

Whole body passive heating versus dynamic lower body exercise: A comparison of peripheral hemodynamic profiles

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- 1 Whole body passive heating versus dynamic lower body exercise: A comparison of peripheral
- 2 hemodynamic profiles
- 3
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25 New & Noteworthy

26

- **27** Passive heating and exercise increase blood flow through arteries generating a frictional force,
- 28 termed shear rate, which is associated with positive vascular health. Few studies have compared
- 29 the increase in arterial blood flow and shear rate elicited by passive heating to dynamic
- 30 continuous exercise. We found thirty minutes of whole-body passive hot water immersion (42 °C
- bath) increased femoral artery blood flow and shear rate equivalent to exercising at a moderate
- **32** intensity (\sim 57% HR_{max}).

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34

35 <u>ABSTRACT</u>

36

37 Passive heating has emerged as a therapeutic intervention for the treatment and prevention of 38 cardiovascular disease. Like exercise, heating increases peripheral artery blood flow and shear 39 rate which is thought to be a primary mechanism underpinning endothelium mediated vascular 40 adaptation. However, few studies have compared the increase in arterial blood flow and shear rate between dynamic exercise and passive heating. In a fixed crossover design study, 15 41 42 moderately trained healthy participants $(25.6 \pm 3.4 \text{ years})$ (5 female) underwent 30 minutes of 43 whole body passive heating (42 °C bath), followed on a separate day by 30 minutes of semirecumbent stepping exercise performed at two workloads corresponding to the increase in cardiac 44 output (Qc) (Δ 3.72 l·min⁻¹) and heart rate (HR) (Δ 40 bpm) recorded at the end of passive 45 heating. Results: At the same Qc (Δ 3.72 l·min⁻¹ vs 3.78 l·min⁻¹), femoral artery blood flow (1599 46 ml/min vs 1947 ml/min) (p=0.596) and shear rate (162 s⁻¹ vs 192 s⁻¹) (p=0.471) measured by 47 48 ultrasonography were similar between passive heating and stepping exercise. However, for the same HR matched intensity, femoral blood flow (1599 ml·min⁻¹ vs 2588 ml·min⁻¹) and shear rate 49 (161s⁻¹ vs 271s⁻¹) were significantly greater during exercise, compared with heating (both 50 51 P=<0.001). The results indicate that, for moderately trained individuals, passive heating increases 52 common femoral artery blood flow and shear rate similar to low intensity continuous dynamic 53 exercise (29% VO_{2max}), however exercise performed at a higher intensity (53% VO_{2max}) results in significantly larger shear rates towards the active skeletal muscle. 54 55 Key Words: passive heating, dynamic exercise, leg blood flow, ultrasound, shear rate 56 57 58 59 60 61

62

63 INTRODUCTION

64

65 Cardiovascular disease (CVD) is the leading cause of death worldwide. In 2016, the worldwide mortality rate for CVD was $\sim 31\%$, a figure greater than the nine other leading causes 66 67 of death combined (46). Exercise is often cited as the most effective non-pharmacological intervention for the prevention and management of CVD (29). The protective effects of exercise 68 69 on vascular structure and function are partly mediated by the frictional force generated between 70 the endothelium and increased blood flow (shear), resulting in nitric oxide (NO) dependent 71 endothelial vasodilation (16). Unfortunately, a large majority of the global population fails to 72 adhere to the recommended guidelines for physical activity (45), which has led to the 73 unprecedented rise in global obesity and consequently, CVD (4). Therefore, alternative therapies 74 which reduce the risk of CVD are widely sought. 75 Passive heating has emerged as a potential therapy for CVD with the notion it may induce 76 vascular adaptations comparable to exercise (3). Indeed, Brunt et al. (2016) reported 77 improvements in brachial artery flow mediated dilation, and superficial femoral artery (SFA) 78 compliance and stiffness, following 8 weeks of passive hot water immersion in young healthy 79 sedentary volunteers. Additionally, Romero et al. (2017) reported improvements in macro- and 80 microvascular function in healthy elderly individuals following lower limb hot water immersion; 81 and, Bailey et al. (2016) reported improvements in brachial flow mediated dilatation (~1.71%), as well as a reduction in heat induced hypoperfusion in healthy females following 8 weeks of 82 83 passive lower body hot water immersion (42°C). These benefits, observed in healthy volunteers, are now being translated into clinical populations. Neff et al. (2016) reported a reduction in mean 84 85 arterial pressure in peripheral artery disease (PAD) patients following leg thermotherapy via a water perused suit (48°C). Similarly, Thomas et al. (2017) reported decreases in blood pressure, 86 increases in popliteal antegrade shear rate, and reductions in both central and peripheral pulse 87 88 wave velocity in PAD patients following 30 minutes of passive lower limb hot water immersion 89 (42°C). 90 Although early evidence demonstrates the therapeutic benefits of passive heating, a key question 91 will be to what extent passive heating provides benefits relative to the most effective non-

92 pharmacological intervention for CVD, exercise. Mechanistically, heating increases cutaneous

93 blood flow to support heat dissipation for maintenance of temperature homeostasis, with exercise

94 increasing skeletal muscle perfusion to meet the metabolic demands of movement. Thus, both

95 interventions rely on increasing conduit artery blood flow and shear rate, which have been 96 established as a primary driver mediating vascular adaptation (42). Yet it is not entirely clear what intensity of dynamic exercise should be performed to match vascular shear to passive 97 heating, which must be ascertained before studies can be designed to directly compare chronic 98 99 passive heating with exercise training. To our knowledge, only one study has directly compared 100 local blood flow and shear rate responses between exercise and passive heating. Thomas et al. (2016) reported a \sim 232% (181 s⁻¹) increase in superficial femoral artery (SFA) shear rate 101 following 30 minutes of passive heating, compared with only a $\sim 146\%$ (104 s⁻¹) increase 102 103 following 30 minutes of treadmill running at \sim 65% of maximum heart rate. These results suggest 104 whole body passive heating may provide a greater vascular shear stimulus compared with exercise. However, as acknowledged by the authors, shear rates within the SFA i.e. the "active" 105 106 skeletal muscle, were likely severely underestimated in the exercise condition, as shear was 107 quantified ~5-10 minutes after each trial. After passive heating, core and skin temperature remain 108 elevated with a persistent reduction in downstream resistance maintaining conduit artery blood 109 flow and shear rate (37). Whereas after exercise, the rapid reduction in oxygen demand causes a near instantaneous reduction in perfusion due to tight metabolic flow coupling. 110

While not providing a direct comparison to passive heating, several studies suggest that shear rates towards active skeletal muscle during exercise (e.g. forearm during handgrip) (14), (quadricep during leg kicking) (33), (quadricep during leg kicking) (43; 44), maybe similar, or substantially higher than values reported for passive heating (28), (35). Indeed, Dawson et al. (2017) reported a mean brachial shear value of 283 s⁻¹ during 30 minutes of cycling at 80% maximum heart rate and Padilla et al. (2011) found similar values of 260 s⁻¹ after 60 minutes semi-recumbent cycling at 120 watts.

118 Therefore, the aim of this study was to directly compare both brachial and femoral artery 119 blood flow and shear rate during passive hot water immersion with dynamic lower body exercise 120 performed at a matched cardiovascular demand. To account for the different cardiac responses 121 between heating (heart rate only) and exercise (stroke volume and heart rate), we compared 122 exercise at two clamped workloads, which corresponded to 1) the increase in cardiac output (Qc) 123 and 2) increase in heart rate (HR) recorded at the end of whole body heating. We hypothesized 124 that increases in femoral blood flow and shear rate would be similar when Oc during exercise 125 was matched to heating, but significantly lower when HR during exercise was matched to heating. As Qc typically increases by 3-4 liters during whole body heating (13; 8), primarily 126

127	mediated by an increase in HR approximately 40 beats min ⁻¹ (13;12), we hypothesized that this
128	would equate to approximately ~45-50% of maximum HR (~195 - 200 bpm) during running
129	exercise in young healthy population. Finally, brachial artery blood flow and shear rates would
130	be significantly higher after 30 minutes of heating compared with both Qc and HR matched
131	exercise, due to exercise induced vasoconstriction in non-active skeletal muscle (18).
132	
133	METHODS
134	
135	Ethical Approval
136	Written informed consent was provided from all participants following detailed verbal
137	explanations of the experimental protocol which included information regarding all potential
138	risks. The study conformed to the standards set out by the Declaration of Helsinki, except for
139	registration in a database, and was approved by the ethics committee of the University of
140	Innsbruck.
141	
142	Participants

143 Ten male and five female participants $(25.6 \pm 3.4 \text{ years, height, } 1.76 \pm 0.1 \text{ m, weight, } 73.6 \pm 9.3 \text{ m})$

144 kg; VO_{2max} , $54.4 \pm 7.8 \text{ ml} \cdot \text{kg} \cdot \text{min}^{-1}$) were recruited from the University of Innsbruck. All

145 participants were healthy, non-smokers and free from cardiovascular, metabolic and

146 neuromuscular diseases. Female participants were tested during the early follicular phase of their

147 cycle, this included participants taking oral contraceptives who were tested during the placebo

- **148** phase of their cycle (low hormone)
- 149

150 <u>Experimental Design</u>

All participants underwent two trials in a fixed crossover study design (Figure 1). Participants were instructed to abstain from strenuous exercise and avoid consumption of caffeine and alcohol for 24 hours prior to each testing day. We advised participants to keep diet consistent between trials and made every effort to perform the stepping test at the same time of day as the heating trial. The first trial involved 30 minutes of passive hot water immersion (PHWI) in a 42 °C bath. This was followed on a separate day by 30 minutes of graded semi-recumbent stepping exercise. All participants undertook the heating trial prior to the stepping trial. A fixed order was required

158 as the cardiovascular demand elicited by passive heating (Δ Qc and Δ HR) was used to target the

159 intensity of stepping exercise. The heating trial was undertaken in a physiology laboratory during

160 the winter months (October-January) and the exercise trial was performed during the summer

161 months (May-June) in an environmentally controlled (Ta, 22 °C, relative humidity 40%) chamber

162 (Küba Blue Line DE Professional, Kelvion Holding GmbH, Germany) at the University of

- **163** Innsbruck.
- 164

165 <u>Heating Trial</u>

166 Upon arrival, participants changed into swimwear (males shorts; females sport shorts and bra) 167 and self-inserted a rectal probe (DeRoyal, Powell, TN, USA) 15 cm past the anal sphincter for 168 monitoring of core temperature. Body weight was recorded (Kern DS 150k1, Kern & Sohn, 169 Germany) and participants positioned themselves on a semi-recumbent bed. After 170 instrumentation and 20 minutes of quiet rest, baseline continuous haemodynamic data were 171 recorded over one minute in a semi recumbent posture. HR was measured from a three-lead 172 electrocardiogram (Tram-rac, Solar 8000M, GE- Marquette, USA). Forearm skin temperature 173 and blood flow were obtained via an integrated thermistor and laser-Doppler flowmeter (Moor 174 Instruments, Devon, UK). Arterial pressure was measured, in duplicate, via electro-175 sphygmomanometry (Tango, SunTechMedical Instruments Inc., USA) with a microphone placed 176 over the brachial artery to detect Korotkoff sounds. Oc was measured and stroke volume 177 calculated via inert gas rebreathing (Innovision DK-5260, Denmark) (10), alongside a measure of 178 VO₂ Thereafter, brachial and common femoral artery blood flow and shear rate were measured 179 using ultrasound (see below). Following acquisition of baseline variables, participants were 180 transferred to a hot bath (42 °C) and immersed up to the height of the mid-sternum with both 181 arms rested at heart level outside the bath (Figure 1), identical to the semi-recumbent posture 182 during baseline measurements. After 30 minutes of heating, brachial blood flow, Qc and VO_2 183 were measured, and the bath was partially drained. Once the water drained to just below the iliac 184 crest, femoral blood flow was measured (2-3 mins) whilst the participant remained seated in the 185 bath.

186

187 <u>Stepping Exercise</u>

188 On a subsequent day, upon arrival to the environmental chamber, participants were positioned on189 the same semi-recumbent medical bed used for the heating trial. To enable participants to

190 perform stepping exercise in the same semi-recumbent position, a cardio-stepper (Ergospect

191 medical technology, Innsbruck, Austria) was custom-fixed onto the end of the bed (Figure 2). 192 Following instrumentation, which replicated the heating trial and included measurement of core 193 temperature via a rectal probe (DeRoyal, Powell, TN, USA), in addition to forearm skin 194 temperature and blood flow via an integrated thermistor and laser-Doppler flowmeter (Moor 195 Instruments, Devon, UK), baseline heamodynamic data were recorded following 20 minutes of 196 quiet semi-recumbent rest. This included, blood pressure measured in duplicate by electro-197 sphygmomanometry and Qc and VO₂ assessed by inert gas rebreathing, as during the heating 198 trial. Participants were then instructed to begin stepping exercise and intensity was manipulated 199 such that Qc increased to levels recorded at the end of passive heating. Based on pilot testing, the 200 workload was increased until heart rate was elevated by 20 beats per min, in order to account for 201 the exercise-induced increase in stroke volume that is not apparent with passive heating. 202 Thereafter Qc and VO_2 were determined via a inter-gas rebreathent to confirm the correct intensity 203 had been obtained (Qc matched) and the workload was adjusted if required. After five minutes of 204 steady-state exercise at the target workload, continuous haemodynamics and blood pressure 205 measurements were taken alongside simultaneous assessment of brachial and femoral artery 206 blood flow. Subsequently, exercise intensity was increased until HR reached the subject specific 207 target, that was determined from the value recorded at the end of the passive heating trial. 208 Exercise at this workload was maintained for 25 minutes at which a rebreathe was performed for 209 determination of Qc and VO₂ alongside measurements of blood pressure (in duplicate) and 210 brachial and femoral blood flow during exercise. To assess blood flow kinetics post-exercise, 211 participants were given a three-second countdown and told to stop exercising where upon a 212 support was placed immediately under the left leg to allow complete relaxation of the limb and 213 facilitate a five-minute continuous measure of femoral blood flow.

214

215 <u>Ultrasound</u>

Brachial and common femoral artery blood flow of the left arm and leg were measured using a 9MHz linear-array Doppler probe (iE33, Philips, Netherlands) by continuous duplex vascular
sonography (iE33, Philips, Netherlands). Arterial diameter was imaged using two-dimensional B
mode over 30 seconds and measured offline during diastole (in triplicate) by the same
investigator. Anatomical landmarks visible during B-mode measurements of diameter were noted
to ensure probe placement remained consistent between baseline and all subsequent recordings,

as well as between trials. Thereafter, the time average mean blood velocity (TAMV) was

recorded at an insonation angle of 60° for between 30-60 seconds and imported into Labchart via 223 224 a Doppler audio converter (Penn State, Hershey, Pennsylvania, USA) (Herr et al. 2010). Antegrade and retrograde blood flows were derived from the TAMV and recorded in separate 225 226 channels in labchart (see Figure 3 for an example of individual flow profiles). Ultrasound 227 assessments in both trials were conducted by the same investigators. Local arm and leg cooling 228 was applied to the skin (fan and wet towels) if diastolic blood flow appeared elevated during resting baseline measures in order to limit the effect of skin temperature and skin blood flow on 229 the assessment of skeletal muscle blood flow (23). 230

231

232 <u>Maximal Exercise Test</u>

All but two participants (due to unrelated injuries) completed a treadmill (HP cosmos, Pulsar,

234 Germany) maximal exercise test for determination of VO_{2max} . The test commenced at a speed of 8 **235** km·h⁻¹ with a ~1% incline. Each minute speed was increased by 1 km·h⁻¹ until 12 km·h⁻¹,

236 thereafter incline was increased by $\sim 1\%$ every 30 seconds until participants reached volitional

exhaustion. Breath-by-Breath gas analysis was continuously sampled using an open spirometric

238 system (Oxycon Pro, CareFusion GmbH, Hoechbach, Germany), which was calibrated prior to

each measurement according to the manufacturer's guidelines. HR was determined by chest belt

240 (Wear Link, Polar, Kempele, Finland) and transmitted to the spirometric device. VO_{2max} was

241 defined as the highest 30s average in oxygen uptake and maximal heart rate (HR_{max}) as the

242 highest 10s average during the test.

243 A previous study by Bachler et al. (2017) compared VO_{2max} between treadmill running and the

244 cardio stepper used in the current study, whereby VO_{2max} was ~23% higher on the treadmill in

similarly trained participants (54.4 ml·kg·min⁻¹ versus 54.7 ml·kg·min⁻¹). Therefore an estimated

246 stepping VO2max of 42.1 ml·kg·min⁻¹ was used to determine the percentage workload during Qc

247 and HR matched intensities while stepping.

248 <u>Data analysis</u>

All continuous measurements were sampled at 250 Hz (Powerlab, ADInstruments, Oxford, UK)

250 and analyzed via an offline data acquisition system (LabChart 8; AD Instruments; Oxford, UK).

251 Brachial and femoral artery blood flow were expressed in ml min⁻¹ using the equation below

252 (equation 1), where TAMV was recorded in cm s^{-1} , vessel diameters in cm and 60 was used to

253 convert from $ml \cdot s^{-1}$ to $ml \cdot min^{-1}$. Antegrade and retrograde blood flow were also derived using the

same equation. Mean arterial pressure (MAP) was calculated from systolic and diastolic values

- **255** from automated measures obtained via electro sphygmomanometer (equation 2). Total blood
- **256** vessel shear rate, including antegrade and retrograde shear were calculated and expressed in s^{-1} .
- **257** Oscillatory shear index (OSI), which represents the direction and magnitude of flow between
- **258** systole and diastole was also calculated. Values range between 0 (no oscillations) to 0.5 (high
- - **259** oscillations).

Equation 1: Blood flow = TAMV x $\pi \left(\frac{\operatorname{artery \, diameter \, (mm)}}{2}\right)^2 x 60$ Equation 2: MAP = DB + $\frac{1}{3} \times (SB - DP)$

Equations 3: shear rate = $4 \times (\frac{TAMV}{diameter})$

Equation 4: $OSI = \frac{retrograde shear}{(antegrade shear+retrograde shear)}$

260

261 <u>Statistical Analyses</u>

A total of 20 participants, consisting of eleven males and nine females undertook the heating trial.
One male volunteer was unable to complete the heating trial and was excluded from analysis. The
same ten males and five females undertook the stepping exercise trial. Four females dropped out
of the exercise trial, two due to external sporting injuries and two moved away from the area.
Therefore, a total of 15 participants took part in both the heating and stepping exercise trials and
were included in the analysis.

268 To identify the independent effect of passive heating, all outcome variables were compared between baseline rest and 30 minutes passive heating using either a paired samples t-269 270 test, or a Wilcoxon rank test. Comparisons for the exercise trial were made using a repeated 271 measures analysis of variance (ANOVA) with Bonferroni correction to determine changes 272 between baseline, the QC_{MATCHED} (5 mins) intensity and the HR_{MATCHED} intensity (30 mins). To 273 compare passive heating to exercise, change scores were calculated for each variable relative to 274 the appropriate baseline control value and compared using repeated measures ANOVA with 275 Bonferroni correction. To examine the time course of femoral blood flow offset kinetics post 276 exercise, a repeated measures ANOVA was performed with multiple Bonferroni corrections to 277 determine difference from the cessation of exercise (time point 0) until the end of the 278 measurement recording (300 seconds). All values are expressed as mean \pm standard deviation 279 with statistical significance set at $P \le 0.05$.

Statistical analysis including Levene test of homogeneity of variance was performed using 280

281 SPSS version 25 (SPSS Inc., IBM, Chicago, IL, USA) and Prism 8 (GraphPad Software Inc., La

- 282 Jolla, CA, USA).
- 283
- 284 RESULTS
- 285
- 286 Maximal Exercise Test
- Average VO_{2max} was $54.7 \pm 7.89 \text{ ml} \cdot \text{kg} \cdot \text{min}^{-1}$ with maximum HR recorded at $189 \pm 6 \text{ beats} \cdot \text{min}^{-1}$ 287 ¹. This equated to an estimated stepping VO_{2max} of 42.1 ml·kg·min⁻¹.
- 288
- 289 Passive hot water immersion
- 290 Thirty minutes of PHWI increased core temperature by 1.3 ± 0.37 °C (P ≤ 0.001) and Qc by $3.72 \pm$
- 1.9 l·min⁻¹ (P<0.001), which was entirely mediated by an increase in HR (38 ± 15 beats·min⁻¹) 291
- (P<0.001) as stroke volume (P=0.884) showed little change (Table 2). 292
- PHWI increased femoral artery blood flow by $1303 \pm 363 \text{ ml} \cdot \text{min}^{-1}$ (P<0.001) and 293
- brachial artery blood flow by $210 \pm 64 \text{ ml} \cdot \text{min}^{-1}$ (P<0.001). The increase in blood flow through 294
- both conduit arteries corresponded with increased antegrade shear (P<0.001) and decreased 295
- 296 retrograde shear (P<0.001) (Table 1).

297 Stepping exercise

- Stepping exercise was successfully matched to the two targeted intensities. The first exercise 298
- workload (Oc_{MATCHED}) matched the increase in Oc during PHWI (Ä PHWI:- Ä 3.72 vs 299
- Qc_{MATCHED};- Ä 3.78 l·min⁻¹, P≥0.999) and the second intensity (HR_{MATCHED}) matched the 300
- increase in HR recorded at the end of PHWI (Ä PHWI;- Ä 38 vs HR_{MATCHED:-} Ä 40 beats min⁻¹, 301
- P≥0.999). 302

Stepping exercise performed at the Qc_{MATCHED} intensity (absolute Qc, 11.41 l·min⁻¹) 303 increased femoral blood flow by $1470 \pm 464 \text{ ml} \cdot \text{min}^{-1}$ (P<0.001) which manifested to an increase 304 in femoral antegrade shear rate by \ddot{A} 144 ± 56 s⁻¹ (P<0.001) and decreased retrograde shear rate 305 by $\ddot{A} 7 \pm 4 \text{ s}^{-1}$ (P<0.001). When exercise was performed at the HR_{MATCHED} intensity (absolute 306 HR, 103 beats \cdot min⁻¹), femoral blood flow increased by Ä 2123 ± 524 ml min⁻¹ (P<0.001) with 307

antegrade shear rate increasing by $\ddot{A} 221 \pm 58 \text{ s}^{-1}$ (P<0.001) and retrograde shear rate decreasing by $\ddot{A} 8 \pm 4 \text{ s}^{-1}$ (P<0.001).

Brachial blood flow did not change during stepping at the Qc_{MATCHED} ($\ddot{A} \ 2 \pm 12 \ \text{ml} \cdot \text{min}^{-1}$, P=0.753), but both brachial antegrade shear rate ($\ddot{A} \ 32 \pm 22 \ \text{s}^{-1}$, P<0.001) and retrograde shear rate ($\ddot{A} \ 23 \pm 18 \ \text{s}^{-1}$, P<0.001) increased. Stepping exercise performed at the HR_{MATCHED} intensity increased brachial blood flow ($\ddot{A} \ 72 \pm 61 \ \text{ml} \cdot \text{min}^{-1}$, P≤0.001), with a further increase in antegrade shear rate ($\ddot{A} \ 119 \pm 78 \ \text{s}^{-1}$, P<0.001), but only a small increase in retrograde shear rate from baseline ($\ddot{A} \ 8 \pm 21 \ \text{s}^{-1}$, P<0.179).

316 <u>Comparison of passive hot water immersion versus stepping exercise</u>

317 The increase in femoral blood flow and total shear rate were similar after 30 minutes of PHWI compared to five-minutes stepping exercise performed at the QC_{MATCHED} intensity, however 318 319 femoral blood flow and shear rate were significantly greater during exercise performed at the HR_{MATCHED} intensity (both P<0.001, Figure 2A). Femoral antegrade shear rate was similar 320 between heating and exercise performed at the QC_{MATCHED} intensity $(161 \pm 58 \text{ s}^{-1} \text{ vs } 193 \pm 55 \text{ s}^{-1})$, 321 P=0.559), but was significantly greater when exercise was performed at the HR_{MATCHED} intensity 322 $(161 \pm 58 \text{ s}^{-1} \text{ vs } 270 \pm 59 \text{ s}^{-1}, P < 0.001)$. Femoral retrograde shear rate decreased to 0 s^{-1} (P < 0.001) 323 following 30 minutes of passive heating, which was almost identical to the shear rate recorded 324 after five minutes of exercise at the Qc matched exercise intensity $(-0.6 \pm 1.1 \text{ s}^{-1})$. Femoral 325 retrograde shear rate remained at a similar value when recorded at 30 minutes of exercise at the 326 $HR_{MATCHED}$ (-0.05 ± 0.09 s⁻¹) intensity. 327

328 In contrast to the femoral artery, brachial blood flow and total shear rate were 329 significantly higher after 30 minutes of PHWI compared with exercise performed at both the QCMATCHED and HRMATCHED intensities (both P<0.001, Figure 2C). Moreover, PHWI caused a 330 greater increase in brachial antegrade shear rate $(245 \pm 64 \text{ s}^{-1})$ than both the QC_{MATCHED} $(32 \pm 22 \text{ s}^{-1})$ 331 ⁻¹) and HR_{MATCHED} (119 \pm 78 s⁻¹) intensities (P<0.001) and caused a reduction in brachial 332 retrograde shear rate (\ddot{A} 10.7 s⁻¹). In contrast, stepping exercise performed at the QC_{MATCHED} 333 intensity resulted in an increase in retrograde shear rate ($\ddot{A} - 23 \pm 18 \text{ s}^{-1}$, P<0.001), which although 334 decreased by Ä 14.7 s⁻¹ after 25 minutes at the HR_{MATCHED} intensity, remained significantly 335 greater than the reduction caused by PHWI (PHWI, Ä 10.7 s⁻¹ vs HR_{MATCHED}, Ä -8.0 s⁻¹). 336

337 <u>*Time course of post exercise femoral shear rate.</u>*</u>

- **338** Peak femoral artery shear rate measured during exercise was 271 ± 59 s⁻¹. Thirty seconds after
- **339** stopping exercise, shear rate was significantly lower ($200 \pm 50 \text{ s}^{-1}$, P<0.001) and continued to
- decrease at each subsequent 30 second time point. At three minutes (180 sec), shear rate started
- **341** to plateau, but continued to decline slowly and was less than half the peak value when recorded
- **342** five minutes post exercise (99 \pm 35 s⁻¹, Figure 3B).
- 343

344 DISCUSSION

345 The main findings of the present study were that, 30 minutes of PHWI, which elicits a substantial increase in core temperature (+1.3 °C) in young moderately trained individuals, increased 346 347 femoral artery blood flow and altered shear patterns similar to those observed when performing 348 low to moderate intensity exercise (equivalent to approximately~29% stepping VO_{2max}). When 349 stepping exercise was performed at a higher HR_{MATCHED} intensity, (approximately ~53% stepping VO_{2max}) femoral blood flow and shear rate were substantially greater during exercise compared 350 351 with PHWI (both P<0.001, Figure 2). In contrast, brachial blood flow and total shear rate were 352 significantly higher after 30 minutes of PHWI compared with exercise performed at both the 353 Qc_{MATCHED} and HR_{MATCHED} intensities (both P<0.001, Figure 2). Thus, while sauna bathing and 354 other forms of heat therapy may be a beneficial strategy to improve vascular function and 355 cardiovascular risk factors (22), its recommendation should be presented with the caveat that 356 exercise may elicit substantially greater vascular hemodynamics in active limbs.

357

358 Whole body passive hot water immersion

359 Heat therapy is an emerging strategy that has been found to reduce vascular stiffness and 360 endothelial dysfunction in young healthy individuals (5) and consequently could potentially be 361 applied for treatment and prevention of CVD. Passive exposure to heat increases skin and core 362 temperature, decreases downstream vascular resistance and thus increases blood flow through 363 conduit arteries. As a result, shear forces are produced on the endothelium, which can trigger an 364 array of molecular pathways and alter both antiatherogenic and proatherogenic genes to favor 365 vascular health (24). Indeed, we found that sitting in a hot bath (42 °C) for 30 minutes increased skin (~430%) and rectal temperature (1.3 °C), decreased total peripheral resistance (~32%) and 366 367 substantially elevated blood flow through the common femoral (~442%) and brachial (~488%) arteries. Ultimately, this produced substantial shear rates on both arteries (common femoral, 161 368 \pm 59 s⁻¹; and brachial, 312 \pm 76 s⁻¹), with similar degrees of shear (126 s⁻¹) being observed in the 369 common femoral artery when core temperature was elevated to 38.4°C via a water perfused suit 370 (7). Others have also measured increases in arterial shear within the brachial (28, 260 s⁻¹), 371 superficial femoral (40, 265 s⁻¹) (33, 387 s⁻¹,) and popliteal (39, 89 s⁻¹), arteries during passive 372 heating. While large differences exist in the absolute shear rates between studies, they are most 373 374

likely due to the choice of artery and crucially the ultimate increase in skin and core temperature.

375 Indeed, Chiesa et al. (2016) have documented that a dose response relationship exists between leg376 temperature and blood flow through the common femoral and superficial femoral arteries.

377 Nonetheless, other potential differences include the interaction between postural unloading of the

378 baroreceptors and heightened sympathetic nerve activity (25) as well as hydrostatic forces due to

- **379** the level of water immersion between studies.
- 380

381 <u>Semi-recumbent stepping exercise</u>

Aerobic exercise is known to produce a range of vascular benefits including improved vessel 382 compliance, blood perfusion and increased NO bioavailability (17), all of which are suspected to 383 384 be mediated through an increase in vascular shear stress (16). However, despite these potential 385 benefits, relatively few studies have been able to simultaneously quantify shear rates in vascular 386 territories perfusing non-active and active skeletal muscle during continuous dynamic exercise 387 involving a large muscle mass. Quantifying blood flow and patterns of shear rate during this type 388 of exercise is important as it represents a more ecologically valid form of rhythmic exercise, 389 whereby stroke volume and arterial pulse pressure are increasing alongside a reduction in total 390 peripheral resistance. Similar to previous studies (28), we observed that short (5 minute) bouts of 391 mild intensity lower limb exercise caused minimal changes in brachial blood flow and total shear 392 rate (non-active skeletal muscle), but with prolonged (30 minutes) moderate intensity exercise, 393 brachial blood flow and total shear rate increased slightly. The reason blood flow is increased 394 towards non-active tissue with prolonged lower limb exercise is likely due to the observed 395 increase in downstream forearm vascular conductance, mediated by the slight increase in core 396 and skin temperature and dilation of the cutaneous circulation to aid thermoregulation (37).

397 To the best of our knowledge, the current study is the only investigation to measure 398 blood flow and shear rate in the common femoral artery (active skeletal muscle) during true 399 dynamic lower body exercise. Stepping for 25 minutes at the highest workload caused an increase in cardiac output to 14.8 l·min⁻¹ (heart rate 103 beats·min⁻¹; stroke volume 143 ml), a rise in 400 arterial pulse pressure (47 mmHg) and a decrease in total peripheral resistance (5.93 mmHg·ml⁻ 401 ¹·min⁻¹). Under the current experimental conditions, mean femoral blood flow and shear rates 402 were recorded at 2588 ml·min⁻¹ and 271 s⁻¹ respectively. Similar values for femoral blood flow 403 $(2480 \text{ ml}\cdot\text{min}^{-1})$ and shear rates (254 s^{-1}) have been observed during ~3 minutes of progressive 404 rhythmic knee extension exercise at similar cardiovascular workloads (43; 44). Together, these 405

406 findings demonstrate that dynamic leg exercise is a profound stimulus to elevate vascular shear407 stress towards the active tissue.

408

409 <u>Comparison of passive hot water immersion with stepping exercise</u>

410 As passive hot water immersion has emerged as a potential non-pharmacological therapeutic 411 strategy to increase shear rate and thus improve vascular function, it is important to ascertain its 412 potential, relative to other interventions such as exercise. In order to contextualize the shear rates 413 during passive heating, we compared passive heating to exercise at an intensity matched to the 414 increase in Qc and HR measured at the end of PHWI (i.e. matched for cardiovascular demand). 415 Both PHWI and exercise reduced retrograde blood flow and retrograde shear rate in the femoral 416 artery (i.e. in active skeletal muscle) to almost zero in all conditions. However, PHWI only 417 produced an increase in antegrade blood flow and shear rate similar to the Qc_{MATCHED} trial, which was equivalent to low intensity running exercise (approximately ~27% VO_{2max}). Indeed, stepping 418 performed at the HR_{MATCHED} intensity (approximately ~45% running VO_{2max}) produced a ~77% 419 420 greater increase in femoral antegrade shear rate compared with PHWI. These data have three 421 important implications. First, they outline the importance of measurement timing by contrasting the findings reported by Thomas et al. (2016) in which higher shear rates were observed in the 422 SFA following 30 minutes of passive heating (259 s^{-1}) compared with 30 minutes of treadmill 423 running (175 s⁻¹). Thomas et al. (2016) measured femoral blood flow 5-10 minutes after exercise, 424 425 which likely resulted in a substantial underestimation of exercise blood flow and shear rate due to the rapid reduction in blood flow following cessation of muscle contraction (30). Indeed, we 426 demonstrated that peak shear rate decreased by 71 s⁻¹, within as little as 30 seconds post exercise 427 and was reduced by 172 s⁻¹ when measured five minutes after exercise (Figure 3B). 428

429 Second, the first five minutes of exercise performed at the QCMATCHED intensity did not 430 increase core temperature, yet increased femoral flow and shear rate equivalent to 30 minutes of 431 passive heating, thus demonstrating the effectiveness of exercise hyperemia at increasing shear 432 rate in response to muscle contraction. Had the Qc_{MATCHED} intensity been extended to 30 minutes, 433 core temperature may have increased slightly, but likely contributing minimally to femoral blood 434 flow relative to the exercise hyperemia. Indeed, 30 minutes of stepping exercise at almost twice 435 the intensity (HR_{MATCHED}) only increased core temperature by 0.39 °C, which was far lower than 436 PHWI (+1.31 °C), suggesting low intensity exercise may be a more tolerable intervention for

providing a vascular shear stimulus. In this regard, emphasis should be placed on the potential of 437 438 dynamic exercise to increase vascular shear. Studies using the thermodilution technique have reported leg blood flow values of ~5.81 l·min⁻¹ (35), ~5.57 l·min⁻¹ (21), ~5.58 l·min⁻¹ (1), ~8.0 439 $1 \cdot \min^{-1}$ (26) and ~8.8 $1 \cdot \min^{-1}$ (36) during leg extensor exercise of various workloads. Moreover, 440 single leg blood flow values of ~9.10 1 min^{-1} (6) and ~12.52 1 min^{-1} (14) have been measured 441 442 during maximal diagonal striding (cross-country skiing) and cycling exercise respectively. As retrograde blood flow and thus velocity through the common femoral artery is minimal during 443 444 rhythmic exercise (see table 2, although likely higher during muscle contraction with resistance 445 type exercise such as leg kicking), a reasonable estimation of vascular shear rate can be 446 recalculated from these blood flow values, assuming a common femoral artery diameter of 0.958 447 cm for modestly trained individuals (current study) and a diameter of 1.053 cm for elite level 448 athletes (20) (Figure 8). Using these values, vascular shear appears to be linearly related to workload and substantially greater values can be achieved compared to passive heating. 449 Moreover, common femoral artery blood flow is increased up to $1.24 \, \text{l}\cdot\text{min}^{-1}(7)$ when core 450 451 temperature is elevated by 1.5 °C, yet no further increases can be obtained despite core temperature being elevated by 2.0 °C. Thus, it appears that passive heating has a celling effect for 452 453 increasing vascular shear in the femoral artery somewhere comparable to exercising at a mild to 454 moderate intensity (Figure 4).

Third, in contrast to the femoral artery, PHWI did provide a larger shear stimulus in the brachial artery compared to exercise, suggesting that passive heating increases flow and shear more globally to vascular beds throughout the body than exercise. This observation suggests that passive heat therapy may theoretically show greater improvements in brachial artery function (i.e. flow mediated dilation) compared to moderate lower body exercise training. However, for a more accurate comparison, passive heat therapy should be compared to longer durations of intense exercise where both core and forearm skin temperatures would be substantially elevated.

462

463 <u>Clinical Implications</u>

In terms of vascular shear, our data quantifies that being exposed to a high degree of whole body
heat stress (+1.3°C) is comparable to low intensity lower body exercise (approx. (~27% running
VO_{2max}) in young healthy individuals. These data should help guide future studies aiming to
compare chronic passive heating with exercise training, alongside future studies aiming to define
and prescribe suitable doses of heating and exercise training in clinical populations. Additionally,

the data place passive heating into context with exercise and may help the general populationunderstand the powerful benefits of even low intensity exercise.

471 That being said, we fully accept that passive heating provides a whole-body shear stimulus the 472 equivalent of low intensity exercise, supporting its application in clinical and elderly populations 473 who are limited in their capacity to exercise. Indeed, Thomas et al. (2017) demonstrated 474 comparable increases in popliteal shear rates following 3 minutes of treadmill walking at 3km/h 475 with a $\sim 10\%$ incline (standard test used for diagnosis of PAD) and 30 minutes of lower limb 476 heating in PAD patients. Critically, these patients report claudication during short bouts of 477 walking, (10 mins) making heating a much more suitable intervention that provides a comparable 478 shear stimulus to exercise. Furthermore, heating can be endured for a prolonged period compared 479 with exercise in this population, thereby potentially providing greater vascular adaptations.

480 *Limitations*

481 During the exercise trial, we successfully matched both exercise intensities to the intended 482 cardiovascular loads of passive heating. However, we did not match the duration of each 483 workload relative to the time spent under passive heating. For example, the QC_{MATCHED} intensity 484 was only measured after 5 minutes. While this may limit our interpretation slightly, it is unlikely 485 that exercising at such a relative low steady-state workload ($\sim 27\%$ running VO_{2max}) for a further 486 25 minutes would have substantially altered the physiological response we observed after 5 minutes. Furthermore, we did not match core temperature changes between trials, which would 487 488 separate differences in heat induced increases in blood flow from exercise. Another limitation is 489 that we did not perform repeat measures of shear rate in the brachial and femoral artery post 490 heating. After passive heating, core and skin temperature remain elevated with a persistent 491 reduction in downstream vascular resistance, which is in opposition to acute mild to moderate 492 intensity exercise. Therefore, to precisely compare passive heating with exercise, several hours of 493 shear need to be recorded during and after varying degrees of heat stress and exercise at various 494 intensities/durations. In addition, we did not assess thermal comfort which would provide support 495 for out statement regarding the tolerability of passive heating compared with exercise. However, 496 we based this statement on the substantially elevated core and skin temperatures which are 497 associated with thermal comfort (11). Furthermore, we referred to participants performing 498 stepping exercise at 27 % (Qc_{MATCHED}) and 45% (HR_{MATCHED}) of their running VO₂ max, 499 however this was an estimate based on the comparison of VO₂ max scores between treadmill

500 running, cycle ergometry and cardio-stepping (Bachlet et al. 2017). Whilst we did not measure 501 VO₂ max on our stepper, the paper referenced used young healthy participants who were well matched in terms of cardiorespiratory fitness (treadmill VO₂ max 54.4 ml·kg·min⁻¹) compared 502 with out participants (54.7 ml·kg·min⁻¹), therefore we felt justified in making this comparison but 503 504 should emphasise that these values were estimates. Finally, we acknowledge the time gap 505 between trials (4 months) may have biased our measurements due to potential changes in lifestyle 506 factors or training habits. The time gap was due to logistically issues concerning the location and 507 transportation of expensive equipment between laboratories. However, all participants were 508 instructed to maintain their normal day-to-day training habits between trials, with emphasis 509 placed on replicating their diet and daily activities 24 hours before testing sessions. We also assessed baseline parameters before both trials and found similar values for Qc, HR and stroke 510 511 volume (Table 3) providing some evidence that training status remained relatively consistent 512 between trials. Moreover, our measurements of blood flow and shear rates at rest and during 513 exercise are in line with previous literature, suggesting these effects are likely minimal (32). 514 Finally, our study was conducted in a young healthy population and we acknowledge that the 515 practical application of many passive heating interventions are targeted at elderly or clinical 516 populations unable to exercise, consequently future studies should focus on such groups to 517 determine whether these populations have similar increases in blood flow and shear rate under 518 matched conditions.

519

520 <u>Conclusion</u>

These findings suggest that whilst whole body heating provides a shear rate stimulus that matches
low intensity exercise, it may be more challenging to endure compared with time matched
exercise owing to the greater increase in core temperature (1.31 ± 0.37 vs 0.39 ± 0.19 °C).
Furthermore, if higher exercise intensities can be performed, exercise provides a substantially
greater shear stimulus toward the active skeletal muscle, which is likely to confer superior
vascular adaptations in young healthy individuals.

527

528

529 ADDITONAL INFORMATION

530 *Competing Interests*

531	None of the	authors have any	conflicts of interests.
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- *Author Contributions*
- 534 Conception/design of the work: JSL, WKC III, JPM, SAR. Acquisition/analysis of data for the
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552 <u>REFERENCES</u>

- 553 1. Andersen P, and Saltin B. Maximal perfusion of skeletal muscle in man. *J Physiol* 366:
 554 233-249, 1985.
- 555 2. Bachler S, Mosbach F, Gatterer H, Burtscher M & Mairer K. Evaluation of an MR556 conditional "cardio stepper" for cardiopulmonary exercise testing. *Cardiothoracic and Vascular*557 *Sciences* 2: 108-112, 2017.
- 558 3. Bailey TG, Cable NT, Miller GD, Sprung VS, Low DA, and Jones H. Repeated Warm
 559 Water Immersion Induces Similar Cerebrovascular Adaptations to 8 Weeks of Moderate-
- 560 Intensity Exercise Training in Females. Int J Sports Med 37: 757-765, 2016.
- 561 4. Biswas A, Oh PI, Faulkner GE, Bajaj RR, Silver MA, Mitchell MS, and Alter DA.
 562 Sedentary time and its association with risk for disease incidence, mortality, and hospitalization
 563 in adults: a systematic review and meta-analysis. *Ann Intern Med* 162: 123-132, 2015.
- 5. Brunt VE, Howard MJ, Francisco MA, Ely BR, and Minson CT. Passive heat therapy
 improves endothelial function, arterial stiffness and blood pressure in sedentary humans. J *Physiol* 594: 5329-5342, 2016.
- 567 6. Calbet JA, Jensen-Urstad M, van Hall G, Holmberg HC, Rosdahl H, and Saltin B.
 568 Maximal muscular vascular conductances during whole body upright exercise in humans. J
 569 *Physiol* 558: 319-331, 2004.
- 570 7. Chiesa ST, Trangmar SJ, and González-Alonso J. Temperature and blood flow
 571 distribution in the human leg during passive heat stress. *J Appl Physiol (1985)* 120: 1047-1058,
 572 2016.
- 573 8. Crandall CG, and Wilson TE. Human cardiovascular responses to passive heat stress.
 574 *Compr Physiol* 5: 17-43, 2015.
- 575 9. Dawson EA, Cable NT, Green DJ, and Thijssen DHJ. Do acute effects of exercise on vascular function predict adaptation to training? *Eur J Appl Physiol* 118: 523-530, 2018.
- 577 10. Fontana P, Boutellier U, and Toigo M. Reliability of measurements with Innocor during
 578 exercise. *Int J Sports Med* 30: 747-753, 2009.
- 579 11. Frank SM, Raja SN, Bulcao CF, and Goldstein DS. Relative contribution of core and
 580 cutaneous temperatures to thermal comfort and autonomic responses in humans. *J Appl Physiol*581 (1985) 86: 1588-1593, 1999.
- 582 12. Gagnon D, Schlader ZJ, and Crandall CG. Sympathetic activity during passive heat stress
 583 in healthy aged humans. *J Physiol* 593: 2225-2235, 2015.
- 584 13. Ganio MS, Overgaard M, Seifert T, Secher NH, Johansson PI, Meyer MA, and Crandall
 585 CG. Effect of heat stress on cardiac output and systemic vascular conductance during simulated
- hemorrhage to presyncope in young men. Am J Physiol Heart Circ Physiol 302: H1756-1761,
 2012.
- 588 14. González-Alonso J, and Calbet JA. Reductions in systemic and skeletal muscle blood
 589 flow and oxygen delivery limit maximal aerobic capacity in humans. *Circulation* 107: 824-830, 2003.
- **591** 15. Green DJ, Bilsborough W, Naylor LH, Reed C, Wright J, O'Driscoll G, and Walsh JH.
- 592 Comparison of forearm blood flow responses to incremental handgrip and cycle ergometer
 593 exercise: relative contribution of nitric oxide. *J Physiol* 562: 617-628, 2005.
- **594** 16. Green DJ, Hopman MT, Padilla J, Laughlin MH, and Thijssen DH. Vascular Adaptation **595** to Exercise in Humans: Role of Hemodynamic Stimuli. *Physiol Rev* 97: 495-528, 2017.
- to Exercise in Humans: Role of Hemodynamic Stimuli. *Physiol Rev* 97: 495-528, 2017.
 Green DJ, Maiorana A, O'Driscoll G, and Taylor R. Effect of exercise training on
- 617. Green DJ, Malorana A, O Driscoll G, and Taylor K. Effect of exercise training617. endothelium-derived nitric oxide function in humans. J Physiol 561: 1-25, 2004.

- **598** 18. Hansen AB, Moralez G, Romero SA, Gasho C, Tymko MM, Ainslie PN, Hofstätter F,
- 599 Rainer SL, Lawley JS, Hearon CM. Mechanisms of sympathetic restraint in human skeletal
- 600 muscle during exercise: role of á-adrenergic and nonadrenergic mechanisms. *American Journal*
- 601 *of Physiology-Heart and Circulatory Physiology* 319, 2020.
- Herr MD, Hogeman CS, Koch DW, Krishnan A, Momen A, and Leuenberger UA. A real time device for converting Doppler ultrasound audio signals into fluid flow velocity. *Am J*
- 604 *Physiol Heart Circ Physiol* 298: H1626-1632, 2010.
- 605 20. Huonker M, Schmid A, Schmidt-Trucksass A, Grathwohl D, and Keul J. Size and blood
 606 flow of central and peripheral arteries in highly trained able-bodied and disabled athletes. *J Appl*607 *Physiol* (1985) 95: 685-691, 2003.
- 608 21. Jorfeldt L, and Wahren J. Leg blood flow during exercise in man. *Clin Sci* 41: 459-473,
 609 1971.
- 610 22. Laukkanen T, Khan H, Zaccardi F, and Laukkanen JA. Association between sauna
- bathing and fatal cardiovascular and all-cause mortality events. *JAMA Intern Med* 175: 542-548,
 2015.
- 613 23. Limberg JK, Casey DP, Trinity JD, Nicholson WT, Wray DW, Tschakovsky ME, Green
- **614** DJ, Hellsten Y, Fadel PJ, Joyner MJ, and Padilla J. Assessment of resistance vessel function in
- 615 human skeletal muscle: guidelines for experimental design, Doppler ultrasound, and
- 616 pharmacology. *Am J Physiol Heart Circ Physiol* 318: H301-H325, 2020.
- 617 24. Miller VM, and Burnett JC. Modulation of NO and endothelin by chronic increases in
 618 blood flow in canine femoral arteries. *Am J Physiol* 263: H103-108, 1992.
- 619 25. Modin AIu. [Effect of gravity on blood flow and distribution in large vessels of healthy
 620 human subjects]. *Fiziol Cheloveka* 29: 47-52, 2003.
- 621 26. Mortensen SP, Dawson EA, Yoshiga CC, Dalsgaard MK, Damsgaard R, Secher NH, and
- 622 González-Alonso J. Limitations to systemic and locomotor limb muscle oxygen delivery and
 623 uptake during maximal exercise in humans. *J Physiol* 566: 273-285, 2005.
- 624 27. Neff D, Kuhlenhoelter AM, Lin C, Wong BJ, Motaganahalli RL, and Roseguini BT.
- **625** Thermotherapy reduces blood pressure and circulating endothelin-1 concentration and enhances
- 626 leg blood flow in patients with symptomatic peripheral artery disease. *Am J Physiol Regul Integr*627 *Comp Physiol* 311: R392-400, 2016.
- **628** 28. Padilla J, Simmons GH, Vianna LC, Davis MJ, Laughlin MH, and Fadel PJ. Brachial
- artery vasodilatation during prolonged lower limb exercise: role of shear rate. *Exp Physiol* 96: 1019-1027, 2011.
- 631 29. Piepoli MF, Hoes AW, Agewall S, Albus C, Brotons C, Catapano AL, Cooney MT, Corrà
 632 U, Cosyns B, Deaton C, Graham I, Hall MS, Richard Hobbs FD, Løchen ML, Löllgen H,
- 633 Marques-Vidal P, Perk J, Prescott E, Redon J, Richter DJ, Sattar N, Smulders Y, Tiberi M, Bart
- 634 van der Worp H, van Dis I, and Monique Verschuren WM. 2016 European Guidelines on
- 635 cardiovascular disease prevention in clinical practice. *Rev Esp Cardiol* (*Engl Ed*) 69: 939, 2016.
- 636 30. Poole DC, Behnke BJ, and Musch TI. The role of vascular function on exercise capacity637 in health and disease. *J Physiol* 2020.
- 638 31. Poole DC, Gaesser GA, Hogan MC, Knight DR, and Wagner PD. Pulmonary and leg
- 639 VO2 during submaximal exercise: implications for muscular efficiency. *J Appl Physiol (1985)*640 72: 805-810, 1992.
- 641 32. Proctor DN, Shen PH, Dietz NM, Eickhoff TJ, Lawler LA, Ebersold EJ, Loeffler DL, and
- **642** Joyner MJ. Reduced leg blood flow during dynamic exercise in older
- **643** endurance-trained men. J Appl Physiol (1985) 85: 68-75, 1998.

- 644 33. Rådegran G. Ultrasound Doppler estimates of femoral artery blood flow during dynamic
 645 knee extensor exercise in humans. *Journal of Applied Physiology* 83: 1383–1388, 1997.
- 646 34. Richardson RS, Kennedy B, Wagner PD. High Muscle Blood Flows Are Not Reduced By
 647 Recruitment Of Additional Muscle Mass. *Medicine & Science in Sports & Exercise* 27, 1995.
- 648 35. Romero SA, Gagnon D, Adams AN, Cramer MN, Kouda K, and Crandall CG. Acute limb
 649 heating improves macro- and microvascular dilator function in the leg of aged humans. *Am J*650 *Physiol Heart Circ Physiol* 312: H89-H97, 2017.
- 651 36. Rowell LB, Saltin B, Kiens B, and Christensen NJ. Is peak quadriceps blood flow in
 652 humans even higher during exercise with hypoxemia? *Am J Physiol* 251: H1038-1044, 1986.
- 653 37. Simmonds GH, Wong BJ, Holowatz LA, Kenny LW. Changes in the control of skin
 654 blood flow with exercise training: where do cutaneous vascular adaptations fit in? *Experimental*655 *Physiology* 96: 822-828, 2011.
- 656 38. Thijssen DH, Dawson EA, Black MA, Hopman MT, Cable NT, and Green DJ. Brachial
 657 artery blood flow responses to different modalities of lower limb exercise. *Med Sci Sports Exerc*658 41: 1072-1079, 2009.
- 659 39. Thijssen DH, Rowley N, Padilla J, Simmons GH, Laughlin MH, Whyte G, Cable NT, and
 660 Green DJ. Relationship between upper and lower limb conduit artery vasodilator function in
- 661 humans. J Appl Physiol (1985) 111: 244-250, 2011.
- **662** 40. Thomas KN, van Rij AM, Lucas SJ, and Cotter JD. Lower-limb hot-water immersion
- acutely induces beneficial hemodynamic and cardiovascular responses in peripheral arterial
 disease and healthy, elderly controls. *Am J Physiol Regul Integr Comp Physiol* 312: R281-R291,
 2017.
- 41. Thomas KN, van Rij AM, Lucas SJ, Gray AR, and Cotter JD. Substantive hemodynamic
 and thermal strain upon completing lower-limb hot-water immersion; comparisons with treadmill
 running. *Temperature (Austin)* 3: 286-297, 2016.
- 42. Tinken TM, Thijssen DH, Hopkins N, Dawson EA, Cable NT, and Green DJ. Shear stress
 mediates endothelial adaptations to exercise training in humans. *Hypertension* 55: 312-318, 2010.
- 43. Walther G, Nottin S, Dauzat M, and Obert P. Femoral and axillary ultrasound blood flow
 during exercise: a methodological study. *Med Sci Sports Exerc* 38: 1353-1361, 2006.
- 44. Walther G, Nottin S, Karpoff L, Pérez-Martin A, Dauzat M, and Obert P. Flow-mediated
 dilation and exercise-induced hyperaemia in highly trained athletes: comparison of the upper and
- 675 lower limb vasculature. *Acta Physiol (Oxf)* 193: 139-150, 2008.
- 45. World Health Organisation (2010). Physical Activity and Adults. Available at: https://www.who.int/dietphysicalactivity/factsheet_adults/en/
- **678** 46. World Health Organisation (2018). The top 10 causes of death. Available at:
- 679 https://www.who.int/news-room/fact-sheets/detail/the-top-10-causes-of-death
- 680

681

682 Table and Figure Captions

Figure 1. Schematic of experimental design and set up of heating and exercise trial. Note valuesare estimated to demonstrate matching of intensity and do not reflect those recorded in study.

000

Figure 2. Comparison of the changes in femoral and brachial blood flow (Panel A), alongside
changes in mean, antegrade and retrograde shear rate in the femoral (Panel B) and brachial (Panel
C) arteries after 30 minutes PHWI (42 °C) versus semi recumbent stepping exercise performed at
two workloads equivalent to the increase in cardiac output (Qc_{MATCHED}) and heart rate
(HR_{MATCHED}) measured at the end of 30 minutes passive heating (n=15).

691

692 Figure 3. (A) Example of beat-by-beat blood flow and shear in the common femoral artery and 693 heart rate during exercise and recovery. On occasion, a transient loss (2-3 seconds) of the femoral 694 flow waveform at the end of exercise was caused by placing support under the participant's leg. 695 (B) Common femoral shear rate measured immediately after cessation of stepping exercise at the 696 HR_{MATCHED} intensity (0) and plotted in 30 second intervals over 5 min of recovery (n=15). All 697 values significantly different from 0 (**P<0.001). FA, common femoral; BF, blood flow; and 698 HR, heart rate.

Figure 4. Common femoral artery shear rate measured during semi recumbent rest (grey), 699 following 30 minutes PHWI (42 °C) (orange), after stepping exercise at Qc_{MATCHED} and 700 HR_{MATCHED} intensities (blue). The purple bar represents common femoral shear rate recalculated 701 702 from blood flow data measured using the thermodilution technique. Participants were endurance trained and were cycling at 70 W which produced a mean blood flow of 3.15 l·min⁻¹ (Proctor et 703 al. 1988). No femoral diameter was reported in the study therefore, using the mean diameter 704 705 reported in the present study (0.958), we re-calculated shear rate using the rearranged equation 706 (see below) to derive blood velocity (TAMV) and then applied blood velocity values to estimate 707 shear rate. Additional data include superficial femoral artery shear rate during leg extension 708 exercise in national level cyclists and swimmers (black) (Walther et al. 2008). The brown bar 709 represents common femoral artery shear rate recalculated based on blood flow values reported during knee extension exercise performed at 70 W for one participant. Peak blood flow values 710 were reported as 7.22 l·min⁻¹ (Rådegran, 1997), however common femoral artery diameter was 711 not reported therefore using a femoral diameter reported in elite road cyclist (10.053 mm), we 712

713 used the same method mentioned previously to re-estimate shear rate. Finally, the same approach

714 was used to estimate shear rate from (Calbet et al. 2004) and (Gonzailez-Alonso & Calbet, 2003),

715 who reported single leg blood flow values of $9.10 \, \mathrm{l \cdot min^{-1}}$ and $12.52 \, \mathrm{l \cdot min^{-1}}$ in healthy trained

716 males during cross country skiing and cycling, respectively. The same diameter reported in elite

717 cyclists was used (10.053 mm) for calculation of shear as participants in these studies were also

718 elite level athletes.

719

Table 1. Brachial and femoral artery blood flow and shear rate patterns pre and post 30 minutes
of passive hot water (42 °C) immersion.

722 Data are mean ± standard deviation. n =15. % Ä calculated from baseline. OSI, oscillatory shear
723 index.

724

Table 2. Brachial and femoral blood flow and shear rate patterns measured during stepping
exercise matched to the increase in cardiac output and heart rate measured after 30 minutes
passive hot water (42 °C) immersion.

728 Data are mean ± standard deviation. n =15, P<0.05*, P<0.01** compared to baseline. P<0.01†
729 Qc matched compared to HR Matched. % Ä calculated from baseline. Abbreviations: OSI,
730 oscillatory shear index.

731

Table 3. Change in cardiovascular variables at rest and during stepping exercise matched to the
increase in cardiac output and heart rate measured after 30 minutes passive hot water (42 °C)
immersion.

735 Data are mean \pm standard deviation. n=15.* $P \le 0.05$ difference to baseline. $\# P \le 0.05$ difference**736** between heating and Qc matched. $\dagger P \le 0.05$ difference between heating and HR matched.**737** Abbreviations: Qc, cardiac output;- HR, heart rate;- SV, stroke volume;- TPR, total peripheral**738** resistance;- SBP, systolic blood pressure;- DBP, diastolic blood pressure;- MAP, mean arterial**739** pressure;- T_{REC}, rectal temperature. Data for forearm skin temp, forearm blood flow and forearm**740** vascular conductance are from 14 participants. Note the discrepancy in HR and SV calculation of**741** Qc is because HR is presented from resting steady-state ECG, whereas SV is calculated from the

- 742 measured cardiac output and HR during the rebreathing procedure, which tends to increase HR
- **743** slightly.

744













	Baseline	Heating	% Д	P-Value
Brachial artery			-	-
Blood Flow $(ml \cdot min^{-1})$	44 ± 11	11 $254 \pm 67 + 47$		< 0.001
Total Shear Rate (s ⁻¹)	$56 \pm 13 \qquad \qquad 312 \pm 76$		+ 457	< 0.001
Antegrade Shear Rate (s ⁻¹)	69 ± 15	312 ± 74	+ 352	< 0.001
Retrograde Shear Rate (s ⁻¹)	-13 ± 12	-2 ± 4	-2 ± 4 - 85	
OSI	0.270 ± 0.265	0.008 ± 0.016	- 97	0.002
Conductance (ml·min ⁻¹ ·100 mmHg ⁻¹)	65 ± 16	368 ± 79	+ 466	< 0.001
Diameter (cm)	0.398 ± 0.048	0.404 ± 0.054		0.342
Femoral artery				
Blood Flow (ml·min ⁻¹)	295 ± 126	1599 ± 402	+ 442	< 0.001
Total Shear Rate (s ⁻¹)	28 ± 9	161 ± 59	+ 475	< 0.001
Antegrade Shear Rate (s ⁻¹)	34 ± 10	161 ± 59	+374	< 0.001
Retrograde Shear Rate (s ⁻¹)	-5 ± 3	0 ± 1	-100	< 0.001
OSI	0.200 ± 0.169	0.002 ± 0.007	-99	< 0.001
Conductance (ml·min ⁻¹ ·100 mmHg ⁻¹)	382 ± 165	2051 ± 439	+437	< 0.001
Diameter (cm)	0.957 ± 0.117	0.959 ± 0.118		0.852

	Baseline	Qc Matched	% Δ	HR Matched	% Δ	ANOVA B. Value
D				· · · ·		P-value
Brachial artery						
Blood Flow $(ml \cdot min^{-1})$	37 ± 19	39 ± 21	+5	$109 \pm 62^{**}$ †	+195	0.0004
Total Shear Rate (s^{-1})	59 ± 24	67 ± 30	+ 14	$168 \pm 84^{**}$ †	+185	0.0003
Antegrade Shear Rate (s ⁻¹)	69 ± 21	101 ± 26 **	+46	188 ± 74 **†	+172	< 0.0001
Retrograde Shear Rate (s^{-1})	-10 ± 8	$-33 \pm 23^{**}$	-230	-18 ± 19	+80	0.0023
OSI	0.22 ± 0.24	$0.65 \pm 0.56 **$	+195	0.15 ± 0.19 **	-32	0.0011
Conductance $(ml \cdot min^{-1} \cdot 100 mmHg^{-1})$	56 ± 33	47 ± 30	-16	$117 \pm 68*$ †	+109	0.0018
Diameter (cm)	0.387 ± 0.053	0.379 ± 0.048		0.378 ± 0.054		0.0391
<u>Femoral artery</u>						
Blood Flow $(ml \cdot min^{-1})$	374 ± 108	$1844 \pm 468 **$	+393	$2588 \pm 527 ** \ddagger$	+592	< 0.0001
Total Shear Rate (s ⁻¹)	41 ± 15	$193\pm60^{\boldsymbol{\ast\ast}}$	+370	271 ± 59 **†	+561	< 0.0001
Antegrade Shear Rate (s^{-1})	49 ± 14	$193\pm59^{\boldsymbol{**}}$	+294	270 ± 59 **†	+451	< 0.0001
Retrograde Shear Rate (s ⁻¹)	-8 ± 4	$-0.6 \pm 1**$	-93	-0.05 ± 0.09 **	-99	< 0.0001
OSI	0.23 ± 0.18	0.01 ± 0.01 **	-96	0.00 ± 0.00 **	-100	0.0002
Conductance $(ml \cdot min^{-1} \cdot 100 mmHg^{-1})$	445 ± 128	2121 ± 497**	+377	$2725 \pm 567 ** \ddagger$	+512	<0.0001
Diameter (cm)	0.931 ± 0.085	0.980 ± 0.096		0.938 ± 0.107		0.5581

	Hea	ting	Exerci	ANOVA P-Value		
	Baseline Heating	Δ Heating	Baseline Exercise	ΔQc Matched	Δ HR Matched	
$Qc (l \cdot min^{-1})$	7.66 ± 1.4	$3.72 \pm 1.9*$	7.68 ± 1.9	$3.78 \pm 2.00*$	$7.02 \pm 6.00 \ddagger$	0.03
Heart rate (beats min ⁻¹)	65 ± 10	$38\pm15*$	63 ±9	18 ± 10 *#	40 ±32*	< 0.0001
SV (ml)	111 ± 20	1 ±19	122 ± 26	20 ±24*#	21 ±57*	0.19
Femoral TPR (mmHg·ml·min ⁻¹)	10.6 ± 2.6	-3.4*	11 ± 1.9	$8 \pm 1.1*$	$5 \pm 1.1*$ †	0.02
Femoral conductance $(ml \cdot min^{-1} \cdot 100 mmHg^{-1})$	382 ± 165	$1669 \pm 415 *$	445 ± 128	$1675\pm495\texttt{*}$	2279 ± 569* †	0.0009
SBP (mmHg)	111 ± 9	$28 \pm 21*$	116 ± 9	$11 \pm 12.33 * \#$	$43 \pm 19*$ †	< 0.0001
DBP (mmHg)	61 ± 6	$-13 \pm 10^{*}$	68 ± 8	$-5 \pm 5*#$	-4 ± 14	0.03
MAP (mmHg)	77 ± 6	0 ± 11	84 ± 7	1 ± 5	$11 \pm 11*$ †	0.0027
Pulse pressure (mmHg)	49 ± 11	41*	48 ± 9	$63 \pm 13*\#$	$95 \pm 21*$	< 0.0001
Rectal temperature (°C)	36.97 ± 0.20	$1.31 \pm 0.37*$	36.88 ± 0.22	$0.03\pm0.11\#$	0.39 ± 0.19 *†	< 0.0001
Forearm skin Temp (°C)	29.45 ± 1.03	$3.71 \pm 1.39*$	28.44 ± 0.87	$0.00\pm0.40^{\#}$	0.66 ± 1.02 *†	< 0.0001
Forearm skin blood flow (PU)	51 ± 41	$132 \pm 79*$	30 ± 10	14 ± 20 *#	$74 \pm 47*$	0.003
Forearm cutaneous vascular conductance (a.u ¹ ·100 mmHg ⁻¹)	67 ± 52	$114 \pm 68*$	35 ± 12	$48 \pm 48*$	$75 \pm 54*$	0.06
VO_{2} (ml·kg·min ⁻¹)	5.36 ± 1.43	$0.92 \pm 1.40 \texttt{*}$	4.51 ± 1.41	7.79 ± 2.76 *#	$16.52 \pm 7.57 * \ddagger$	< 0.0001