

Aortic stiffness contributes to greater pressor responses during static hand grip exercise in healthy young and middle-aged normotensive men

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1	Aortic stiffness contributes to greater pressor responses during static hand grip
2	exercise in healthy young and middle-aged normotensive men
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23 Abstract

24 Central arterial stiffness can influence exercise blood pressure (BP) by increasing the rise in 25 arterial pressure per unit increase in aortic inflow. Whether central arterial stiffness influences 26 the pressor response to isometric handgrip exercise (HG) and post-exercise muscle ischemia 27 (PEMI), two common laboratory tests to study sympathetic control of BP, is unknown. We 28 studied 46 healthy non-hypertensive males (23 young and 23 middle-aged) during HG (which 29 increases in cardiac output [Qc]) and isolated metaboreflex activation PEMI (no change or 30 decreases in Qc). Aortic stiffness (aortic pulse wave velocity [aPWV]; applanation tonometry 31 via SphygmoCor) was measured during supine rest and was correlated to the pressor 32 responses to HG and PEMI. BP (photoplethysmography) and muscle sympathetic nerve 33 activity (MSNA) were continuously recorded at rest, during HG to fatigue (35% maximal 34 voluntary contraction) and 2-minutes of PEMI. aPWV was higher in middle-aged compared to 35 young males (7.1±0.9 vs 5.4±0.7 m/s, P<0.001). Middle-aged males also exhibited greater increases in systolic pressure ($\Delta 30\pm 11$ vs 10±8 mmHg) and MSNA ($\Delta 2313\pm 2006$ vs 36 37 1387±1482 %/min) compared to young males during HG (both, P<0.03); with no difference in 38 the Qc response (P=0.090). Responses to PEMI were not different between groups. Sympathetic transduction during these stressors (MSNA-diastolic pressure slope) was not 39 40 different between groups (P>0.341). Middle-aged males displayed a greater increase in SBP 41 per unit change of $\dot{Q}c$ during HG (Δ SBP/ $\Delta\dot{Q}c$; 21±18 vs 6±10 mmHg/L/min, P=0.004), with a 42 strong and moderate relationship between the change in systolic (r=0.53, P<0.001) and diastolic pressure (r=0.34, P=0.023) and resting aPWV, respectively; with no correlation 43 during PEMI. Central arterial stiffness can modulate pressor responses during stimuli 44 associated with increases in cardiac output and sympathoexcitation in healthy males. 45

46 Introduction

47 Static handgrip exercise (HG) is often employed in the laboratory or clinic to study autonomic 48 adjustments that regulate arterial blood pressure (BP). Notably, the basis of this approach is 49 that the arterial BP response to HG is underpinned primarily by reflex increases in sympathetic vasomotor outflow (i.e. muscle sympathetic nerve activity [MSNA] (Fisher et al., 50 51 2015). Hence, the magnitude of the exercise pressor reflex response provides an index of 52 activation of MSNA. Furthermore, an exaggerated increase in arterial blood pressure with 53 exercise is predictive of the future diagnosis of hypertension (Kayrak et al., 2010; Matthews 54 et al., 1993; Schultz et al., 2015) and cardiovascular events (Lewis et al., 2008), as reviewed 55 previously (Schultz et al., 2017). The cold pressor test is another classical laboratory/clinical test used to elicit increases in MSNA and arterial BP (Victor et al., 1987). Recently, Borner 56 57 and colleagues showed that resting aortic stiffness (aortic pulse wave velocity [aPWV]) 58 positively correlates with the change in BP induced by a cold pressor test in a cohort of young 59 and older individuals (Borner et al., 2017). This finding raises the intriguing possibility that, in addition to reflex sympathoexcitation, the arterial BP response to CPT is influenced by central 60 61 arterial stiffness.

62 Stiffer central arteries contribute to a greater pressor response during increases in 63 cardiac output (Qc) due to a reduced ability of the central arteries to distend and 64 accommodate aortic inflow. This is demonstrated as a greater rise in arterial pressure per unit increase in flow during whole body dynamic exercise (Miyai et al., 2021; Sarma et al., 65 66 2020; Thanassoulis et al., 2012). Therefore, age-related central artery stiffening (McEniery 67 et al., 2005) could contribute to greater pressor responses during HG in older, compared to 68 younger, individuals as suggested previously (Lalande et al., 2014). However, it is unknown 69 whether arterial stiffness correlates with the pressor responses to HG; if the correlation exists, 70 previously reported differences in pressor responses to small muscle mass exercise (i.e. HG) 71 may be confounded by differences in arterial stiffness rather than reflecting a difference in 72 the effectiveness of sympathetic nerve activity in eliciting vasoconstriction. Thus, the 73 knowledge of whether arterial stiffness contributes to pressor responses to HG is highly-74 relevant for the field and interpretation of these tests.

Herein, we assessed whether arterial stiffness correlates with pressor responses in healthy non-hypertensive young and middle-aged men during HG and post-exercise muscle ischaemia (PEMI). HG is associated with an increase in Qc and sympathoexcitation while PEMI is associated with sympathoexcitation and no change, or even a decrease, in Qc (Kiviniemi et al., 2012). The use of these two stimuli in younger and middle-aged individuals, known to have differences in aortic stiffness (Talbot et al., 2020; Wakeham et al., 2022), 81 facilitates investigation as to whether arterial stiffness influences the pressor responses to

82 changes in Qc.

83 Methodology

84 All data presented were collected during 2015-2017 as part of a cross-sectional study in 23 85 young $(23 \pm 3 [18 - 30]$ years) and 23 middle-aged $(55 \pm 4 [50 - 63]$ years) endurance-trained 86 and recreationally-active healthy men designed to address several *a-priori* research aims 87 (Lord et al., 2020; Talbot et al., 2020; Wakeham et al., 2022; Wakeham et al., 2019). These prior aims were to assess the independent effects of age and habitual exercise on integrative 88 89 cardiovascular control by studying men only due to the well-known sex differences in 90 autonomic support of blood pressure, MSNA, BP, sympathetic transduction at rest or during 91 HG and PEMI in both younger and older age (Best et al., 2014; Christou et al., 2005; Hart et 92 al., 2011; Jarvis et al., 2011; Vianna et al., 2012), as reviewed previously (Joyner et al., 2015).

93 We recruited forty-six males, all were non-smokers, normotensive and reported no 94 chronic diseases. Recruitment criteria were either young (18-30 years) or middle-aged (50-95 65 years) men who were recreationally-active (\leq 3hours of structured physical activity for \geq 2 96 or \geq 10years for the young and middle-aged men, respectively) or endurance-trained (Young: 97 \geq 50 miles of moderate to intensity training for \geq 2 years; Middle-aged: \geq 25 miles of moderate 98 to intensity training for ≥ 10 years). Participants were requested to abstain from caffeine, 99 alcohol, nutritional supplements, and heavy exercise for 24 hours prior to testing; furthermore, 100 participants arrived having fasted for the previous 6 hours. All participants provided written 101 informed consent. The study conformed to the Declaration of Helsinki, except for registration 102 as a clinical trial and was approved by the Cardiff Metropolitan University School of Sport and 103 Health Sciences Research Ethics Committee (16/7/02R).

104 Participants attended the laboratory on two occasions, first for a screening visit and 105 second for the experimental visit. Study visits were separated by a minimum of 24 hours. 106 During the screening visit, a PWV and cardiorespiratory fitness ($\dot{V}O_2$ Peak) were assessed, as 107 described previously (Wakeham et al., 2022; Wakeham et al., 2019). aPWV was assessed 108 via applanation tonometry (SphygmoCor) measuring the transit time between the foot of the 109 carotid and femoral arterial waveforms divided by the path length (measuring tape). \dot{VO}_2 Peak 110 was assessed with an incremental (ramp, 20 Watts/min) test to exhaustion on a cycle 111 ergometer. During the experimental visit, heart rate (electrocardiography), beat-by-beat blood 112 pressure (finger photoplethysmography; FinometerPro, FMS, Groningen, Netherlands), 113 multiunit MSNA (microneurography; Nerve Traffic Analyzer, Model 663 C, University of Iowa, 114 lowa City, IA) and left ventricular volumes (2-D echocardiography; Vivid E9, GE Medical, Norway) were recorded at rest (Wakeham et al., 2019) and during HG and PEMI. Briefly, 115

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116 following a 1-minute baseline, participants performed HG at 35% of their maximal voluntary 117 contraction with their left hand (MLT004/D, ADInstruments, Oxford, UK; 5 participants were 118 left hand dominant). When a participant was unable to maintain the required force for >3 119 seconds, a cuff positioned around the left forearm (to enable ultrasonography at the level of 120 the brachial artery; data not shown) was rapidly inflated (E20 Rapid Cuff Inflation System, D.E. 121 Hokanson, Bellevue, USA) to suprasystolic pressure (220mmHg), to elicit a period of PEMI 122 for two minutes. To account for inter-individual variability in time to task failure. HG duration 123 was divided into five equal quintiles (20%, 40%, 60%, 80%, 100% HG duration); whereas 124 PEMI was assessed in quartiles (30 second bins).

125 Data Acquisition and Analyses

Echocardiographic images were acquired with a 4-MHz array probe (Vivid E9, GE Medical, Norway) over five cardiac cycles by an experienced cardiac sonographer (RNL) and stored for analysis off-line. Left ventricular stroke volume was derived in the single plane from apical 4-chamber views in 2D echocardiograms as the as the absolute difference between end diastolic and systolic volumes using commercially available software (EchoPAC, BT12 GE Medical, Norway).

132 The beat-by-beat arterial pressure waveform was calibrated against the average of 133 three systolic (SBP) and diastolic (DBP) blood pressures, measured at rest, using a manual 134 sphygmomanometer. We calculated pulse pressure (PP [SBP-DBP]). Heart rate was 135 determined from the R-R interval recorded in the electrocardiogram (Lead II). The assessment 136 of stroke volume and arterial pressure permitted the calculation of Qc (heart rate x stroke 137 volume), total vascular conductance (TVC, Qc/mean arterial pressure [MAP]) and total 138 peripheral resistance (TPR, MAP/ Qc). Stroke volume could not be obtained in 4 middle-aged 139 males, therefore stroke volume and associated hemodynamics are reported for 42 individuals 140 during all stimuli (23 young and 19 middle-aged males).

MSNA (raw and integrated) signals, arterial BP, electrocardiogram, and grip force data were sampled at 1 kHz using analog-to-digital data acquisition hardware (Powerlab 8/35, ADInstruments). Multiunit bursts of integrated MSNA were inspected independently by two researchers (DJW/JPM) and verified. To account for variation in the microelectrode position, and the effect this has on MSNA burst amplitude, the height of the largest burst appearing under resting conditions was assigned a value of 100 units. All other bursts were expressed relative to this value.

MSNA was quantified as burst frequency (bursts/min), burst incidence (bursts/100hb;
hb, heartbeats), mean burst amplitude (%) and total activity (burst amplitude x burst frequency,
%/min). We were unable to record MSNA from one middle-aged participant. MSNA total

activity was used as the primary index of sympathetic responses, as it accounts for changes
in both burst occurrence and burst size. Thirty seconds of representative MSNA and blood
pressure data are presented from one young (Figure 1, A-C) and one middle-aged male
(Figure 1, D-F) during baseline, at the end of SHG and at the end of PEMI.

155 Sympathetic transduction to pressure was assessed via the calculation of slopes from 156 linear regression analyses (Halliwill et al., 1996) between MSNA burst frequency and DBP 157 and MSNA total activity and DBP, respectively (see representative data in Figure 2). DBP 158 was used as it is reproducible, and a target variable of sympathetic outflow; furthermore, DBP 159 indicates systemic vascular responses (Briant et al., 2016). To generate sympathetic vascular 160 transduction slopes, the relationship between MSNA burst frequency and total activity was 161 plotted against DBP, for each quintile (HG) or quartile (PEMI). The baseline value for each 162 variable was included in the regression analyses. Due to similar between-group differences 163 when sympathetic vascular transduction was determined using MSNA burst frequency and 164 total activity, only data using total activity are reported.

165 Statistical Analysis

166 All analyses were completed using IBM SPSS (version 26, IBM statistics, Armonk, NY). Data 167 were tested for normality (Shapiro-Wilks), sphericity (Mauchly's test) and the presence of significant outliers (≥ 3 standard deviations). Participant characteristics, HG duration and 168 169 sympathetic transduction were compared via independent samples t-tests. Neural and 170 haemodynamic data were analysed using a linear mixed effects model including subject as a 171 random factor, to determine main effects of group (young versus middle-aged) and condition 172 (pre-test baseline versus final quintile of HG or final quartile of PEMI). In the event of a 173 significant interaction, SIDAK post hoc-multiple comparisons were performed. The relationship 174 between variables was assessed via Pearson product-moment correlation coefficients. 175 Statistical significance was defined at a level of P < 0.05. Values are reported in text as mean 176 ± standard deviation (SD).

177 Results

- 178 Middle-aged males were older (55 \pm 4 vs 23 \pm 3 years, *P* < 0.001), of greater stature (179.1 \pm
- 179 5.4 vs 175.1 \pm 6.7 cm, *P* < 0.05), had a higher body fat percentage (22.0 \pm 7.8 vs 14.8 \pm 7.0
- 180 %, P < 0.05; via bioelectrical impedance) and aPWV (7.1 ± 0.9 vs 5.4 ± 0.7 m/s, P < 0.001)
- 181 compared to young males. Body mass (72.3 ± 11.6 vs 72.8 ± 12.9 kg, P = 0.896), $\dot{V}O_2$ Peak
- 182 (43.8 ± 10.9 vs 50.1 ± 14.6 mL/kg/min, *P* = 0.108) and HG duration (250 ± 108 vs 190 ± 136
- 183 seconds, P = 0.119) were not different between groups.
- 184 Baseline cardiovascular haemodynamics and neural activity

- 185 Heart rate, stroke volume, cardiac output and TVC were lower in middle-aged males; whereas,
- 186 TPR, blood pressure (all) and MSNA burst frequency, burst incidence and total activity were
- higher (main effects of group, P < 0.05; Figure 2 and Table 1).

188 Handgrip exercise

In response to HG, in both groups, heart rate, Qc, total vascular conductance (TVC), and all indices of blood pressure and MSNA increased, whereas stroke volume decreased (main effects of condition, P < 0.05) and TPR did not change. However, compared to their younger counterparts, there were greater increases in MSNA total activity, systolic blood pressure (SBP) and MAP in middle-aged males (group*condition interactions, P < 0.05; Table 1 and Figure 2). Sympathetic transduction to pressure was not different (P = 0.341) between young (0.004 ± 0.008 mmHg·%·min⁻¹) and middle-aged groups (0.006 ± 0.004 mmHg·%·min⁻¹).

196 Post-exercise muscle ischaemia

197 Heart rate, stroke volume, cardiac output and TVC were lower and TPR, systolic blood 198 pressure (SBP), and MSNA (except burst amplitude) were higher in middle-aged males during 199 PEMI (main effects of group, P < 0.05). Stroke volume, Qc and total vascular conductance 200 (TVC) decreased during PEMI; whereas, TPR, blood pressure (all) and MSNA (all) increased 201 (main effects of condition, P < 0.05), with no significant effect of age on the neural or 202 haemodynamic responses to PEMI (Table 1 and Figure 3). Sympathetic transduction to 203 pressure was not different (P = 0.807) between young (0.003 ± 0.004 mmHg·%·min⁻¹) and 204 middle-aged groups (0.002 \pm 0.007 mmHg·%·min⁻¹).

205 Correlation analysis

With all data pooled, aPWV exhibited a significant positive linear correlation with the systolic, diastolic and mean pressor responses during HG, but not PEMI (Figure 4). Furthermore, there were significant correlations between aPWV and the change in pulse pressure for both HG (r = 0.54, P < 0.001) and PEMI (r = 0.55, P < 0.001); there was no significant correlation between aPWV and baseline pulse pressure (r = 0.03, P = 0.849). In line with this, middle-aged men displayed a greater increase in SBP per unit change of Qc during HG than young men (Δ SBP/ Δ Qc; 21 ± 18 vs 6 ± 10 mmHg/L/min, P = 0.004), likely due to the higher aPWV.

When age was included as a covariate for partial correlational analyses, there were no longer any significant relationships between aPWV and the pressor responses during HG (SBP: r = -0.03, P = 0.865; MAP: r = -0.04, P = 0.781; DBP: r = -0.03, P = 0.845; PP: r = -0.01, P = 0.942). However, there were no significant relationships between aPWV and pressor responses to HG in young (SBP: r = 0.19, P = 0.396; MAP: r = -0.001, P = 0.995; DBP: r = -0.11, P = 0.628; PP: r = 0.36, P = 0.092) or middle-aged (SBP: r = -0.12, P = 0.614; MAP: r = 219 -0.04, P = 0.881; DBP: r = 0.07, P = 0.765; PP: r = -0.25, P = 0.280) men when assessed 220 separately. Furthermore, all correlations remained (either significant or non-significant) when 221 adjusted for either $\dot{V}O_2$ Peak or baseline MAP (Table 2).

222 Discussion

223 Herein, middle-aged men displayed a greater increase in SBP, MAP and MSNA total activity, 224 during HG; whereas, there was no effect of age on the arterial pressure and sympathetic 225 neural responses to PEMI. Since Qc increases during HG and decreases during PEMI, we 226 speculate that arterial stiffening, such as that induced by ageing, exaggerates the pressor 227 response during HG. Moreover, when data from young and middle-aged men were combined, 228 a positive relationship exists between baseline aPWV and the absolute change in SBP, MAP 229 and DBP during HG. Notably, there is no relationship between baseline aPWV and the pressor 230 responses to PEMI. Considering these findings together, we suggest that the arterial BP 231 response to HG is influenced by central arterial stiffness. Hence, there is potential for vascular 232 stiffness to be a confounding factor in studies utilizing HG exercise to study neural control and 233 autonomic regulation of blood pressure.

234 The magnitude of the systolic and mean pressor responses during HG, but not PEMI, 235 were greater for middle-aged males. Several possible mechanisms may contribute to the 236 exaggerated pressor response to HG in middle-aged males. First of which may be higher 237 central arterial stiffening, which is commonly observed in western society (McEniery et al., 238 2005). Central arterial stiffening would exaggerate the pressor response during increases in 239 Qc, due to a reduced ability of the central arteries to distend and accommodate aortic inflow, 240 thereby increasing arterial pressure per unit increase in flow. Indeed, the middle-aged males, 241 who exhibited a greater aPWV (i.e., index of arterial stiffness), presented with a greater 242 change in systolic for a given change in Qc, compared to the younger males. Furthermore, in 243 response to HG, during which cardiac output increased, we identified a significant positive 244 relationship between aPWV and the absolute change in SBP, DBP and MAP (Figure 3A, 3C 245 and 3E). This suggests that those with greater arterial stiffness exhibit more exaggerated 246 pressor responses to increases in Qc, further highlighting the important role of arterial 247 stiffness. However, during PEMI, where cardiac output fell compared to baseline, there was 248 no significant relationship between baseline aPWV and the pressor responses (Figure 3B, 3D 249 and 3F). Importantly, to highlight the effect of age, and associated arterial stiffening, when 250 including age as a covariate in partial correlational analyses, there were no longer any 251 significant relationships between aPWV and the pressor responses during HG (SBP: r = -252 0.027, P = 0.865; MAP: r = -0.044, P = 0.781; DBP: r = -0.031, P = 0.845), suggesting that 253 this association is a function of differences in age. As a rtic stiffening appears to correlate with the pressor response during sympathoexcitation associated with increases in cardiac output
(i.e. exercise), studies comparing groups with known differences in aPWV (i.e. age) should
consider including measurement of arterial stiffness to include in covariate analysis.

257 An alternative mechanism mediating greater pressor responses during HG in middle-258 aged men could be the greater increase in MSNA total activity eliciting greater decreases in 259 vascular conductance (or increases in resistance) and ultimately DBP. Although there was no 260 significant effect of age on the response of total vascular conductance (or resistance) during 261 HG, there was a trend for a greater DBP response with age (group*condition interaction, P = 262 0.050; Table 1). This is of relevance as DBP is a target variable of vascular sympathetic activity 263 (Briant et al., 2016).Indeed, sympathetic transduction to pressure was similar between young 264 and middle-aged men, suggesting the greater increase in total MSNA activity would result in 265 a greater increase in BP, compared to young individuals. However, there was no correlation 266 between the change in MSNA and the change in arterial pressure (SBP: r = 0.278, P = 0.06; 267 MAP: r = 0.024, P = 0.114; DBP: r = 0.017, P = 0.262). Thus, it appears unlikely that the 268 greater increase in MSNA total activity contributed to the larger increase in arterial pressure 269 with age during HG.

270 Our findings contrast previous studies that report no effect of age on the MSNA or 271 pressor responses to HG (Greaney et al., 2013; Houssiere et al., 2006; Krzeminski et al., 272 2012; Markel et al., 2003; Momen et al., 2004; Ng et al., 1994; Tan et al., 2013). Nevertheless, 273 only the change in MSNA total activity was greater with age and there were no age-related 274 effects on the response of MSNA burst frequency, amplitude or incidence. Although we cannot 275 determine the reasons for the disparity between our findings and those of other studies, it 276 likely reflects between study sample differences in either hemodynamic (Watanabe et al., 277 2014) or genetic (Notay et al., 2018) factors. Furthermore, the influence of comparing absolute 278 or relative responses in previous data also may partly contribute to differences in study 279 findings. However, the greater response of MSNA and blood pressure here occurred despite 280 higher MSNA in middle-aged men with no difference in blood pressure between groups at 281 baseline. Thus, it is unlikely that baseline differences influenced the age-related difference in 282 the responses observed here. Together, it appears that the interaction between increases in 283 cardiac output, higher aortic stiffness and greater elevations in sympathetic vasomotor outflow 284 contribute to greater increases in arterial pressure during HG in middle-aged compared to 285 young males.

286 Notably, we observed no significant effect of age on changes in MSNA burst frequency 287 or amplitude during HG, suggesting it is the interaction of burst rate and size which culminates 288 in a greater increase in MSNA total activity in middle-aged males. The contributing 289 mechanism(s) to the larger increase in MSNA total activity during HG with age are unclear but 290 appear to be independent of afferent feedback from skeletal muscle metaboreceptors, as the 291 augmented MSNA response was not maintained during PEMI. A greater feedforward (central 292 command) or feedback (muscle mechanoreflex) signal could contribute to the greater level of 293 sympathoexcitation with age during HG. In addition, venous compliance decreases (Monahan 294 et al., 2001) and pulmonary vascular stiffness increases (Dawes et al., 2016) with age; 295 accordingly, during increases in Qc, there could be a greater activation of the 296 sympathoexcitatory venous distention and pulmonary baroreceptor reflexes (Moore et al., 297 2022). Despite these effects being difficult to isolate in humans, future studies should attempt 298 to investigate the mechanisms contributing to this greater level of sympathoexcitation during 299 HG with age.

300 Methodological Considerations

301 There are several strengths of our study, including the comprehensive assessment of 302 cardiovascular responses to exercise using echocardiography and microneurography, as well 303 as characterizing arterial stiffness in a large sample size. However, our inclusion of males only 304 represents a major limitation of our study, and this limits the generalisability of our findings. 305 Future studies employing a balanced sex ratio are required to assess whether these same 306 correlations exist in both sexes. Also, resting aortic stiffness was measured on a separate day 307 to the sympathetic and hemodynamic responses to HG and PEMI to address a separate a-308 priori study aim (Wakeham et al., 2022). However, the pressor responses to HG and PEMI 309 and resting aPWV have been reported to have good reproducibility (Dillon et al., 2020; Yasmin 310 et al., 1999). Accordingly, the difference in days of assessment of aortic stiffness and HG and 311 PEMI responses is unlikely to affect the conclusions of this study.

312 Conclusion

313 This study provides new information regarding the influence of aortic stiffness on pressor 314 responses during HG in men. The greater pressor response to HG in middle-aged men likely 315 occurs due to central artery stiffening, which increases arterial pressure per unit increase in 316 flow (i.e. Qc). The systolic pressor difference between groups suggests central artery stiffening 317 plays a larger role than does the greater increase in MSNA in middle-aged men, as there was 318 no difference in the DBP or TPR response. These findings suggest that pressor responses to 319 stimuli associated with increases in Qc and sympathoexcitation (e.g. exercise) are likely to be 320 exaggerated in healthy middle-aged men, who exhibit higher arterial stiffness.

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324 **Declarations**

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- 328 Sport Health and Exercise Sciences, Bangor University.
- 329 Conflicts of interest/Competing interests
- 330 Not Applicable.

331 Availability of data and material

332 Data are available from the corresponding author upon reasonable request.

333 Code availability

Not Applicable.

335 Author Contributions

All testing was completed at the Cardiff School of Sport and Health Sciences, Cardiff 336 337 Metropolitan University, Cardiff, Wales, UK. D.J.W., C.J.P., R.S., and J.P.M contributed to 338 conception and design of the work and acquisition, analysis and interpretation of the data and writing of the manuscript. R.N.L., J.S.T., F.M.L., B.A.C., T.G.D. and L.L.S., contributed to 339 340 acquisition, analysis and interpretation of the data and critically revised the manuscript. All 341 authors approved the final version of the manuscript and agree to be accountable for all 342 aspects of the work. All persons included as an author qualify for authorship, and all those 343 who qualify for authorship are listed.

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480 Figures and Tables

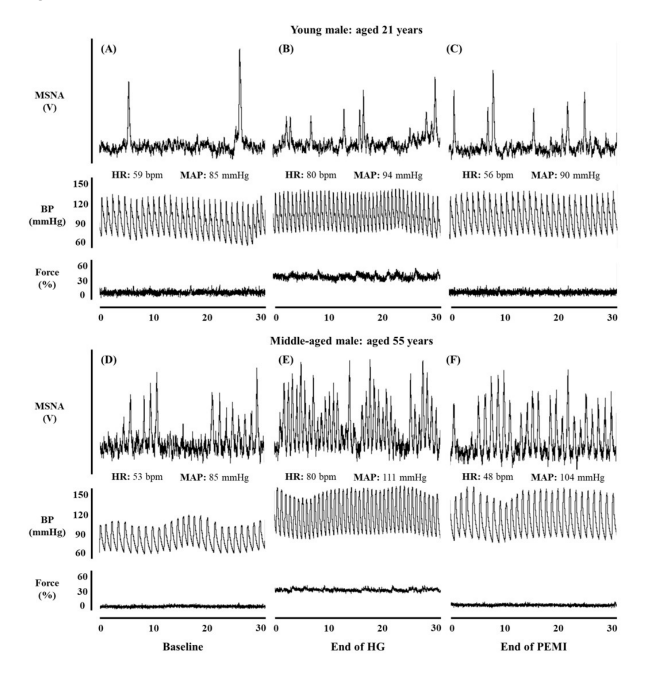
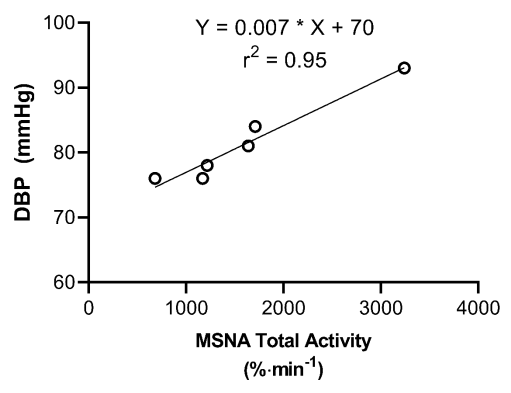
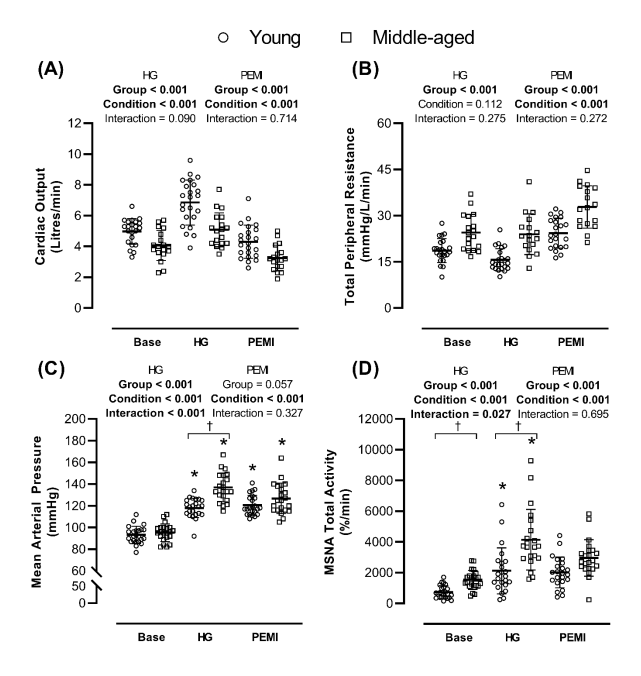


Figure 1 - Representative raw data from one young and one middle-aged man. Thirty seconds of raw muscle sympathetic nerve activity (MSNA) and blood pressure (BP) data are shown from one young (panels A, B and C) and one middle-aged male participant (panels D, E and F) at baseline, the end of static hand grip (HG) and the end of post-exercise muscle ischemia (PEMI).



486 Figure 2 - A representative sympathetic vascular transduction slope from one young participant.

487 Sympathetic vascular transduction was assessed as the slope of the linear relationship between MSNA
 488 total activity and diastolic blood pressure during hand grip exercise.



490 Figure 3 - The sympathetic and haemodynamic responses to HG and PEMI in young and middle-491 aged men. The changes in cardiac output (Panel A) and total peripheral resistance (Panel B) were not 492 different between groups. The increases in mean arterial pressure (Panel C) and MSNA Total Activity 493 (Panel D) during HG were greater in middle-aged men when compared to younger counterparts. There 494 were no significant group differences in the sympathetic or haemodynamic responses to PEMI. Notes: 495 Significant main effects or interactions are shown in bold above each respective panel, as determined 496 via separate linear mixed models for HG and PEMI. Symbols display results from SIDAK post hoc 497 analyses; * and \dagger represent within-group and between-group differences (P < 0.05), respectively. 498 Abbreviations: Base, baseline; HG, static hand grip exercise; MSNA, muscle sympathetic nerve activity; 499 PEMI, post-exercise muscle ischaemia.

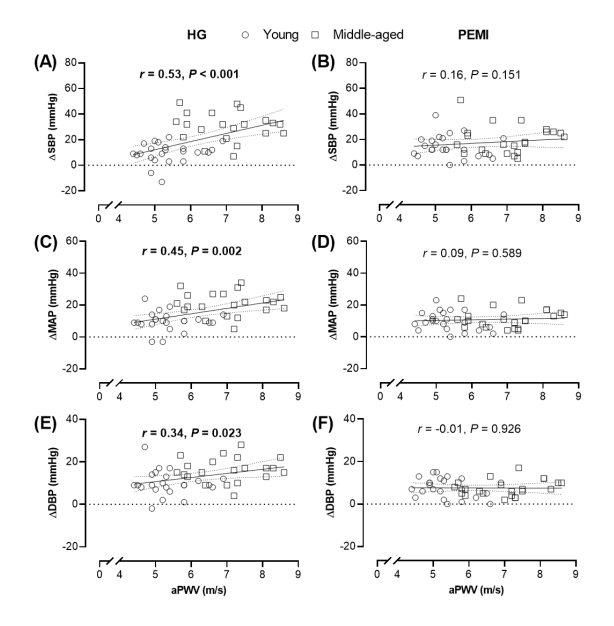


Figure 4 - Correlations between baseline aortic stiffness and the delta pressor responses to HG and PEMI. aPWV significantly correlated with the change in systolic, mean and diastolic pressure from rest to the final quintile of HG (Panels A, C and E, respectively). However, there were no significant correlations between aPWV and pressor responses to the final 30-seconds of PEMI (Panels B, D and F). Abbreviations: aPWV, aortic pulse wave velocity; DBP, diastolic blood pressure; HG, static handgrip exercise; MAP, mean arterial pressure; PEMI, post-exercise muscle ischaemia; SBP, systolic blood pressure.

				HG			PEMI				
Variable	Group	n	Baseline	End HG	Group <i>P</i>	Condition P	Interaction P	End PEMI	Group <i>P</i>	Condition P	Interaction P
Haemodynamic											
Heart Rate	Y	23	57 ± 15	86 ± 18	0.007	< 0.001	0.431	55 ± 13	0.044	0.528	0.830
(bpm)	MA	22	52 ± 10	76 ± 17	0.007			47 ± 7			
Stroke Volume	Y	23	92 ± 16	80 ± 12	< 0.001	< 0.001	0.974	79 ± 14	< 0.001	< 0.001	0.945
(ml)	MA	18	79 ± 10	67 ± 10	< 0.001			66 ± 9			
TVC	Y	23	0.056 ± 0.014	0.067 ± 0.014	< 0.001	0.028	0.108	0.042 ± 0.010	< 0.001	< 0.001	0.689
(l/min [/] mmHg)	MA	18	0.043 ± 0.010	0.045 ± 0.013				0.031 ± 0.008			
SBP	Y	23	124 ± 9	133 ± 10 *	< 0.001	< 0.001	< 0.001	138 ± 11	0.038	< 0.001	0.325
(mmHg)	MA	22	127 ± 11	158 ± 17 *†				147 ± 18			
DBP	Y	23	78 ± 10	87 ± 9	0.006	< 0.001	0.050	85 ± 8	0.315	< 0.001	0.894
(mmHg)	MA	22	80 ± 7	96 ± 8				87 ± 7			
Sympathetic activ	vity										
MSNA BF	Y	23	13 ± 8	26 ± 15	< 0.001	< 0.001	0.147	26 ± 10	< 0.001	< 0.001	0.449
(bursts/min)	MA	22	27 ± 11	47 ± 14				37 ± 11			
MSNA BI	Y	23	26 ± 18	35 ± 22		0.007	0.623	50 ± 23	< 0.001	< 0.001	0.504
(bursts/100hb)	MA	22	54 ± 24	68 ± 17	< 0.001			72 ± 21			
MSNA BA	Y	23	58 ± 11	81 ± 27	0.007	< 0.001	0.302	77 ± 21	0.359	< 0.001	0.525
(%)	MA	22	60 ± 11	94 ± 46	0.207			84 ± 38			

Data are presented as mean ± standard deviation. Significant main effects or interactions are highlighted in bold, determined via separate linear mixed models for HG and PEMI. Symbols display results from SIDAK post hoc analyses; * and † represent within-group and between-groups differences, respectively. *Abbreviations: BA, burst amplitude; BF, burst frequency; BI, burst incidence; DBP, diastolic blood pressure; HG, hand grip exercise; MA, middle-aged; MAP, mean arterial pressure; MSNA, muscle sympathetic nerve activity; n, sample size; PEMI, post-exercise muscle ischaemia; SBP, systolic blood pressure; TVC, total vascular conductance; Y, young.*

	∆SBP (mmHg)	∆DBP (mmHg)	∆PP (mmHg)	∆MAP (mmHg)				
HG								
aPWV, m/s	r = 0.53	r = 0.34	r = 0.54	r = 0.45				
(unadjusted)	P < 0.001	P = 0.023	P < 0.001	P = 0.002				
aPWV, m/s	r = 0.53	r = 0.34	r = 0.53	r = 0.45				
(adj VO2peak)	P < 0.001	P = 0.027	P < 0.001	P = 0.002				
aPWV, m/s	r = 0.52	r = 0.38	r = 0.53	r = 0.47				
(adj BL MAP)	P < 0.001	P = 0.012	P < 0.001	P = 0.002				
PEMI								
aPWV, m/s	r = 0.16	r = -0.01	r = 0.55	r = 0.09				
(unadjusted)	P = 0.151	P = 0.926	P < 0.001	P = 0.589				
aPWV, m/s	r = 0.20	r = 0.05	r = 0.57	r = 0.139				
(adj VO2peak)	P = 0.191	P = 0.728	P < 0.001	P = 0.374				
aPWV, m/s	r = 0.14	r = 0.05	r = 0.52	r = 0.101				
(adj BL MAP)	P = 0.365	P = 0.770	P < 0.001	P = 0.521				

Table 2 - Correlation Matrix

Unadjusted data are presented as Pearson's product-moment correlation coefficients, with associated significance values. Adjusted correlations for VO2peak (ml/kg/min) and baseline MAP were performed using Partial correlation analysis within SPSS v20 (IBM Corp). Significant correlations are presented in bold. *Abbreviations: aPWV, aortic pulse wave velocity; BL, baseline; HG, handgrip exercise; MAP, mean arterial pressure; PEMI, post-exercise muscle ischemia; VO2peak, peak oxygen uptake.*