \times 10^3 \text{ s}^{-1}$  (95% CI, -3.86 to 38.62),  $P=0.01$ ].  
321 There was no effect of fitness on the shear rate responses after exercise ( $P=0.78$ ) (Table 3a and  
322 3b).

323

### 324 **Heart rate and blood pressure responses after exercise**

325 There was a condition x time interaction for HR, SBP and MAP (Table 3a and 3b;  $P<0.01$ ).  
326 Heart rate was elevated by  $9 \text{ b}\cdot\text{min}^{-1}$  (95% CI, 8 to 12) and by  $13 \text{ b}\cdot\text{min}^{-1}$  (95% CI, 11 to 15) 10  
327 min after moderate-intensity and high-intensity exercise, respectively, compared to rest. MAP  
328 was 5 mmHg (95% CI, 3 to 8) and 6 mmHg (95% CI, 3 to 9) higher 10-min after moderate- and  
329 high-intensity exercise, respectively, compared to rest.

330

331

### 332 **Discussion**

333 To our knowledge, this is the first study to investigate the acute effects of exercise intensity and  
334 cardiorespiratory fitness on endothelial function in healthy, elderly adults. The main findings  
335 from this study indicate that the acute effects of exercise on brachial FMD are dependent on both  
336 the intensity of exercise and cardiorespiratory fitness in the elderly. We observed an immediate  
337 increase in FMD following MICE that normalised after 60 min in both fitness groups. In  
338 contrast, FMD decreased immediately and 60 min following HIIE in the lower-fit, whereas FMD  
339 increased after 60 min in the higher-fit participants. We also observed reductions in FMD in both  
340 groups following prolonged rest in control.

341

342 The FMD response to acute exercise is suggested to be *biphasic* (15), with an inverse  
343 relationship between exercise-intensity and the recovery in brachial artery endothelium-  
344 dependent function observed in some (11, 32) but not all studies (4, 58). We attempted to capture  
345 the time-course response by measuring FMD immediately (10 min post) and 60 min after  
346 exercise in the elderly and found an exercise intensity-dependent decrease in endothelial function  
347 immediately after high-intensity exercise, which is consistent with previous findings in young  
348 (11, 32), hypertensive (38) and peripheral arterial disease patients (34). Conversely, we found an  
349 immediate increase in endothelial function after short-term moderate-intensity exercise, which  
350 has been observed in one (32), but not all (4, 11) studies in younger individuals, and following  
351 30 min of walking exercise in healthy middle-aged adults (13). The immediate improvement in  
352 FMD after MICE of 40% PPO in this study contrasts the finding of no-change in FMD following  
353 cycling exercise at 50% HR<sub>max</sub> in albeit, younger healthy individuals (11). This difference in  
354 findings may be due to the degree of baseline endothelial dysfunction in elderly compared to  
355 younger adults, with greater improvements in acute FMD observed after exercise in coronary  
356 artery disease patients with a lower baseline FMD (14). Moreover, the increase in FMD after  
357 moderate-intensity exercise normalised after 60 min which is similar in younger adults (32).

358

359 In line with the suggested effect of higher-intensity exercise (>70% HR<sub>max</sub>) on the *bi-phasic*  
360 FMD response, we observed an increase in FMD 60 min after HIIE compared to normalisation  
361 of FMD after MICE in the higher-fit elderly adults. This contrasts a report by Currie *et al.*  
362 (2012), who found an increased FMD after both high- and moderate-intensity exercise in  
363 coronary artery disease patients. However, unlike the study by Currie and colleagues, our  
364 exercise protocols were duration and work matched, which is important as the dose of exercise

365 affects FMD independent of intensity (32). Our study reports intensity-dependent, dose-matched  
366 differences in the *bi-phasic* FMD response in elderly adults. We provide further support that  
367 exercise intensity modulates acute endothelial function (4, 11, 18, 32), in elderly healthy adults.

368

369 The rationale for assessing the acute response of endothelial function to exercise relates to the  
370 potential impact of repeated bouts of exercise on vascular adaptation (23), but whether the  
371 immediate increase or decrease in FMD after exercise in this study is important for future  
372 vascular adaptation in the elderly is unknown. Padilla *et al.* (2011) suggest recurring periods of  
373 exercise-induced transient endothelial impairment may represent a beneficial stimulus that  
374 contributes to longer-term improvements in vascular function and structure, a concept known as  
375 *hormesis*. That is, the initial challenge, e.g. acute reductions in FMD, leads to activation of  
376 beneficial adaptive processes (42). The acute exercise-intensity dependent reductions in FMD we  
377 observed in this study may be linked to the recent observation that HIIE training is likely more  
378 effective in improving conduit artery endothelial function compared to MICE (47), therefore  
379 improving FMD immediately after moderate-intensity exercise (which normalised after 60 min)  
380 may not lead to beneficial long-term vascular adaptation with training. Interestingly, we  
381 observed that cardiorespiratory fitness modulates the *bi-phasic* response of FMD to high-, but  
382 not moderate-intensity exercise in the elderly. The lack of a *bi-phasic* response in the lower fit  
383 individuals after high-intensity exercise, with sustained reductions in FMD, may be the signal  
384 required for future vascular adaptation observed following training and increases in fitness (42,  
385 61).

386

387 Our study is the first to directly assess the effect of cardiorespiratory fitness levels on acute  
388 changes in FMD following exercise in the elderly. The positive relationship between exercise  
389 training and endothelial function is well established (39, 40), whilst cardiorespiratory fitness is  
390 related to training status (36) and can be modified through changes in routine physical activity  
391 (25, 41). In support of this, acute reductions in FMD have been reported in sedentary, but not  
392 active adults after both leg-press exercise (44), and maximal running (29). Whether the  
393 similarities observed in the reduced FMD response after HIIE in the present study reflect the low  
394 overall physical activity levels or the impact of low activity on reductions in cardiorespiratory  
395 fitness is not known.

396

397 The mechanisms responsible for exercise-induced, intensity-dependent changes in FMD have  
398 been proposed to include alterations in oxidative stress, inflammation, shear stress and shear  
399 pattern, blood pressure, baseline artery diameter, sympathetic nerve activity and vasoconstrictors  
400 (15). As we did not assess mechanisms of FMD changes, we can only speculate on the possible  
401 causes. We did not report any differences in blood pressure between exercise intensities so this is  
402 unlikely to be the cause of our observed differences, whilst we covariate controlled for exercise-  
403 induced changes in artery diameter and shear stress. The altered FMD response between exercise  
404 intensities may be linked to NO bioavailability (50), and shear stress patterns during exercise as  
405 this is known to directly contribute to changes in FMD (20, 66, 69). Large increases in brachial  
406 antegrade shear stress occur during cycling exercise (20) and are associated with improved FMD  
407 (69), whilst increases in oscillatory shear and/or retrograde flow lead to reductions in FMD (53).  
408 Increases in oscillatory flow are observed early during cycling exercise (20), but may also be  
409 augmented in interval compared to continuous exercise in this study, due to the stop-start nature

410 of the high-intensity modality. This may explain the immediate improvement in FMD after  
411 MICE compared to the reduced FMD immediately following HIIE. Similarly, reductions in FMD  
412 immediately after exercise of higher-intensity, and not moderate-intensity exercise, may be due  
413 to dose-dependent increases in brachial artery blood flow and the production of reactive oxygen  
414 species (18, 30) endothelin-1 expression (27) or increased sympathetic nervous activity (26). An  
415 increase in NO bioavailability, even in the presence of large changes in reactive oxygen species,  
416 may explain the differing responses we observed in FMD after high-intensity exercise between  
417 the high- and low-fit groups. In line with this, arterial compliance is compromised in elderly  
418 adults with a lower-, but not higher- cardiorespiratory fitness during similar increases in  
419 inflammation (31). Hence, a higher fitness in elderly adults may be associated with improved  
420 anti-inflammatory mechanisms, such as BH4 synthesis (3) that preserves NO bioavailability  
421 when exposed to increases in reactive oxygen species. We acknowledge that these proposed  
422 mechanisms are speculative and should form the basis of future investigations.

423

424 Our findings highlight the *exercise paradox*, where those who are at the greatest risk of adverse  
425 responses to acute exercise, have the most to gain from regular exercise and activity (37). Elderly  
426 individuals with low endothelial function who exhibit further reductions in FMD after higher-  
427 intensity exercise may be at increased cardiovascular risk. In this study, FMD was reduced  
428 significantly 60 min after high-intensity exercise in the elderly lower-fit individuals, compared to  
429 those with a higher fitness. The acute reduction in FMD was not observed following MICE in  
430 either group or recovery time-point. Whether the acute reduction is necessary to induce vascular  
431 adaptation (see *hormesis*, discussed above) (42, 61) and represents a potential danger period  
432 where the vascular system may be less responsive to stress is unknown. However, higher fitness

433 in this study did attenuate the reduction in FMD observed following high-intensity exercise,  
434 suggesting there may be an adaptive or tolerance response with improvements in  
435 cardiorespiratory fitness.

436

437 Studies investigating the acute effect of exercise intensity on endothelial function do not  
438 commonly assess FMD across the same measurement period using a non-exercise control. This  
439 study is unique in that it offers the opportunity to assess changes in FMD during extended  
440 periods of sedentary time in the elderly. We observed a reduction in brachial artery FMD after  
441 ~120 min of “sedentary time” which is not reported in younger individuals after 6 hours of  
442 prolonged sitting (48). As sitting time increases all-cause and cardiovascular mortality risk in  
443 older adults (37), the vascular effects of prolonged sitting warrants investigation. In line with  
444 recent evidence (48), we showed that reductions in FMD with sedentary time can be attenuated  
445 with short-term moderate-intensity exercise. However, we also found that high-intensity exercise  
446 in lower-fit individuals led to a similar decline in FMD to that of prolonged supine rest. This  
447 suggests that prescribing moderate-intensity in lower-fit elderly individuals might be considered  
448 before progressing to higher-intensity exercise as cardiorespiratory fitness improves.

449

450 A modest association exists between cardiorespiratory fitness and basal endothelial function,  
451 independent of age and health status (39). Similarly, aerobically trained middle-aged and older  
452 adults have preserved endothelial function compared to those who are sedentary (16, 17, 40, 45,  
453 49), however in this study investigating FMD in the elderly there was no difference in resting  
454 brachial artery FMD between lower- and higher-fit groups. This may be due to normalised FMD  
455 in the higher-fit following increases in artery diameter and structural remodelling observed with

456 exercise training (35, 68) with a tendency for a larger arterial diameter in the higher-fit compared  
457 to the lower-fit group. It is also possible that a “ceiling” effect exists on basal FMD in the  
458 elderly, as no improvements in FMD were reported following short-term training in older,  
459 higher-fit adults despite increases in  $VO_{2peak}$  (19).

460

#### 461 **Study limitations**

462 In future studies, it would be interesting to have prolonged FMD measurements e.g. 2h-24h after  
463 exercise to establish whether the *bi-phasic* pattern is delayed or persistent in the lower-fit  
464 compared to higher-fit individuals, particularly after high-intensity exercise. A limitation of our  
465 study is that we did not include measures of potential mechanisms involved in the changes in  
466 FMD we observed, such as the inflammatory response to exercise. However, the study was  
467 designed to explore whether cardiorespiratory fitness had an effect on acute FMD following  
468 moderate- and high-intensity exercise that were matched for workload. Nonetheless, as  
469 individuals are now living longer, and age-associated reductions in endothelial function become  
470 a growing concern for CVD, it is necessary to investigate the effect of exercise as a potential  
471 therapy on those that would benefit the most e.g. elderly individuals of a lower cardiorespiratory  
472 fitness.

473

#### 474 **Clinical relevance**

475 Ischemic events typically occur in the elderly who have known cardiovascular risk factors and/or  
476 disease. It is known that regular physical activity and exercise training throughout the lifespan  
477 has cardio-protective and vascular effects. Recently, HIIE has become popular for its potential  
478 for additional cardiovascular benefits in a shorter bouts of exercise, including the ability to

479 improve endothelial function (47). However, in the elderly who are of a lower fitness and/or  
480 those who already exhibit vascular dysfunction, this type of exercise may need to be treated with  
481 caution due to the potential that vascular dysfunction is transiently exacerbated. Importantly,  
482 whether the differences in the FMD response to different acute exercise intensities reported here  
483 has longer-term consequences on endothelial function and/or CV risk in healthy elderly  
484 individuals needs to be determined.

485

## 486 **Conclusions**

487 In conclusion, the present study illustrates the effect of exercise intensity on acute FMD  
488 responses in the elderly. Furthermore, we highlight the importance of cardiorespiratory fitness on  
489 the acute FMD response following high-intensity exercise. Increases in FMD after moderate-  
490 intensity exercise normalised quickly. Conversely, there was prolonged benefit in FMD after  
491 high-intensity exercise in those with a higher-fitness, whereas lower-fitness individuals exhibited  
492 sustained endothelial dysfunction. This decrease in FMD may represent the signal for an  
493 adaptive vascular response and/or endothelial fatigue in untrained elderly individuals. Further  
494 studies on the acute and training effects of exercise intensity on endothelial function will be  
495 important to establish the link between changes in FMD with acute exercise and the potential for  
496 chronic adaptation with exercise training in the elderly.

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## References

- 506  
507  
508 1. **ACSM.** *ACSM's Guidelines for Exercise Testing and Prescription.* Baltimore, USA:  
509 2010.
- 510 2. **Ainslie PN, and Bailey DM.** Your ageing brain: the lows and highs of cerebral  
511 metabolism. *The Journal of Physiology* 591: 1591-1592, 2013.
- 512 3. **Antoniades C, Cunnington C, Antonopoulos A, Neville M, Margaritis M,**  
513 **Demosthenous M, Bendall J, Hale A, Cerrato R, Tousoulis D, Bakogiannis C, Marinou K,**  
514 **Toutouza M, Vlachopoulos C, Leeson P, Stefanadis C, Karpe F, and Channon KM.**  
515 Induction of vascular GTP-cyclohydrolase I and endogenous tetrahydrobiopterin synthesis  
516 protect against inflammation-induced endothelial dysfunction in human atherosclerosis.  
517 *Circulation* 124: 1860-1870, 2011.
- 518 4. **Atkinson CL, Carter HH, Dawson EA, Naylor LH, Thijssen DH, and Green DJ.**  
519 Impact of handgrip exercise intensity on brachial artery flow-mediated dilation. *Eur J Appl*  
520 *Physiol* 115: 1705-1713, 2015.
- 521 5. **Atkinson G.** The dependence of FMD% on baseline diameter: a problem solved by  
522 allometric scaling. *Clinical science (London, England : 1979)* 125: 53-54, 2013.
- 523 6. **Atkinson G, and Batterham AM.** Allometric scaling of diameter change in the original  
524 flow-mediated dilation protocol. *Atherosclerosis* 226: 425-427, 2013.
- 525 7. **Atkinson G, and Batterham AM.** The clinical relevance of the percentage flow-  
526 mediated dilation index. *Curr Hypertens Rep* 17: 4, 2015.
- 527 8. **Atkinson G, and Batterham AM.** The percentage flow-mediated dilation index: a large-  
528 sample investigation of its appropriateness, potential for bias and causal nexus in vascular  
529 medicine. *Vascular medicine (London, England)* 18: 354-365, 2013.
- 530 9. **Atkinson G, Batterham AM, Thijssen DH, and Green DJ.** A new approach to improve  
531 the specificity of flow-mediated dilation for indicating endothelial function in cardiovascular  
532 research. *Journal of hypertension* 31: 287-291, 2013.
- 533 10. **Bailey TG, Birk GK, Cable NT, Atkinson G, Green DJ, Jones H, and Thijssen DHJ.**  
534 Remote ischemic preconditioning prevents reduction in brachial artery flow-mediated dilation  
535 after strenuous exercise. *American Journal of Physiology - Heart and Circulatory Physiology*  
536 303: H533-H538, 2012.
- 537 11. **Birk GK, Dawson EA, Batterham AM, Atkinson G, Cable T, Thijssen DH, and**  
538 **Green DJ.** Effects of exercise intensity on flow mediated dilation in healthy humans. *Int J*  
539 *Sports Med* 34: 409-414, 2013.
- 540 12. **Black MA, Cable NT, Thijssen DH, and Green DJ.** Importance of measuring the time  
541 course of flow-mediated dilatation in humans. *Hypertension* 51: 203-210, 2008.
- 542 13. **Cosio-Lima LM, Thompson PD, Reynolds KL, Headley SA, Winter CR, Manos T,**  
543 **Lagasse MA, Todorovich JR, and Germain M.** The acute effect of aerobic exercise on  
544 brachial artery endothelial function in renal transplant recipients. *Prev Cardiol* 9: 211-214, 2006.
- 545 14. **Currie KD, McKelvie RS, and Macdonald MJ.** Brachial artery endothelial responses  
546 during early recovery from an exercise bout in patients with coronary artery disease. *BioMed*  
547 *research international* 2014: 591918, 2014.
- 548 15. **Dawson EA, Green DJ, Cable NT, and Thijssen DH.** Effects of acute exercise on flow-  
549 mediated dilatation in healthy humans. *Journal of applied physiology (Bethesda, Md : 1985)*  
550 115: 1589-1598, 2013.

- 551 16. **DeSouza CA, Shapiro LF, Clevenger CM, Dinunno FA, Monahan KD, Tanaka H,**  
552 **and Seals DR.** Regular aerobic exercise prevents and restores age-related declines in  
553 endothelium-dependent vasodilation in healthy men. *Circulation* 102: 1351-1357, 2000.
- 554 17. **Franzoni F, Ghiadoni L, Galetta F, Plantinga Y, Lubrano V, Huang Y, Salvetti G,**  
555 **Regoli F, Taddei S, Santoro G, and Salvetti A.** Physical activity, plasma antioxidant capacity,  
556 and endothelium-dependent vasodilation in young and older men. *American journal of*  
557 *hypertension* 18: 510-516, 2005.
- 558 18. **Goto C, Higashi Y, Kimura M, Noma K, Hara K, Nakagawa K, Kawamura M,**  
559 **Chayama K, Yoshizumi M, and Nara I.** Effect of different intensities of exercise on  
560 endothelium-dependent vasodilation in humans: role of endothelium-dependent nitric oxide and  
561 oxidative stress. *Circulation* 108: 530-535, 2003.
- 562 19. **Grace FM, Herbert P, Ratcliffe JW, New KJ, Baker JS, and Sculthorpe NF.** Age  
563 related vascular endothelial function following lifelong sedentariness: positive impact of  
564 cardiovascular conditioning without further improvement following low frequency high intensity  
565 interval training. *Physiological reports* 3: 2015.
- 566 20. **Green DJ, Bilsborough W, Naylor LH, Reed C, Wright J, O'Driscoll G, and Walsh**  
567 **JH.** Comparison of forearm blood flow responses to incremental handgrip and cycle ergometer  
568 exercise: relative contribution of nitric oxide. *J Physiol* 562: 617-628, 2005.
- 569 21. **Green DJ, Hopman MTE, Padilla J, Laughlin MH, and Thijssen DHJ.** Vascular  
570 adaptation to exercise in humans: The role of hemodynamic stimuli. *Physiological reviews* In  
571 Press Sept 13.: 2016.
- 572 22. **Green DJ, Jones H, Thijssen D, Cable NT, and Atkinson G.** Flow-Mediated Dilation  
573 and Cardiovascular Event Prediction. *Hypertension* 57: 363-369, 2011.
- 574 23. **Green DJ, Maiorana A, O'Driscoll G, and Taylor R.** Effect of exercise training on  
575 endothelium-derived nitric oxide function in humans. *J Physiol* 561: 1-25, 2004.
- 576 24. **Greyling A, Schreuder TH, Landman T, Draijer R, Verheggen RJ, Hopman MT,**  
577 **and Thijssen DH.** Elevation in blood flow and shear rate prevents hyperglycemia-induced  
578 endothelial dysfunction in healthy subjects and those with type 2 diabetes. *Journal of applied*  
579 *physiology (Bethesda, Md : 1985)* 118: 579-585, 2015.
- 580 25. **Haskell WL, Lee IM, Pate RR, Powell KE, Blair SN, Franklin BA, Macera CA,**  
581 **Heath GW, Thompson PD, and Bauman A.** Physical activity and public health: Updated  
582 recommendation for adults from the American College of Sports Medicine and the American  
583 Heart Association. *Circulation* 116: 1081-1093, 2007.
- 584 26. **Hijmering ML, Stroes ES, Olijhoek J, Hutten BA, Blankestijn PJ, and Rabelink TJ.**  
585 Sympathetic activation markedly reduces endothelium-dependent, flow-mediated vasodilation. *J*  
586 *Am Coll Cardiol* 39: 683-688, 2002.
- 587 27. **Himburg HA, Dowd SE, and Friedman MH.** Frequency-dependent response of the  
588 vascular endothelium to pulsatile shear stress. *American journal of physiology Heart and*  
589 *circulatory physiology* 293: H645-653, 2007.
- 590 28. **Hwang C-L, Yoo J-K, Kim H-K, Hwang M-H, Handberg EM, Petersen JW, and**  
591 **Christou DD.** Novel all-extremity high-intensity interval training improves aerobic fitness,  
592 cardiac function and insulin resistance in healthy older adults. *Experimental gerontology* 82:  
593 112-119, 2016.
- 594 29. **Hwang IC, Kim KH, Choi WS, Kim HJ, Im MS, Kim YJ, Kim SH, Kim MA, Sohn**  
595 **DW, and Zo JH.** Impact of acute exercise on brachial artery flow-mediated dilatation in young  
596 healthy people. *Cardiovascular ultrasound* 10: 39, 2012.

- 597 30. **Hwang J, Ing MH, Salazar A, Lassegue B, Griendling K, Navab M, Sevanian A, and**  
598 **Hsiai TK.** Pulsatile versus oscillatory shear stress regulates NADPH oxidase subunit expression:  
599 implication for native LDL oxidation. *Circulation research* 93: 1225-1232, 2003.
- 600 31. **Jae SY, Yoon ES, Jung SJ, Jung SG, Park SH, Kim BS, Heffernan KS, and Fernhall**  
601 **B.** Effect of cardiorespiratory fitness on acute inflammation induced increases in arterial stiffness  
602 in older adults. *Eur J Appl Physiol* 113: 2159-2166, 2013.
- 603 32. **Johnson BD, Padilla J, and Wallace JP.** The exercise dose affects oxidative stress and  
604 brachial artery flow-mediated dilation in trained men. *Eur J Appl Physiol* 112: 33-42, 2012.
- 605 33. **Jones H, Green DJ, George K, and Atkinson G.** Intermittent exercise abolishes the  
606 diurnal variation in endothelial-dependent flow-mediated dilation in humans. *American journal*  
607 *of physiology Regulatory, integrative and comparative physiology* 298: R427-432, 2010.
- 608 34. **Joras M, and Poredos P.** The association of acute exercise-induced ischaemia with  
609 systemic vasodilator function in patients with peripheral arterial disease. *Vascular medicine*  
610 *(London, England)* 13: 255-262, 2008.
- 611 35. **Laughlin MH, Newcomer SC, and Bender SB.** Importance of hemodynamic forces as  
612 signals for exercise-induced changes in endothelial cell phenotype. *Journal of applied physiology*  
613 *(Bethesda, Md : 1985)* 104: 588-600, 2008.
- 614 36. **Lin X, Zhang X, Guo J, Roberts CK, McKenzie S, Wu WC, Liu S, and Song Y.**  
615 Effects of Exercise Training on Cardiorespiratory Fitness and Biomarkers of Cardiometabolic  
616 Health: A Systematic Review and Meta-Analysis of Randomized Controlled Trials. *Journal of*  
617 *the American Heart Association* 4: 2015.
- 618 37. **Matthews CE, Moore SC, Sampson J, Blair A, Xiao Q, Keadle SK, Hollenbeck A,**  
619 **and Park Y.** Mortality Benefits for Replacing Sitting Time with Different Physical Activities.  
620 *Med Sci Sports Exerc* 47: 1833-1840, 2015.
- 621 38. **McGowan CL, Levy AS, Millar PJ, Guzman JC, Morillo CA, McCartney N, and**  
622 **Macdonald MJ.** Acute vascular responses to isometric handgrip exercise and effects of training  
623 in persons medicated for hypertension. *American journal of physiology Heart and circulatory*  
624 *physiology* 291: H1797-1802, 2006.
- 625 39. **Montero D.** The association of cardiorespiratory fitness with endothelial or smooth  
626 muscle vasodilator function. *European journal of preventive cardiology* 22: 1200-1211, 2015.
- 627 40. **Montero D, Padilla J, Diaz-Canestro C, Muris DM, Pyke KE, Obert P, and Walther**  
628 **G.** Flow-mediated dilation in athletes: influence of aging. *Med Sci Sports Exerc* 46: 2148-2158,  
629 2014.
- 630 41. **Myers J, McAuley P, Lavie CJ, Despres J-P, Arena R, and Kokkinos P.** Physical  
631 Activity and Cardiorespiratory Fitness as Major Markers of Cardiovascular Risk: Their  
632 Independent and Interwoven Importance to Health Status. *Progress in Cardiovascular Diseases*  
633 57: 306-314, 2015.
- 634 42. **Padilla J, Simmons GH, Bender SB, Arce-Esquivel AA, Whyte JJ, and Laughlin**  
635 **MH.** Vascular Effects of Exercise: Endothelial Adaptations Beyond Active Muscle Beds.  
636 *Physiology* 26: 132-145, 2011.
- 637 43. **Perneger TV.** Whats wrong with Bonferroni adjustments? *British Medical Journal* 316:  
638 1236, 1998.
- 639 44. **Phillips SA, Das E, Wang J, Pritchard K, and Gutterman DD.** Resistance and aerobic  
640 exercise protects against acute endothelial impairment induced by a single exposure to  
641 hypertension during exertion. *Journal of applied physiology (Bethesda, Md : 1985)* 110: 1013-  
642 1020, 2011.

- 643 45. **Pierce GL, Donato AJ, LaRocca TJ, Eskurza I, Silver AE, and Seals DR.** Habitually  
644 exercising older men do not demonstrate age-associated vascular endothelial oxidative stress.  
645 *Aging cell* 10: 1032-1037, 2011.
- 646 46. **Raitakari OT, and Celermajer DS.** Flow-mediated dilatation. *British journal of clinical*  
647 *pharmacology* 50: 397-404, 2000.
- 648 47. **Ramos JS, Dalleck LC, Tjonna AE, Beetham KS, and Coombes JS.** The impact of  
649 high-intensity interval training versus moderate-intensity continuous training on vascular  
650 function: a systematic review and meta-analysis. *Sports medicine (Auckland, NZ)* 45: 679-692,  
651 2015.
- 652 48. **Restaino RM, Holwerda SW, Credeur DP, Fadel PJ, and Padilla J.** Impact of  
653 prolonged sitting on lower and upper limb micro- and macrovascular dilator function. *Exp*  
654 *Physiol* 100: 829-838, 2015.
- 655 49. **Rinder MR, Spina RJ, and Ehsani AA.** Enhanced endothelium-dependent vasodilation  
656 in older endurance-trained men. *Journal of applied physiology (Bethesda, Md : 1985)* 88: 761-  
657 766, 2000.
- 658 50. **Rognmo O, Bjornstad TH, Kahrs C, Tjonna AE, Bye A, Haram PM, Stolen T,**  
659 **Slordahl SA, and Wisloff U.** Endothelial function in highly endurance-trained men: effects of  
660 acute exercise. *J Strength Cond Res* 22: 535-542, 2008.
- 661 51. **Rothman KJ.** No adjustments are needed for multiple comparisons. *Epidemiology* 1: 43-  
662 46, 1990.
- 663 52. **Sawyer BJ, Tucker WJ, Bhammar DM, Ryder JR, Sweazea KL, and Gaesser GA.**  
664 Effects of high-intensity interval training and moderate-intensity continuous training on  
665 endothelial function and cardiometabolic risk markers in obese adults. *Journal of applied*  
666 *physiology (Bethesda, Md : 1985)* 121: 279-288, 2016.
- 667 53. **Schreuder TH, Green DJ, Hopman MT, and Thijssen DH.** Acute impact of retrograde  
668 shear rate on brachial and superficial femoral artery flow-mediated dilation in humans.  
669 *Physiological reports* 2: e00193, 2014.
- 670 54. **Seals DR, Jablonski KL, and Donato AJ.** Aging and vascular endothelial function in  
671 humans. *Clinical science (London, England : 1979)* 120: 357-375, 2011.
- 672 55. **Seals DR, Kaplon RE, Gioscia-Ryan RA, and LaRocca TJ.** You're only as old as your  
673 arteries: translational strategies for preserving vascular endothelial function with aging.  
674 *Physiology (Bethesda, Md)* 29: 250-264, 2014.
- 675 56. **Seals DR, Walker AE, Pierce GL, and Lesniewski LA.** Habitual exercise and vascular  
676 ageing. *J Physiol* 587: 5541-5549, 2009.
- 677 57. **Shechter M, Issachar A, Marai I, Koren-Morag N, Freinark D, Shahar Y, Shechter**  
678 **A, and Feinberg MS.** Long-term association of brachial artery flow-mediated vasodilation and  
679 cardiovascular events in middle-aged subjects with no apparent heart disease. *International*  
680 *journal of cardiology* 134: 52-58, 2009.
- 681 58. **Siasos G, Athanasiou D, Terzis G, Stasinaki A, Oikonomou E, Tsitkanou S,**  
682 **Kolokytha T, Spengos K, Papavassiliou AG, and Tousoulis D.** Acute effects of different types  
683 of aerobic exercise on endothelial function and arterial stiffness. *European journal of preventive*  
684 *cardiology* 2016.
- 685 59. **Skaug EA, Aspenes ST, Oldervoll L, Morkedal B, Vatten L, Wisloff U, and**  
686 **Ellingsen O.** Age and gender differences of endothelial function in 4739 healthy adults: the  
687 HUNT3 Fitness Study. *European journal of preventive cardiology* 20: 531-540, 2013.

- 688 60. **Stanimirovic DB, and Friedman A.** Pathophysiology of the neurovascular unit: disease  
689 cause or consequence? *Journal of cerebral blood flow and metabolism : official journal of the*  
690 *International Society of Cerebral Blood Flow and Metabolism* 32: 1207-1221, 2012.
- 691 61. **Suvorava T, and Kojda G.** Prevention of transient endothelial dysfunction in acute  
692 exercise: a friendly fire? *Thrombosis and haemostasis* 97: 331-333, 2007.
- 693 62. **Tachibana H, Washida K, Kowa H, Kanda F, and Toda T.** Vascular Function in  
694 Alzheimer's Disease and Vascular Dementia. *American journal of Alzheimer's disease and other*  
695 *dementias* 2016.
- 696 63. **Thijssen DH, Black MA, Pyke KE, Padilla J, Atkinson G, Harris RA, Parker B,**  
697 **Widlansky ME, Tschakovsky ME, and Green DJ.** Assessment of flow-mediated dilation in  
698 humans: a methodological and physiological guideline. *American journal of physiology Heart*  
699 *and circulatory physiology* 300: H2-12, 2011.
- 700 64. **Thijssen DH, Dawson EA, Black MA, Hopman MT, Cable NT, and Green DJ.**  
701 Brachial artery blood flow responses to different modalities of lower limb exercise. *Med Sci*  
702 *Sports Exerc* 41: 1072-1079, 2009.
- 703 65. **Thijssen DH, Dawson EA, Black MA, Hopman MT, Cable NT, and Green DJ.**  
704 Heterogeneity in conduit artery function in humans: impact of arterial size. *American journal of*  
705 *physiology Heart and circulatory physiology* 295: H1927-1934, 2008.
- 706 66. **Thijssen DH, Dawson EA, Tinken TM, Cable NT, and Green DJ.** Retrograde flow  
707 and shear rate acutely impair endothelial function in humans. *Hypertension* 53: 986-992, 2009.
- 708 67. **Thijssen DH, Schreuder TH, Newcomer SW, Laughlin MH, Hopman MT, and**  
709 **Green DJ.** Impact of 2-Weeks Continuous Increase in Retrograde Shear Stress on Brachial  
710 Artery Vasomotor Function in Young and Older Men. *Journal of the American Heart*  
711 *Association* 4: 2015.
- 712 68. **Tinken TM, Thijssen DH, Black MA, Cable NT, and Green DJ.** Time course of  
713 change in vasodilator function and capacity in response to exercise training in humans. *J Physiol*  
714 586: 5003-5012, 2008.
- 715 69. **Tinken TM, Thijssen DH, Hopkins N, Black MA, Dawson EA, Minson CT,**  
716 **Newcomer SC, Laughlin MH, Cable NT, and Green DJ.** Impact of shear rate modulation on  
717 vascular function in humans. *Hypertension* 54: 278-285, 2009.
- 718 70. **Vogel RA, Corretti MC, and Plotnick GD.** Effect of a single high-fat meal on  
719 endothelial function in healthy subjects. *The American journal of cardiology* 79: 350-354, 1997.
- 720 71. **Walker AE, Kaplon RE, Pierce GL, Nowlan MJ, and Seals DR.** Prevention of age-  
721 related endothelial dysfunction by habitual aerobic exercise in healthy humans: possible role of  
722 nuclear factor kappaB. *Clinical science (London, England : 1979)* 127: 645-654, 2014.
- 723 72. **Welsch MA, Dobrosielski DA, Arce-Esquivel AA, Wood RH, Ravussin E, Rowley C,**  
724 **and Jazwinski SM.** The association between flow-mediated dilation and physical function in  
725 older men. *Med Sci Sports Exerc* 40: 1237-1243, 2008.
- 726 73. **Woodman RJ, Playford DA, Watts GF, Cheetham C, Reed C, Taylor RR, Puddey**  
727 **IB, Beilin LJ, Burke V, Mori TA, and Green D.** Improved analysis of brachial artery  
728 ultrasound using a novel edge-detection software system. *Journal of Applied Physiology* 91: 929-  
729 937, 2001.

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732 **Tables**

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734 **Table 1. Participant characteristics.**

735 Data are presented as mean±SD. Significance value  $P \leq 0.05$ . CRF, cardiorespiratory fitness; BMI, body mass index;  
 736 SBP, systolic blood pressure; DBP, diastolic blood pressure;  $VO_{2peak}$ , peak oxygen uptake; RER, respiratory  
 737 exchange ratio

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739 **Table 2. Comparison of baseline FMD indices between testing visits.**

740 Data are presented as mean±SD. Significance value  $P \leq 0.05$ . FMD, flow-mediated dilation;  $SRAUC$ , shear rate area-  
 741 under-the-curve.

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743 **Table 3. Flow-mediated dilation and hemodynamic indices at rest, 10 min and 60 min**  
 744 **following control or acute exercise in lower-fit elderly.**

745 Data are presented as mean±SD for a) lower-fit and b) higher-fit. Significance value  $P \leq 0.05$ . A fitness x time x  
 746 condition significant interaction was observed for FMDmm ( $P=0.04$ ), FMD% ( $P=0.01$ ) and 'adjusted  
 747 FMD%' ( $P=0.04$ ). For clarity, post-hoc  $P$  values are reported in the text only. \*significantly different to baseline  
 748 #significantly different to control °significantly different between moderate- and high-intensity. FMD; flow-mediated  
 749 dilation;  $SRAUC$ , shear rate area-under-the-curve; TTP, time-to-peak diameter; SBP, systolic blood pressure; DBP,  
 750 diastolic blood pressure; MAP, mean arterial pressure.

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752 **Figure**

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754 **Figure 1. Delta FMD % from baseline at a) 10-minutes post and b) 60-minutes post in**  
 755 **control, moderate-intensity and high-intensity exercise in both lower-fit (open-bars) and**  
 756 **higher-fit (dark bars) elderly individuals.**

757 Error bars represent SD. Significance value  $P \leq 0.05$ . Post hoc analysis revealed <sup>a</sup> control 60-min  $\Delta$ FMD% was  
 758 significantly reduced compared to exercise ( $P=0.01$ ), <sup>b</sup>  $\Delta$ FMD% significantly increased 10-min after moderate-  
 759 intensity compared to high-intensity exercise ( $P=0.02$ ), <sup>c</sup>  $\Delta$ FMD% significantly improved in the higher-fit compared  
 760 to the lower-fit group 60-min after high-intensity exercise ( $P=0.01$ ). FMD, Flow-mediated dilation.

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<b>Table 1.</b>	<b>All (n=47)</b>	<b>Lower-CRF (n=27)</b>	<b>Higher CRF (n=20)</b>	<b>P value (lower vs. higher)</b>
<b>Demographics</b>				
Age (years)	70±5	72±5	69±5	0.05
Hypertensive (%)	31	29	26	-
<b>Anthropometric measurements</b>				
Height (m)	1.74±0.08	1.72±0.08	176±0.09	0.27
Weight (kg)	76.4±11.5	76.3±12.5	76.5±10.3	0.96
BMI (kg.m <sup>-2</sup> )	25.3±3.4	25.5±3.4	24.9±3.3	0.52
Body fat (%)	24.7±5.9	25.8±6.0	23.3±5.8	0.17
Waist:Hip ratio	0.92±0.08	0.92±0.08	0.92±0.07	0.71
<b>Hemodynamic variables</b>				
Resting heart rate (bpm)	55±7	58±7	52±7	0.005
Brachial SBP (mm Hg)	125±15	124±14	126±12	0.66
Brachial DBP (mm Hg)	72±8	72±9	72±7	0.87
<b>Medication classification</b>				
ARB / ACE inhibitors (%)	23	22	19	-
Antiplatelets (%)	6	7	4	-
Beta-blockers (%)	4	7	0	-
Calcium channel blockers (%)	11	7	11	-
Statins (%)	30	40	11	-
<b>Cardiorespiratory fitness</b>				
VO <sub>2</sub> peak : Absolute (L.min <sup>-1</sup> )	2.22±0.63	1.85±0.39	2.71±0.56	<0.001
Relative (mL.kg <sup>-1</sup> .min <sup>-1</sup> )	29.0±6.96	24.3±2.9	35.4±5.5	<0.001
Peak heart rate (bpm)	151±15	146±15	156±10	0.02
Age-predicted (%)	100±10	102±12	97±6	0.08
RER (AU)	1.18±0.11	1.19±0.13	1.16±0.08	0.16
Peak Power (Watts)	160±40	140±30	190±40	<0.001



<b>Table 2.</b>	<b>CONTROL</b>	<b>MODERATE- INTENSITY</b>	<b>HIGH- INTENSITY</b>	<b><i>P</i> value (condition)</b>
<b>Baseline FMD test</b>				
<b>Diameter (mm)</b>	4.82±0.62	4.81±0.66	4.81±0.58	0.79
<b>FMD (mm)</b>	0.02±0.01	0.02±0.01	0.02±0.01	0.32
<b>FMD (%)</b>	4.71±1.57	4.86±1.58	4.89±1.45	0.50
<b>FMD SR<sub>AUC</sub> (10<sup>3</sup> s<sup>-1</sup>)</b>	13.8±5.7	13.7±7.6	14.6±7.1	0.29

3 a) LOW-FIT	CONTROL (NO EXERCISE)			MODERATE-INTENSITY CONTINUOUS EXERCISE			HIGH-INTENSITY INTERVAL EXERCISE		
	Pre	Post	Post (60 min)	Pre	Post	Post (60 min)	Pre	Post	Post (60 min)
<b>Flow-mediated dilation</b>									
<b>Diameter (mm)</b>	4.6±0.6	4.6±0.6	4.5±0.6*	4.6±0.6	4.7±0.6* <sup>#</sup>	4.6±0.6	4.6±0.6	4.7±0.6* <sup>#</sup>	4.6±0.7
<b>FMD (mm)</b>	0.02±0.01	0.02±0.01	0.02±0.01	0.02±0.01	0.03±0.01* <sup>#a</sup>	0.02±0.01	0.02±0.01	0.02±0.01	0.02±0.01
<b>Rest blood flow (mL.s<sup>-1</sup>)</b>	1.2±0.7	1.2±0.6	0.8±0.7*	1.2±0.6	1.8±0.9*	0.8±0.6	1.2±0.7	2.1±1.4* <sup>#</sup>	0.9±0.6
<b>Peak blood flow (mL.s<sup>-1</sup>)</b>	4.8±2.2	4.5±2.3	4.0±2.6*	4.8±2.0	5.5±2.1* <sup>#</sup>	4.7±2.6	5.2±2.8	6.0±2.5* <sup>#a</sup>	4.9±2.8
<b>FMD SR<sub>AUC</sub> (10<sup>3</sup> s<sup>-1</sup>)</b>	14.1±5.9	13.4±7.4	13.3±6.5*	15.0±8.2	17.6±8.1* <sup>#</sup>	14.7±8.0	15.5±7.0	18.3±7.6* <sup>#a</sup>	15.0±7.9
<b>TTP diameter (s)</b>	66±27	67±35	74±36*	72±31	64±27	73±46	69±34	71±32	67±40
<b>FMD (%)</b>	4.7±1.6	4.4±1.7	4.1±1.6*	4.7±1.6	5.4±1.9* <sup>#</sup>	4.8±1.7	4.8±1.4	4.0±2.2* <sup>#a</sup>	4.1±1.3* <sup>a</sup>
<b>Adjusted FMD (%)</b>	4.5±1.6	4.2±1.5	4.0±4.6*	4.5±1.9	5.1±1.7* <sup>#</sup>	4.5±1.7	4.9±1.4	3.9±2.1* <sup>#a</sup>	4.2±1.2* <sup>a</sup>
<b>Heart rate and blood pressure</b>									
<b>Heart rate (bpm)</b>	59±10	56±8	55±7	58±7	68±9*	58±6	58±8	71±13* <sup>#a</sup>	59±8
<b>SBP (mm Hg)</b>	124±15	130±15	129±15	125±14	133±13*	126±15	124±12	132±14*	124±11
<b>DBP (mm Hg)</b>	72±9	76±9	74±9	73±9	75±9	74±11	73±9	76±10	74±9
<b>MAP (mm Hg)</b>	87±8	91±9	90±9	88±10	93±9*	89±12	88±10	93±11*	88±9

3 b) HIGH-FIT	CONTROL (NO-EXERCISE)			MODERATE-INTENSITY CONTINUOUS EXERCISE			HIGH-INTENSITY INTERVAL EXERCISE		
	Pre	Post	Post (60 min)	Pre	Post	Post (60 min)	Pre	Post	Post (60 min)
<b>Flow-mediated dilation</b>									
Diameter (mm)	5.0±0.7	4.9±0.6	5.0±0.6	5.0±0.7	5.1±0.7* <sup>#</sup>	5.0±0.6	4.9±0.5	5.1±0.6* <sup>#</sup>	5.0±0.6
FMD (mm)	0.02±0.01	0.02±0.01	0.02±0.01	0.02±0.01	0.03±0.01* <sup>#a</sup>	0.02±0.01	0.02±0.01	0.02±0.01	0.03±0.01* <sup>#a</sup>
Rest blood flow (mL.s <sup>-1</sup> )	1.1±0.9	0.9±0.6	0.7±0.6*	1.2±0.9	1.9±1.0* <sup>#</sup>	1.0±0.8	1.2±0.9	2.2±1.1* <sup>#a</sup>	1.0±0.6
Peak blood flow (mL.s <sup>-1</sup> )	5.0±2.7	4.4±2.7	3.5±1.9*	4.7±2.6	5.1±2.4* <sup>#</sup>	4.9±2.0	5.0±2.9	6.2±1.9* <sup>#a</sup>	4.7±2.2
FMD SR <sub>AUC</sub> (10 <sup>3</sup> s <sup>-1</sup> )	10.2±5.6	10.1±5.9	9.3±5.6*	11.6±6.5	13.7±7.3* <sup>#</sup>	12.0±3.5	13.2±7.1	15.5±7.3* <sup>#a</sup>	12.7±5.2
TTP diameter (s)	57±24	61±26	69±33*	60±21	54±18	56±23	62±32	58±32	58±27
FMD %	4.8±1.6	4.4±1.0	4.1±1.3	5.1±1.5	6.1±2.5* <sup>#a</sup>	4.9±1.3	4.9±1.5	5.0±2.6	5.7±2.0* <sup>#a</sup>
Adjusted FMD (%)	4.6±1.4	4.4±1.1	3.8±1.6	5.0±1.6	5.9±2.0* <sup>#a</sup>	4.6±1.6	4.9±1.4	4.8±2.3	5.5±1.6* <sup>#a</sup>
<b>Heart rate and blood pressure</b>									
Heart rate (bpm)	51±7	48±6	49±8	52±7	61±8*	52±6	52±7	64±7* <sup>#a</sup>	53±6
SBP (mm Hg)	126±12	133±13	132±12	127±12	136±11*	125±13	126±10	135±12*	125±13
DBP (mm Hg)	72±7	75±8	75±8	72±7	76±7	72±8	73±9	76±7	72±8
MAP (mm Hg)	87±7	90±8	89±8	88±8	93±8*	86±10	87±6	94±7*	87±8

Fitness x time  $P = 0.37$

Condition x fitness  $P = 0.04$

Condition x time  $P < 0.01$

Fitness x condition x time  $P = 0.01$

