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1	Characterisation of extracellular redox enzyme concentrations in response
2	to exercise in humans
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35 Abstract

36 Redox enzymes modulate intracellular redox balance and are secreted in response to 37 cellular oxidative stress, potentially modulating systemic inflammation. Both aerobic and 38 resistance exercise are known to cause acute systemic oxidative stress and inflammation; 39 however, how redox enzyme concentrations alter in extracellular fluids following bouts of either type of exercise is unknown. Recreationally active males (n=26, mean \pm SD: age 28 \pm 40 41 8 years) took part in either: 1) two separate energy-matched cycling bouts: one of moderate intensity (MOD) and a bout of high intensity interval exercise (HIIE) or 2) an eccentric-based 42 43 resistance exercise protocol (RES). Alterations in plasma (study 1) and serum (study 2) 44 peroxiredoxin (PRDX)-2, PRDX-4, superoxide dismutase-3 (SOD3), thioredoxin (TRX-1), 45 TRX-reductase and interleukin (IL)-6 were assessed before and at various timepoints after 46 exercise. There was a significant increase in SOD3 (+1.5 ng/mL) and PRDX-4 (+5.9 ng/mL) concentration following HIIE only, peaking at 30- and 60-min post-exercise respectively. 47 48 TRX-R decreased immediately and 60-min following HIIE (-7.3 ng/mL) and MOD (-8.6 ng/mL) respectively. In non-resistance trained males, no significant changes in redox enzyme 49 50 concentrations were observed up to 48 hours following RES, despite significant muscle 51 damage. IL-6 concentration increased in response to all trials, however there was no 52 significant relationship between absolute or exercise-induced changes in redox enzyme 53 concentrations. These results collectively suggest that HIIE, but not MOD or RES increase 54 the extracellular concentration of PRDX-4 and SOD3. Exercise-induced changes in redox 55 enzyme concentrations do not appear to directly relate to systemic changes in IL-6 56 concentration.

57

Abbreviations: ANOVA: Analysis of Variance, BMI: Body Mass Index, CK: Creatine 58 Kinase, ELISA: Enzyme Linked Immunosorbent Assay, EV: Extracellular Vesicle, H₂O₂: 59 60 Hydrogen Peroxide, HIIE: High Intensity Interval Exercise, IL: Interleukin, IPAQ: International Physical Activity Questionnaire, LDH: Lactate Dehydrogenase, MOD: 61 62 Moderate Intensity Exercise, NADH: reduced nicotinamide adenine dinucleotide, ONOO: Peroxynitrite, PBS: Phosphate Buffered Saline, PBSwC: Phosphate Buffered Saline Wash 63 Casein, PRDX: Peroxiredoxin, ROS: Reactive oxygen species, SD: Standard deviation, SOD: 64 Superoxide Dismutase, TLR: Toll-like Receptor, TRX: Thioredoxin, TRX-R: Thioredoxin-65 Reductase, VO_{2MAX}: Maximum oxygen consumption. 66

67 New & Noteworthy

We conducted two studies to characterise changes in redox enzyme concentrations after single bouts of exercise to investigate the emerging association between extracellular redox enzymes and inflammation. We provide evidence that SOD3 and PRDX-4 concentration increased following high intensity aerobic, but not eccentric-based resistance exercise. Changes were not associated with IL-6. The results provide a platform to investigate the utility of SOD3 and PRDX-4 as biomarkers of oxidative stress following exercise. Introduction

93 It is well documented that acute exercise perturbs cellular reduction-oxidation (redox) 94 balance through the increased production of reactive oxygen species (ROS) within actively 95 contracting skeletal muscle (34), as well as other infiltrating cell types (35). Evidence 96 suggests that ROS such as hydrogen peroxide (H_2O_2) and peroxynitrite (ONOO⁻) have 97 important roles in facilitating muscle contractile activity (25) and regulating the expression of 98 genes involved with metabolism and endogenous antioxidant protection (14, 39). Conversely, 99 heightened levels of exercise-induced H₂O₂ at the expense of antioxidant defense systems can 100 elicit oxidative stress, which may limit contractile function and promote fatigue (33). Given 101 this biphasic relationship, studies have previously evaluated alterations in redox balance in 102 response to both aerobic and resistance type exercise. These studies have primarily focused 103 on the quantification of distal markers in extracellular fluids, such as the oxidation 104 biomolecules and/or activity of antioxidant enzymes in plasma (48), serum (31), saliva (11) 105 and urine (41); highlighting exercise duration (3), intensity (17) and muscle-damage (4) as 106 factors governing greater increases. However, criticisms are commonly made with regards to 107 the direct relationship of these markers with the redox state of active tissues during exercise 108 (9). Recent evidence has highlighted that intracellular redox enzymes, such as peroxiredoxin 109 (PRDX) can be secreted from skeletal muscle myocytes (28) and immune cells (40) in 110 response to increasing concentrations of H₂O₂ in vitro. Human studies are also beginning to 111 provide evidence that plasma/ serum PRDX-2 and PRDX-4 concentrations could serve as 112 important biomarkers of intracellular redox state in the context of acute and chronic 113 inflammatory conditions (27, 40).

114 PRDXs are a major family of ubiquitous redox proteins, which modulate intracellular redox balance through a highly reactive cysteine thiolate group. The reaction rate of this 115 116 cysteine is markedly greater than any other thiol-containing protein (50), allowing rapid 117 regulation of cellular H₂O₂, with some evidence to suggest that this may facilitate muscle contraction (26). PRDXs are therefore reliable footprints of intracellular redox state, with 118 119 heightened oxidation of the PRDX cysteine indicative of oxidative stress (37). In addition, 120 upon secretion from immune cells, PRDX can directly bind to toll-like receptor (TLR)-4 to 121 initiate inflammatory cytokine production (e.g. interleukin (IL)-6) (38), providing some 122 support for the association between PRDX and inflammation (27, 40). Recent work has 123 begun to explore changes in the PRDX catalytic cycle in blood cells isolated from humans before and after acute exercise (6, 46, 47). In parallel with increases in soluble markers of 124 125 inflammation (e.g. IL-6 and C-reactive protein), an increase in the oxidation of PRDX (i.e.

126 dimer and over-oxidised states) has been reported following intensive cycling and running 127 exercise (46, 47). To our knowledge, changes in PRDX have yet to be assessed in the context 128 of exercise in humans and represents a potentially unexplored area of exercise and redox 129 biology. Interestingly, PRDX-2 can be secreted in tandem with its enzymatic reducing partners, thioredoxin (TRX-1) and thioredoxin reductase (TRX-R) (20, 40). TRX-1 and TRX-130 131 R are cysteine and selenium based-antioxidant enzymes respectively, with higher reduction 132 potentials than PRDX, thus contributing towards maintaining the antioxidant function of PRDX. In addition, the enzyme superoxide dismutase 3 (SOD3) is an extracellular 133 antioxidant released upon cellular stimulation, providing an immediate change in 134 extracellular antioxidant capacity (15, 20). Given the emerging body of literature supporting a 135 relationship between intracellular oxidative stress, redox enzyme secretion and soluble 136 137 inflammatory markers, the quantification of PRDX-2, PRDX-4, TRX-1, TRX-R and SOD3 in extracellular fluids offers the potential for accurate assessment of changes in oxidative stress 138 139 and inflammation after different types of exercise.

140 Based upon existing knowledge of the factors that can impact acute changes in 141 exercise-induced oxidative stress, we sought to perform two experiments to understand how novel markers, such as PRDX-2, PRDX-4, TRX-1, TRX-R and SOD3 respond to acute 142 143 exercise, and whether relationships exist between changes in inflammation. Specifically, we 144 aimed to characterise how these markers would be impacted by aerobic exercise intensity and 145 eccentric-based resistance exercise. We tested the hypothesis that both protocols would elicit 146 an increase in the concentrations of redox enzymes within plasma/ serum after exercise; with 147 higher exercise intensity causing a larger increase following aerobic exercise.

148

149 Methods

150 Participants

Healthy, untrained participants were recruited for two independent studies (Table 1) Participants in both studies completed the International Physical Activity Questionnaire (IPAQ), which addresses habitual levels of weekly physical activity. Participants gave their informed written consent and all studies were approved by the local Ethical Review Committee, in accordance with the Declaration of Helsinki, 2008. Participants were all nonsmokers and had not taken any antioxidant vitamin supplements or anti-inflammatory drugs for 8 weeks prior to the laboratory visits. All participants were required to refrain from any

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strenuous physical activity, consumption of alcoholic beverages or caffeine for at least twodays prior to the experimental sessions.

160

161 Experimental Sessions

The full workflow for this project is detailed in Figure 1. Experimental sessions took place in the morning (7.00 - 8.00 am start time) under stable climatic conditions (18 - 20°C and humidity between 45 – 55%) and following at least a 10-hour fast. After a period of rest, height (*Seca Alpha, Hamburg, Germany*) and mass (*Tanita, Tokyo, Japan*) were determined.

166 In study 1, participants first visited the laboratory for an assessment of cardiorespiratory fitness ($\dot{V}O_{2MAX}$) using a ramp test to exhaustion on an electromagnetically 167 168 braked cycle ergometer (Lode Excalibur Sport, Groningen, Netherlands). The protocol 169 involved commencing pedalling at 100 Watts, followed by fixed 30-Watt increments every 4 170 minutes. Oxygen uptake was assessed continuously using a breath-by-breath system (Oxycon 171 Pro, Jaeger, Wuerzberg, Germany) and heart rate monitored using a Polar Vantage heart rate 172 monitor (Polar, Kempele, Finland). The test ended when the participant reached volitional 173 exhaustion or when a plateau in oxygen consumption was observed with an increase in workload (49). A final obtained value of rate of oxygen consumption was accepted as \dot{VO}_2 174 MAX and expressed relative to body weight (mL.kg⁻¹min⁻¹). At least one week later, 175 176 participants then undertook the first of two energy and time-matched cycling trials in a 177 randomised order, at least one week apart: a continuous bout of moderate intensity cycling at approximately 60% \dot{VO}_2 (%_{MAX}) for 58 minutes (MOD) and a bout of high intensity interval 178 exercise (HIIE), consisting of 10 x 4-minute intervals at 85% \dot{VO}_2 (%_{MAX}), with 2-minute rest 179 180 intervals. In both trials, oxygen uptake was assessed continuously and power output was adjusted where necessary in order to maintain target $\dot{V}O_2$ and equal energy expenditure 181 182 between MOD and HIIE (study 1). Rating of Perceived Exertion (RPE) was monitored every 183 5 minutes throughout the trials (5).

In study 2 (n = 16), non-resistance trained males undertook an eccentric-based resistance exercise protocol adapted from a previous study by Alemany et al (1). This muscle damaging protocol was performed on a Humac Norm dynamometer (CSMI, Massachusetts, USA). The dynamometer lever arm was programmed to flex the participant's knee from a start position of 10° of flexion to 90° of flexion, thus allowing a range of motion of 80°. The participants began with their leg at the start position and were asked to maximally contract their quadriceps against a resistance while the lever arm moved to the finish position (90° 191 knee flexion). Once at the finish position they were advised to relax their leg and the 192 dynamometer moved them back to the start position to avoid a concentric contraction being 193 performed. The lever arm moved at a set speed of $60^{\circ} \cdot s^{-1}$. The bout consisted of 20 sets of 10 194 repetitions with each set being separated by 1 minute of rest. Visual feedback and verbal 195 encouragement were provided to all participants to maximise torque output for each 196 contraction.

197

198 <u>Blood sampling and Plasma Isolation</u>

For both studies, a catheter (Appleton Woods, Birmingham, UK) was inserted into the 199 200 antecubital vein of the arm prior to exercise to obtain a baseline blood sample after thirty 201 minutes of rest (Pre). The catheter was continually kept clear with isotonic saline solution 202 (0.9% sodium chloride). As indicated in Figure 1, blood samples were then taken immediately, 30 minutes and 60 minutes after both HIIE and MOD (Study 1 - Pre, Post+0, 203 204 Post+30 and Post+60) and immediately, 30 minutes, 3 hours and 48 hours following the muscle damage protocol (Study 2 - Pre, Post-0, Post-30min, Post+3hr and Post+48hr). The 205 206 post+48 hr (Study 2) blood sample was taken via venepuncture. At each time point, 12 mL of 207 blood was drawn into vacutainer tubes containing either potassium ethylene 208 diaminetetraacetic acid in study 1 (Becton, Dickson & Company, Oxford, UK) or no 209 anticoagulant in study 2. In study 1, whole blood was centrifuged at 1525g for 15 minutes, at 210 room temperature. In study 2, whole blood was allowed to clot at room temperature for 20 mins and then centrifuged at 1500g for 15 minutes. The resulting plasma (study 1) and serum 211 212 (study 2) were aliquoted and frozen at -80°C for future analysis of redox enzymes, IL-6, creatine kinase (CK) and lactate dehydrogenase (LDH). Capillary blood samples were 213 214 obtained from the earlobe after 4 min of exercise and then every 6 min thereafter (i.e. end of 215 each HIIE interval) in study 1. These samples were used for analysis of blood glucose and 216 lactate concentrations to verify intensity-dependent differences between each protocol.

217

218 Analytical Procedures

219 PRDX-2, PRDX-4. TRX-1, TRX-R and SOD3 ELISAs

ELISAs for the detection of PRDX-2, PRDX-4, TRX, TRX-R and SOD3 were developed in-house. Commercially available antigens and antibodies (i.e. PRDX-2. PRDX-4, TRX and TRX-R) were purchased from either *Abcam*, Cambridge, UK (ab) or *Sigma Aldrich*, Dorset, UK (SRP). The human SOD3 antigen and rabbit antiserum directed against human SOD3 were developed as previously described (16, 20). Plasma or serum and standards (100 225 µL) were loaded onto individual wells of an ELISA plate (Thermo Scientific F8 polysorp immune wells) and protein left to bind overnight at 4 °C. Wells were then pre-washed with 226 227 PBS wash buffer, supplemented with 0.1% casein (PBSwC, 200 µL) and then blocked with 228 1% casein in PBS (200 µL) for 30 minutes at room temperature, with gentle agitation. Anti-229 human rabbit antibodies for PRDX-2 (ab133481, 1:2000), PRDX-4 (ab59542, 1:2000) and 230 SOD3 (in-house, 1:2000), and anti-human mouse antibodies for TRX-1 (ab16965, 1:8000) 231 and TRX-R (ab16847, 1:1000) were then added to each well, diluted in PBSwC for 45 232 minutes at room temperature. Following this, 100 µL of anti-rabbit (1:5000) or anti-mouse (1:500) IgG Biotin antibodies in PBSwC, and streptavidin-horseradish peroxidase (1:2000 in 233 PBSwC) were added separately to each well, both for 45 minutes, with gentle agitation. 234 Between all stages, all wells were washed three times with PBSwC. Finally, 100 µL of 235 236 3,3',5,5'-tetramethylbenzidine (10ug) was added per well, and the plate left to develop in the 237 dark for 15-25 minutes. Stop solution (1.5mM H₂SO₄, 50 µL) was then added to each well 238 and absorption at 450nm subsequently evaluated by using a plate reader (Multiskan Ascent, 239 Thermo Labsystems). Concentration of each antigen was then determined by comparing 240 absorbance values of recombinant PRDX-2 (ab167977, Abcam), PRDX-4 (ab93947, Abcam), TRX-1 (ab51064, Abcam), TRX-R (SRP6081, Sigma Aldrich) and SOD3 (in-house) proteins 241 242 (0-50 ng/mL). ELISA validation experiments showed no cross-reactivity of the PRDX-2, 243 PRDX-4, TRX-1, TRX-R and SOD3 antibodies with the respective antigens, nor with serum 244 albumin. All values were adjusted for plasma volume, according to previous methods (12).

245

246 Other Analyses

247 In both studies, a cytometric bead array was used to quantify plasma (study 1) and 248 serum (study 2) IL-6 concentrations on a BD C6 Accuri Flow Cytometer (BD Biosciences, 249 Berkshire). In study 1, blood lactate and glucose concentrations were determined 250 immediately following collection using an automated lactate and glucose analyser (Biosen C-251 Line Clinic, EKF-diagnostic GmbH, Barleben, Germany). In study 2, serum CK and LDH 252 concentrations were determined to monitor muscle damage using an automated ABX Pentra 400 system (Horiba UK Ltd, UK). Haematocrit and haemoglobin concentrations were used to 253 254 ascertain plasma volume changes and make appropriate adjustments in plasma redox enzyme and IL-6 concentrations (Beckman Coulter, London, UK). 255

256

257 Statistical Analysis

258 The Shapiro Wilk test was used to test for normality in scale data at all time points. 259 Differences between participant characteristics and the physiological responses to exercise in 260 both studies were assessed using unpaired samples T-tests or non-parametric Mann-Whitney 261 U Tests. The influence of exercise on plasma/ serum PRDX-2, PRDX-4, SOD3, TRX-1, 262 TRX-R and IL-6 concentration was assessed over time by repeated-measures analysis of 263 variance (ANOVA) or non-parametric Wilcoxon signed rank tests, depending variable 264 normality. Post hoc analysis of any significant effect of time or interaction effect (study 1; Group*Time) was performed by a test of simple effects by pairwise comparisons, with 265 Bonferroni correction. Effect sizes for main effects and interaction effects of ANOVA are 266 presented as partial eta² (η^2_p), using Cohen's definition of η^2_p of 0.01, 0.06 and 0.14 for 267 'small', 'medium' and 'large' effects respectively (10). Pearson correlation and Spearman 268 269 rank were used to assess the relationship between parametric and non-parametric data 270 respectively. All values are presented as means ± standard deviation or error (indicated 271 throughout manuscript). Statistical significance was accepted at the p < .05 level. Statistical analyses were performed using SPSS (PASW Statistics, release 23.0, SPSS Inc., Chicago, IL, 272 273 USA).

274

275 **Results**

There was no significant difference in age or BMI between the participants taking part in the two studies, Participants in study 1 (p = 0.004) had significantly higher self-reported physical activity than in study 2.

279

280 Acute physiological responses to HIIE and MOD

For study 1, the physiological responses during each exercise bout are reported in Table 2. Peak \dot{VO}_2 and RPE were significantly greater in HIIE compared to MOD (p < 0.00001), but there were no statistically significant differences in mean \dot{VO}_2 and energy expenditure. Whole blood lactate and glucose data are reported in Table 2. Mean lactate concentration was significantly higher during HIIE than MOD (p < 0.0001), but there was no significant difference in average glucose concentration between trials.

287

288 Effects of eccentric-based resistance exercise on muscle damage markers

Changes in markers of muscle damage are reported in Table 3. A stepwise increase
(Post+48hr > Post+3hr > Post+30min > Post+0 > Pre) in serum CK concentration was

observed over time, peaking above Pre at Post+48hr (p > 0.001). Serum LDH concentration was elevated above Pre at all post-exercise timepoints (p < 0.05), also increasing Post+3hr and Post+48hr, relative to Post+30min (p < 0.05).

294

295 Effects of aerobic and eccentric-based resistance exercise on IL-6 concentration

296 IL-6 data is presented in Figure 3. In study 1, plasma IL-6 increased in both trials 297 (Time effect: F (3) = 15.5, p < 0.0001, $\eta^2 = 0.66$), being elevated above resting values, both 298 immediately (p = 0.004) and Post+30 (p = 0.002), but not Post+60 (Figure 3A). The 299 magnitude of this increase was significantly greater Post-Ex in HIIE (p = 0.031), than MOD (Time x Condition effect: F (3) = 7.0, p < 0.001, $\eta^2 = 0.47$). IL-6 concentration decreased 300 301 Post+30 (p = 0.004) and Post+60 (p = 0.007), relative to Post+0, and Post+60, relative to 302 Post+30 (p = 0.026) in HIIE only. In study 2 (Figure 3B), IL-6 concentration was significantly higher at all timepoints up to three hours, but not 48 hours after exercise, 303 relative to Pre (Time effect: F (4) = 14.3, p < 0.0001, $\eta^2 = 0.30$). 304

305

306 Effects of aerobic exercise on PRDX-2, PRDX-4, TRX-1, TRX-R and SOD3 concentration

307 No differences were observed in resting concentrations of PRDX-2, PRDX-4, TRX-1, 308 TRX-R or SOD3 when quantified in plasma and serum across all trials. Changes in plasma 309 PRDX-2, PRDX-4, TRX-1, TRX-R and SOD3 in response to MOD and HIIE are reported in 310 Figure 2A. There was a significant increase in plasma SOD3 (Trial x Time Effect: F(3,1) =5.3, p = 0.028, η^2 = 0.31) and PRDX-4 following HIIE only (non-parametric tests: all p < 311 312 0.05). SOD3 concentration was elevated above pre-exercise values at all post-HIIE 313 timepoints, peaking at Post+0 (p = 0.015) and Post+30 (p = 0.013), but only significantly 314 higher than MOD at Post+30 (p = 0.05). Plasma SOD3 concentration decreased relative to 315 Post+30 at Post+60 (p = 0.013). Relative to Pre, PRDX-4 concentration increased at Post+30 (p = 0.015) and Post+60 (p = 0.008) following HIIE, with PRDX-4 concentration higher at all 316 317 post-exercise timepoints compared with MOD (p < 0.038). There was a significant decrease in plasma TRX-R concentration in both MOD and HIIE. Relative to Pre, TRX-R significantly 318 319 decreased at Post+0 in HIIE only (p = 0.021), with values significantly less than MOD (p =320 0.011). Following MOD, TRX-R was significantly lower at Post+60, relative to all 321 timepoints (all p < 0.038). There were no statistically significant changes in PRDX-2 and TRX-1 concentration over time in either trial; however, TRX-1 concentration was 322 323 significantly higher in HIIE than MOD Post+60 only (p = 0.021).

324

325 *Effects of eccentric-based resistance exercise on PRDX-2, PRDX-4, TRX-1, TRX-R and* 326 *SOD3 concentration*

Serum redox enzyme concentration changes in response to an eccentric-based resistance exercise protocol are presented in Figure 2B. A trend was observed for a decrease in PRDX-2 concentration Post+30min (-1.12 ng/mL), however this did not reach statistical significance (Time effect: F (4) = 2.3, p = 0.065, $\eta^2 = 0.13$). Similarly, no significant changes were noted in PRDX-4, TRX-R or SOD3 up to 48 hours following eccentric-based resistance exercise. A significant increase in TRX-1 was shown Post+48hr, relative to Post+30min (p = 0.039), but not Pre (p = 0.309).

334 335

336 Discussion

337 The current results have characterised the kinetic responses of endogenous redox enzymes within the extracellular environment after exercise for the first time. We highlight 338 339 novel findings that high intensity aerobic cycling induces a significant increase in SOD3 and PRDX-4 in healthy, untrained males. Similar responses were not observed following 340 341 moderate intensity cycling or muscle damaging resistance exercise. In contrast, plasma TRX-342 R concentration decreased within one hour following moderate and high-intensity cycling 343 exercise, but not resistance exercise. Taken together these findings provide novel insights into 344 the regulation of extracellular redox enzymes in response to exercise.

345 The current data highlights modality and exercise-intensity specific increases in two 346 abundant redox enzymes. In response to aerobic exercise, PRDX-4, but not PRDX-2 347 concentration increased thirty minutes following HIIE and remained elevated until Post+60. 348 The secretory pathways of PRDXs are isoform specific, with endoplasmic reticulum (ER, i.e. PRDX-4) and cytosolic (i.e. PRDX-2) resident isoforms released via classical and non-349 350 classical secretory pathways respectively (8). The current data therefore suggests that 351 exercise may activate the ER-golgi pathway to secrete PRDX-4 in an intensity-dependent 352 manner. SOD3, which is also released via this pathway, increased more rapidly than PRDX-4 353 following HIIE (Post+0), with levels tailing off Post+60, relative to Post+30. SOD3 is an 354 antioxidant enzyme released directly from the cell membrane (15, 20), specifically secreted during exercise to metabolise superoxide anions produced in the extracellular environment to 355 H₂O₂ (30). The different peak concentrations of SOD3 (i.e. Post+0) and PRDX-4 (i.e. 356

357 Post+30) following HIIE may be explained, in part, by a) the membrane proximity of SOD3 358 compared to the ER location of PRDX-4 and b) the appearance of superoxide anions first in the extracellular space following exercise, before their metabolism to H₂O₂, which then 359 360 induced PRDX-4 secretion. This may also be reflective of differential secretion rates of 361 SOD3 and PRDX-4 from various tissues during and following exercise. Both proteins are 362 expressed in skeletal muscle (19), a highly redox active tissue (36); however, PRDX-4 is 363 primarily located in pancreas, liver and heart (21), whereas SOD3 is expressed in the heart and vasculature tissue (42). The association with the vasculature may explain the more rapid 364 365 increase in plasma SOD3 concentration following HIIE. Aside from these increases, a modest decrease was observed in plasma TRX-R after both MOD and HIIE (study 1), with this 366 change being much more rapid in HIIE (Post+0), compared to MOD (Post+60). The 367 368 mechanisms driving a decrease in TRX-R after exercise are unclear at present. The decrease 369 may represent transient homeostatic fluctuations involving uptake of redox enzymes by 370 neighboring cells and tissues, perhaps to regulate intracellular redox balance (23).

371 A finding that was in contrast to our hypothesis was that eccentric-based resistance 372 exercise did not induce an increase in the extracellular concentrations of redox enzymes. The 373 measurement of redox enzymes in plasma and serum is an emerging area of biomedical 374 research, particularly in the context of acute (24) and chronic (13, 43) inflammatory 375 conditions, where PRDXs and TRX-1 have been associated with enhanced cytokine and 376 chemokine production (22, 38). The participants in both studies were relatively inactive, with 377 participants in study 2 in particular, reporting significantly lower levels of habitual physical 378 activity (Table 2) and being unaccustomed to eccentric-based resistance exercise. 379 Unaccustomed eccentric exercise induces significant amounts of acute muscle damage and 380 inflammation (7), as demonstrated by the stepwise increases in CK and LDH concentrations 381 up to 48 hours following our protocol, and IL-6 up to 3 hours post-exercise (Figure 3B). These data suggest that the increase in SOD3 and PRDX-4 observed in study one is unlikely 382 383 due to just a disruption to the plasma membrane, given that no changes were observed 384 following a muscle-damaging bout of resistance exercise. It must be acknowledged that only 385 selective timepoints were measured following the protocol, and perhaps the secretion of 386 redox enzymes occurs between 3- and 48-hours post-exercise. Nevertheless, this study has 387 highlighted for the first time that redox enzyme concentrations do not match that of established markers of muscle damage and inflammation when measured in serum samples 388 389 following an eccentric-based resistance exercise bout. In response to aerobic-based exercise,

390 we have recently demonstrated a positive association between intracellular peroxiredoxin (I-391 IV) over-oxidation in immune cells and plasma IL-6 concentration (47). In the current study, 392 IL-6 concentration increased in an intensity-dependent manner (HIIE > MOD) following 393 aerobic exercise (Figure 3A); however, there were no statistically significant relationships 394 between absolute or exercise-induced changes in PRDX-4 and SOD3 with IL-6. The 395 observations across both studies therefore suggest no relationship between that IL-6 and 396 redox enzymes after exercise. A larger sample size may be needed to adequately address 397 these associations and support the previously documented relationship between plasma/ 398 serum redox enzymes and soluble inflammatory markers (27, 40).

399 The results of the current investigation demonstrate clear differences in the changes in 400 SOD3, TRX-R and PRDX-4 following aerobic vs. eccentric-based resistance exercise. With 401 regards to PRDX-2 and TRX-1, no changes were observed following aerobic or eccentric-402 based resistance exercise. Both PRDX-2 and TRX-1 are cytosolic redox enzymes that contain 403 no N-terminal signal peptide for secretion and thus are released via non-classical pathways, 404 associated with extracellular vesicles (EVs), such as exosomes and nanoparticles (45). 405 PRDX-2 and TRX-1 are detectable in plasma/ serum samples through their association with 406 the exofacial surface of the EV membrane (18, 44); however, their protein levels may be 407 higher due to protein contained within the EVs. This protein would not be detectable by 408 antibodies when enclosed within the lipid membrane during ELISA quantification, as 409 previously shown (32). Indeed, recent evidence has highlighted that a series of leaderless 410 redox enzymes (i,e, PRDX-1, PRDX-2, PRDX-5, PRDX-6, TRX-1, SOD1 and SOD2) are 411 secreted in EVs via a non-classical route following exposure to stress, with classically 412 secreted SOD3, TRX-R and PRDX-4 not detectable within EVs (2). This may explain why plasma/ serum PRDX-2 and TRX-1 concentration did not significantly change following 413 414 muscle-damaging or aerobic exercise. It must be noted that TRX-1 concentration was 415 significantly higher 48 hours after the eccentric-based resistance exercise protocol, relative to **Post+0** (study 2) and also significantly higher at Post+60 in HIE, compared to MOD (study 416 417 1). These findings again underpin intensity-dependent differences, despite in both cases, concentrations not being higher than pre-exercise values. In response to a far more extreme 418 419 bout of exercise, Marumoto et al, (2010) reported a marked increase in TRX-1 levels 420 $(17.9 \pm 1.2 \text{ ng/mL} \text{ at baseline to } 70.1 \pm 6.9 \text{ ng/mL})$ after a 2-day 130km ultra-endurance marathon (29); however, these exercise bouts were substantially different in nature and thus 421 422 hard to directly compare. Even though an Ultramarathon is accompanied by significant amounts of muscle damage, given the findings of study 2, it is unlikely that muscle damage is
the primary cause of TRX-1 secretion in this context. Further work is needed to clarify
whether TRX-1 and PRDX-2 protein levels alter within EVs after conventional bouts (i.e. not
ultra-endurance) of muscle-damaging and aerobic-based exercise.

427 This study has quantified the responses of antioxidant enzymes in the extracellular environment following acute exercise in age and BMI matched individuals from two 428 429 independent exercise studies (Table 1). We must acknowledge that the studies would have benefited from a direct comparison between redox enzyme concentrations and other 430 431 established biomarkers of oxidative stress (e.g. protein carbonyls and F2-isoprostanes). However, due to limited sample volume this analysis was not feasible and should therefore be 432 prioritised as an area of future research. A second limitation is that the quantification of redox 433 434 enzymes and IL-6 were undertaken in both plasma (study 1) and serum (study 2); however, 435 there were no differences in any of these proteins when quantified in pre-exercise samples.

436

437 Conclusion

438 The results of the present study have highlighted that plasma SOD3 and PRDX-4 439 concentration increased in response to acute exercise. Importantly, the secretion of these proteins appears to be intensity and modality dependent, with increases only observed in 440 441 response to high intensity aerobic cycling in untrained individuals. A decrease in TRX-R was also noted following different aerobic exercise bouts, with exercise intensity driving a more 442 443 rapid decrease in TRX-R. Future research is required to pinpoint the precise mechanisms governing the secretion and uptake of redox enzymes, and their role in regulating redox 444 445 balance between tissues after exercise.

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455

456 **Conflict of Interest**

457 None of the authors declare a conflict of interest.

458

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Tables

	Energy-matched Trials (study 1)	Eccentric-based Resistance Exercise (study 2)	Statistical Analysis
Number of Participants	9	16	n/a
Age (years)	29 ±	25 ±	$\mathbf{P} = \mathbf{NS}$
	5	9	
Body Mass Index	24.2 ±	25.3 ±	D NG
(kg/m ²⁾	3.4	4.1	$\mathbf{P} = \mathbf{NS}$
IPAQ (METs-	6683 ±	2540 ±	⁺ P = 0.005
min/week)	3835*	2022	$^{+}P = 0.004$
	3.4 ±		
Watt Max (Watt/kg)	0.5		n/a
[.] VO _{2 max}	44.5 ±		n/a
(mL.kg ⁻¹ .min ⁻¹)	6.4		11/ a

Table 1. Demographics for participants in studies 1 and 2.

Grey boxes indicate missing data.

* Indicates significant difference in comparison to study 2: *P < 0.05, **P < 0.001.

NS P > 0.05.

	Statistical Analysis		
Trial	Continuous cycling for 58 min, predicted 60% \dot{VO}_{2}_{MAX} (MOD)	10 x 4 min cycling intervals, predicted 85% $\dot{VO}_{2 \text{ MAX}}$ (2 min rest intervals. Total time = 58 min, HIIE)	
Mean VO2 MAX (%)	56.5 ± 2.6	58.9 ± 4.3	P = NS
Energy Expenditure (kJ)	2077 ± 340	2072 ± 339	P = NS
Average RPE	12 ± 1	$16 \pm 1^{***}$	****P < 0.0001
Mean Blood Lactate (mmol/L)	1.9 ± 0.6	6.8 ± 1.4	****P<0.0001
Mean Blood Glucose (mmol/L)	3.9 ± 0.3	4.5 ± 0.6	$\mathbf{P} = \mathbf{NS}$

Table 2. Physiological response to aerobic-based exercise (study 1).

* Indicates a significant difference between MOD and HIIE: *** P < 0.0001.

NS P > 0.05.

Table 3. Changes in markers or muscle damage following eccentric-based resistance exercise (study 2). Values are means \pm standard deviation.

	Pre	Post+0	Post+30min	Post+3hr	Post+48hr
Creatine Kinase	147.6 ±	236.1 ±	289.9 ±	560.8 ±	575.9 ±
(Units/ L)	27.1	65.5 *	86.0 *+	273.5 **+#	290.8 **+#\$
Lactate Dehydrogenase	$254.9 \pm$	282.7 ±	274.1 ±	290.3 ±	299.9 ±
(Units/ L)	130.6	70.9 *	77.1 *	77.8 *+	165.2 *+

* Indicates significant difference in comparison to Pre: P < 0.05, P < 0.001.

+ Indicates significant difference in comparison to Post+0: *P < 0.05.

Indicates significant difference in comparison to Post+30min: #P < 0.05.

\$ Indicates significant difference in comparison to Post+3hr: P < 0.05.

Figure 1: Schematic of the two exercise studies. Dark lines represent the exercise session, with lighter lines indicating pre- and post-exercise resting periods. Gaps between dark lines indicate the rest periods during the HIIE trial. Blood samples taken for each study are indicated as arrows.

Figure 2: Changes in redox enzyme concentration in response to two energy-matched cycling bouts (A) - moderate steady state (MOD - black bars) and high intensity interval exercise (HIIE – white bars) and an eccentric-based resistance exercise protocol (B): PRDX-2, PRDX-4, TRX-1, TRX-R and SOD3. Values are means \pm standard error. For Figure 2A: * indicates significant differences relative to Pre: * p<.05. # indicates a significant difference relative to Post+0: # p<.05. \$ indicates a significant difference relative to Post+30: \$ p<.05. + indicates a significant difference between MOD and HIIE: + p<.05. For Figure 2B: ^ indicates a significant difference between Post+30min and Post+48hrs timepoints.

Figure 3: Changes in plasma IL-6 in response to two energy-matched cycling bouts (A): moderate steady state (MOD - black bars) and high interval exercise (HIIE – white bars) and an eccentric-based resistance exercise protocol (B). Values are means \pm standard error. For Figures 3A and 3B: * indicates significant differences relative to Pre: * p<.05; ** p<.001. # indicates a significant difference relative to Post+0: # p<.05. \$ indicates a significant difference between MOD and HIIE: + p<.05.