1	Effect of dietary nitrate supplementation on conduit artery blood flow, muscle
2	oxygenation, and metabolic rate during handgrip exercise
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22 Abstract

Dietary nitrate supplementation has positive effects on mitochondrial and muscle contractile 23 24 efficiency during large muscle mass exercise in humans, and on skeletal muscle blood flow (O)25 in rats. However, concurrent measurement of these effects has not been performed in humans. Therefore, we assessed the influence of nitrate supplementation on \dot{Q} and muscle oxygenation 26 27 characteristics during moderate (40% peak) and severe (85% peak) intensity handgrip exercise in 28 a randomized, double-blind, crossover-design. Nine healthy men (age: 25 ± 2 yrs) completed four 29 constant-power exercise tests (two per intensity) randomly assigned to condition (nitrate-rich 30 (Nitrate) or nitrate-poor (Placebo) beetroot supplementation) and intensity (40% peak or 31 85% peak). Resting mean arterial pressure was lower after Nitrate compared to Placebo (84±4 vs 32 89 ± 4 mmHg; p<0.01). All subjects were able to sustain 10 min of exercise at 40% peak in both 33 conditions. Nitrate had no effect on exercise tolerance during 85% peak (Nitrate: 358±29, 34 Placebo: 341 ± 34 s; p=0.3). Brachial artery \dot{Q} was not different after Nitrate at rest or any time 35 during exercise. Deoxygenated-[hemoglobin+myoglobin] was not different for 40% peak 36 (p>0.05), but was elevated throughout 85% peak (p<0.05) after Nitrate. The metabolic cost (\dot{VO}_2) was not different at end exercise, however, the $\dot{V}O_2$ primary amplitude at the onset of exercise 37 38 was elevated after Nitrate for the 85% peak work rate (96 ± 20 vs 72 ± 12 ml/min; p<0.05) and had a faster response. These findings suggest that an acute dose of Nitrate reduces resting blood 39 pressure and speeds $\dot{V}O_2$ kinetics in young adults, but does not augment \dot{Q} or reduce steady-state 40 41 $\dot{V}O_2$ during small muscle mass handgrip exercise.

42 Keywords: Beetroot juice, NIRS, Kinetics, Dynamic exercise

43 New and Noteworthy

We show that acute dietary nitrate supplementation via beetroot juice increases the amplitude and speed of local muscle $\dot{V}O_2$ on-kinetics parameters during severe- but not moderate-intensity handgrip exercise. These changes were found in the absence of an increased blood flow response, suggesting the increased $\dot{V}O_2$ was attained via improvements in fractional O₂ extraction and/or spatial distribution of blood flow within the exercising muscle.

49

50 Introduction

51 Dietary nitrate supplementation is well documented to have positive effects during large 52 muscle mass exercise in humans (4, 33, 38, 48, 50). These effects include lowering oxygen 53 consumption ($\dot{V}O_2$) (4, 32, 38, 42), speeding $\dot{V}O_2$ kinetics (3, 6, 31, 32) and reducing the ATP 54 cost of work (2, 26) during submaximal exercise, which may translate to the enhanced exercise 55 tolerance found during severe intensity exercise (6, 31, 37, 50). The precise mechanism(s) for 56 these effects still remains uncertain, but they are facilitated through the reduction of the dietary 57 nitrate to nitrite by commensal bacteria in the mouth (40). Once absorbed into the circulatory 58 system, nitrite is readily converted to nitric oxide (NO) in hypoxic (16, 47) and acidic (41) 59 environments, which are expected to be present at the exercising muscle.

Nitric oxide is a potent vasodilator (20, 22, 45); as such it had been proposed that nitrate supplementation may augment blood flow (\dot{Q}) to active muscle. This was first experimentally investigated in rats during submaximal treadmill running (23, 24). These authors found that nitrate supplementation resulted in an increased \dot{Q} to the hindlimb, particularly to muscles composed of greater percentages of type II fibers. These findings demonstrate that nitrate may change the regulation of \dot{Q} relative to $\dot{V}O_2$, as these two variables generally increase in

66	proportion to one another across a range of exercise intensities (1, 43). Recently the effect of
67	nitrate on \dot{Q} was investigated in human subjects (5, 12, 34), but no change in brachial artery
68	blood flow (\dot{Q}_{BA}) was found in healthy, young men and women during light-to-moderate
69	intensity handgrip exercise. These previous studies might not have recruited type II fibers in the
70	younger subjects due to lower intensity exercise, potentially missing the preferential effects of
71	dietary nitrate on higher order fiber types (for review see (30)). It should be noted, two of the
72	aforementioned studies did find improvements in compromised populations (i.e., older adults in
73	hypoxia (12) and 'noncompensators' (5)).

The Importantly, these previous studies in humans using nitrate (12, 34), provided no measure of $\dot{V}O_2$ or fractional O_2 extraction (which can be estimated noninvasively via deoxygenated-

76 [hemoglobin + myoglobin] (deoxy-[Hb + Mb]) and used to estimate $\dot{V}O_2$) (7, 18, 19, 35).

Moreover, the measurements were made after fixed durations of moderate intensity submaximal exercise and during the steady state, leaving the effects of nitrate on local muscle $\dot{V}O_2$ during the exercise onset transient unknown. Since faster $\dot{V}O_2$ kinetics are associated with a reduction in the O₂ deficit (and thus accumulation of fatigue inducing metabolites), these findings carry important implications for patient populations, such as chronic heart failure (CHF), as accumulating evidence suggests nitrate supplementation may be effective for enhancing quality of life through improvements in exercise and/or daily activity tolerance (21, 25, 51).

Therefore, the purpose of this investigation was to resolve whether acute supplementation of nitrate preferentially provided positive effects in small muscle mass exercise during severe intensity exercise, where type II fibers would be recruited and greater hypoxic and acidic muscle environment exists. Specifically, we tested the hypotheses that with nitrate supplementation compared to placebo: 1) \dot{Q}_{BA} would be significantly elevated during severe, but not moderate

intensity exercise; 2) $\dot{V}O_2$ would be elevated during severe intensity exercise and display faster kinetics; and 3) tolerance of exercise (T_{lim}) would be increased during severe intensity exercise.

91

92 Materials and Methods

93 Ten healthy, recreationally active men volunteered for the investigation (mean \pm SD: age: 25 ± 2 yrs; height: 178 ± 4 cm; body mass: 80 ± 10 kg; BMI: 25 ± 3 kg/m²). All experimental 94 95 procedures in the present study were approved by the Institutional Review Board at Kansas State 96 University and conformed to the standards set forth by the Declaration of Helsinki. Prior to 97 participation in the study, all subjects were informed of the protocol, any possible health risks, as 98 well as the probable benefits of the study. All subjects provided written informed consent to 99 participate and completed a medical health history questionnaire to ensure absence of any known 100 cardiovascular or metabolic diseases which would preclude them from the study.

101

102 Experimental Protocol

103 All testing sessions were performed on a custom-built, two-handed handgrip ergometer 104 previously described by our laboratory (7, 8). Briefly, the subjects were seated in an upright 105 position at arm's length from the ergometer with the hands pronated at heart level and directly in 106 front of their torso. All sessions were performed utilizing a 50% duty-cycle (1.5 s contraction, 107 1.5 s relaxation) and fixed 4 cm linear displacement that was maintained via audio cues. All 108 subjects were familiarized with the exercise, audio cues, and duty-cycle prior to the first testing 109 session. During the first visit, subjects performed an incremental test for the determination of peak power (P_{peak}) starting at 1 Watt (W) and increasing at a rate of 0.5 W·min⁻¹. The test was 110 111 performed until volitional exhaustion or after three consecutive contraction cycles in which the

subject was unable to maintain the correct tempo or complete full contractions. P_{peak} was 112 113 recorded as the highest power obtained in which the subjects completed at least 30 s of the stage. 114 The four subsequent visits were randomly assigned to 40 or 85 % P_{peak} (two tests per 115 intensity) and supplemental condition (see Supplementation below; Figure 1). The four constant 116 power tests were performed for 10 min or until exhaustion for 40 and 85 % P_{peak}, respectively. 117 The coefficient of variation for tolerance of exercise (T_{lim}) at 85 % P_{peak} intensity in our 118 laboratory is ~7% (8). All testing sessions were separated by 48 - 72 h and subjects were asked 119 to abstain from vigorous activity, food, and caffeine prior to testing for 12, 3, and 2 h, 120 respectively. Upon arrival to the laboratory, the subjects sat quietly for 15 min, after which 121 resting blood pressure measurements and subsequent plasma samples were obtained (See Figure 122 1). All exercise tests were performed at approximately the same time of day (\pm 1.5 h for each 123 subject) between 1000 and 1500 hours.

124

125 Supplementation

126 The exercise testing sessions were randomly assigned to nitrate or placebo beetroot supplementation conditions (one of each per intensity; i.e., nitrate + 40 % P_{peak} and placebo + 40 127 128 %P_{peak}), creating a randomized, double-blind, crossover study design. In each condition, the 129 subjects consumed beetroot concentrate (2 x 70 ml providing ~13 mmol nitrate) or nitrate-130 depleted beetroot concentrate placebo (2 x 70 ml providing ~0.006 mmol nitrate; both Beet It 131 Sport, James White Drinks, Ipswich, UK). Subjects consumed the shots on their own outside of 132 the laboratory ~2.5 h before testing began to allow for maximal expression of plasma nitrite 133 concentrations ([nitrite]) (49, 50) (See Figure 1). This dose of nitrate was chosen because it was 134 shown to increase T_{lim} with no greater effects seen at higher doses (50). During the study,

135 subjects were asked to abstain from using mouthwash (29) and toothpaste or chewing gum that 136 contained triclosan, as these products serve to reduce the oral bacteria needed to facilitate the 137 conversion of nitrate to nitrite. Each exercise testing session was separated from the others by at 138 least 48 h to allow plasma [nitrite] adequate time to return to pre-supplementation concentrations 139 (50). Subjects were asked to maintain their normal diet with the exception of limiting foods high 140 in nitrate, such as spinach and arugula (39). No subjects reported taking any multivitamins or 141 anti-oxidant supplements. All subjects self-reported compliance with the supplemental protocol. 142 No subjects reported gastrointestinal discomfort; however, when subjects reported typical 143 symptoms (i.e., beeturia or red stools) they were assured this was a typical side effect of the 144 nitrate supplementation.

145

146 Measurements

Venous blood samples (5-6 ml) were separated into 1.5 ml Eppendorf tubes containing 5
µl heparin (concentration 1000U/ml) and centrifuged at 3250 rpm at 4 °C for 5 min within 1 min
of withdrawal. Plasma samples were then pipetted into separate Eppendorf tubes, flash frozen in
liquid nitrogen, and stored at -80 °C until later analysis.

The measurements of plasma nitrate and nitrite were performed within 30 min of thawing via chemiluminescence with a NO analyzer (NOA 280i, Sievers Instruments, Boulder, CO, USA). In order to obtain plasma nitrite levels and to avoid potential reduction of nitrate, potassium iodide in acetic acid was used as a reductant. This reductant has the ability to reduce nitrite to NO but is incapable of reducing higher oxides of nitrogen (i.e., nitrate), thus increasing the specificity for nitrite. Plasma nitrate concentrations were obtained using the same apparatus with the stronger reductant vanadium chloride in hydrochloric acid at a temperature of 95 °C.

This stronger reductant reduces the sum of all nitrogen oxides with an oxidation state of +2 or higher, which is predominately nitrate (μ M), but also includes both nitrite (nM) and nitrosothiols (nM).

161 Resting blood pressure was measured in the left arm using an automated patient monitor 162 (S/5 Light Monitor type F-LM1-03, Datex-Ohmeda General Electric, Finland) which makes use 163 of the oscillometric technique. To increase accuracy, this machine utilizes a 3-lead ECG to 164 monitor heart rate. This measurement was taken in triplicate and a mean value was obtained. 165 Exercising blood pressure was taken from the left ankle using the same patient monitor while the 166 subject was seated at the handgrip ergometer. During the measurement, subjects were asked to 167 remain still and allow their leg to relax. A correction factor (pressure = measured pressure – 168 (distance between the heart and ankle in meters x 76 mm Hg) was used to adjust for the 169 increased hydrostatic pressure present between the ankle and heart (27). Pilot work performed in 170 our lab validated the correction factor with measurements taken from the ankle and arm at heart 171 level. This pilot work also revealed that the increase in blood pressure during 85 % P_{peak} handgrip 172 exercise exceeded the capabilities of the equipment to accurately measure ankle pressure so 173 pressure was only obtained for the 40 % P_{peak} intensity.

The raw blood velocity profiles were measured in the right brachial artery using Doppler ultrasound (Vivid 3, GE Medical Systems, Milwaukee, WI, USA) operating in pulse wave mode at a Doppler frequency of 4.0 MHz with a phased linear array transducer probe operating at an imaging frequency of 6.7 MHz, and were stored for *post-hoc* analysis. For all testing sessions the Doppler gate was set to the full width of the brachial artery to ensure complete insonation and all Doppler velocity measurements were corrected for the angle of insonation, which was adjusted to be less than 60°. Measurements were made at least 3 cm above the antecubital fossa to avoid

bifurcation of the brachial artery. Brachial artery diameters were measured in the transverse axisusing two-dimensional sonography.

183 Muscle and microvascular oxygenation status were measured noninvasively using a 184 frequency-domain multi-distance near infrared spectroscopy (NIRS) system (OxiplexTS, ISS, 185 Champaign, IL, USA) positioned over the belly of the left *flexor digitorum superficialis* (FDS). 186 Details of this technique have been described previously (7, 11). Briefly, this device consists of 187 one detector fiber bundle and eight light-emitting diodes (LED) operating at wavelengths of 690 188 and 830 nm (four LEDs per wavelength). The LED-detector fiber bundle separation distances are 189 2.0, 2.5, 3.0, and 3.5 cm. This NIRS device measures and incorporates the reduced scattering 190 coefficient (μ_s'), measured dynamically, to provide absolute concentrations (μM) for deoxy-[Hb 191 + Mb] and total-[Hb + Mb]. The NIRS probe was calibrated prior to each test according to the 192 manufacturer's specifications. The belly of the FDS of the left arm was identified using palpation 193 and EMG. The NIRS probe was secured along the belly of the FDS and was wrapped with an 194 elastic bandage to prevent shifting of the probe. The placement of the NIRS probe was marked 195 with permanent ink for reproducible positioning throughout the study. The NIRS data were 196 collected at 50 Hz and stored for *post-hoc* analysis.

197 The $\dot{V}O_2$ (ml $O_2 \cdot min^{-1}$) of the FDS was calculated for each minute of exercise using the 198 technique described previously (7), which integrates deoxy-[Hb + Mb] and \dot{Q}_{BA} . It was assumed 199 that the deoxy-[Hb + Mb] signal reflects exclusively deoxy-[Hb] [we acknowledge that the 200 signal contains deoxy-[Mb] as well (17)] and that the entire signal arises only from the muscle 201 (i.e., not from any interposing adipose or skin tissue). With these assumptions the deoxy-[Hb] 202 may be converted into an estimated $\dot{V}O_2$. The deoxy-[Hb] values are in units of µmole heme/*l* 203 tissue, where the tissue is assumed to be muscle. These deoxy-[Hb] units can be converted into

204 µmole heme/l blood using the conversion 1.36% capillary blood volume/muscle volume [derived from 400 cap/mm², 28.3 μ m² CSA, and a coefficient of 1.2 correcting for tortuosity and 205 206 branching of the capillaries (44)]. These units can then be converted into mole O_2/l blood 207 assuming 1 mole O_2 /mole heme and further to $l O_2/l$ blood using the conversion 22.4 $l O_2$ /mole 208 O₂. \dot{V} O₂ values in l O₂/min may then be obtained by multiplying this value by the measured \dot{Q}_{BA} values. This calculation was performed with the understanding that $\dot{Q}_{\rm BA}$ likely overestimates \dot{Q} 209 210 through the capillaries under the NIRS probe. However, because the same calculation (and 211 subsequent assumptions) was used across subjects and the primary comparison was within 212 subjects, the error associated with this assumption was minimized. Further, these assumptions 213 were held constant across both supplemental conditions.

214

215 Data analysis

Mean blood velocity (\dot{V}_{mean} ; cm·s⁻¹) was defined as the time-averaged mean velocity over 216 each 3 s contraction cycle. \dot{Q}_{BA} (ml·min⁻¹) was calculated using the product of \dot{V}_{mean} and vessel 217 cross-sectional area (CSA = πr^2). CSA (cm²) was calculated each minute of exercise using 218 brachial artery diameters measured at the beginning of each minute. The \dot{Q}_{BA} data were analyzed 219 220 using three consecutive contraction cycles (i.e., 9 s) for rest and at the end of each minute of 221 exercise. The NIRS data were first multiplied by 4 to convert the values from hemoglobin 222 equivalents back to total heme units (15) and were subsequently analyzed using 1 s mean values 223 that were converted to 30 s mean bins for resting values and 9 s time-binned mean values at the 224 end of each minute of exercise and at exhaustion. Systolic blood pressure (SBP) and diastolic 225 blood pressure (DBP) were measured at least three times at rest and once every 2 min during

exercise and were then used to calculate MAP. Vascular conductance (VC) (ml·min⁻¹·(100 mmHg)⁻¹) was calculated using the quotient of \dot{Q}_{BA} /MAP, multiplied by 100. Kinetics analyses were conducted for the $\dot{V}O_2$ data using 6 s time-binned mean values over the initial 120 s of exercise and 9 s time-binned mean values at 180 and 240 s with a monoexponential model:

231
$$y(t) = y(b) + A(1 - e^{-(t - TD)/\tau})$$

where y(t) is the \dot{VO}_2 at any point in time, y(b) is the baseline before the onset of exercise, *A* is the primary amplitude of the response, *TD* is the time delay proceeding the increase in, and τ is the time constant. The rate constant (RC) was calculated as *A* divided by τ (giving ml/min/s) to give an indication of the acceleration of the response.

236

237 Statistical analysis

238 All curve fitting and statistical analyses were performed using a commercially available 239 software package (SigmaPlot 12.5, Systat Software, San Jose, CA, USA). Differences in resting values and Tlim were analyzed using Student's paired t-tests. Differences within condition (i.e., 240 241 40% nitrate and 85% nitrate) for resting plasma [NO_x] and MAP were compared and if no 242 differences were found, these values were averaged to represent the mean resting value for that condition. Exercising values (i.e., \dot{Q}_{BA} , deoxy-[Hb + Mb], total-[Hb + Mb], and $\dot{V}O_2$) were 243 244 analyzed using two-way ANOVAs with repeated measures (supplement x time) using Tukey's 245 *post hoc* tests when main effects were detected. \dot{VO}_2 kinetics parameters were analyzed using 246 two-way ANOVA with repeated measures (supplement x work rate) using Tukey's post hoc tests 247 when main effects were detected. Differences were considered significant when p < 0.05. Data 248 are presented as means \pm standard error unless otherwise noted.

249

250 **Results**

251 Ten subjects completed the protocol. One subject was determined to be an outlier based 252 on their 85 % P_{peak} T_{lim} change score (-285 s) being more than 3 SD outside the group mean 253 change score (15 ± 84 s). This subject was removed from all data analyses.

254

255 **Plasma [nitrate] & [nitrite] and resting blood pressure**

Plasma [nitrate] and [nitrite] was measured in six participants (see *Limitations* section below for explanation). Plasma [nitrate] was elevated 26-fold over placebo after acute nitrate supplementation ($784 \pm 32 \text{ vs } 29 \pm 2 \mu \text{M}$, p < 0.001). All subjects demonstrated elevated plasma [nitrite] after acute nitrate supplementation ($456 \pm 60 \text{ vs } 68 \pm 7 \text{ nM}$, p < 0.001, Fig. 2) resulting in a 5.7-fold increase over placebo. Resting blood pressure values are presented in Table 1. Acute nitrate supplementation was associated with a lowering of SBP, DBP, and MAP by 7%, 4%, and 6%, respectively (all p < 0.05) compared to placebo.

263

264 **40 %P**_{peak} exercise

The mean power for 40 % P_{peak} was 2.2 ± 0.1 W. All subjects were able to sustain 10 min of exercise at 40 % P_{peak} in both conditions. \dot{Q}_{BA} increased rapidly from exercise onset in both conditions before approaching a steady-state of approximately 260 ml·min⁻¹ by 240 s. \dot{Q}_{BA} was not different after nitrate supplementation at rest or at any time during exercise compared to placebo (Fig. 3). MAP was measured during exercise in eight of nine subjects. There was no main effect of nitrate on MAP during exercise compared to placebo (p = 0.11, Fig. 4), although MAP was 4 mmHg lower on average throughout exercise before reaching the peak values 90 ± 4 and $98 \pm 5 \text{ mmHg}$ (p = 0.02) for nitrate and placebo, respectively. There was no effect of nitrate on VC (p = 0.14, Fig. 4); both groups increased to end exercise values of 314 ± 58 and 279 ± 28 ml/min/100 mmHg (p = 0.08) for nitrate and placebo, respectively.

275 Deoxy-[Hb + Mb] increased following exercise onset in both conditions, with no 276 differences between conditions. End exercise deoxy-[Hb + Mb] was not different between nitrate 277 and placebo $(154 \pm 15 \text{ vs } 156 \pm 19 \text{ }\mu\text{M}, \text{ }p = 0.83, \text{ Fig. 5})$. Total-[Hb + Mb] was not different after 278 nitrate supplementation at any min during exercise or at the end of exercise compared to placebo 279 $(408 \pm 15 \text{ vs } 402 \pm 25 \text{ }\mu\text{M}, \text{ }p = 0.76, \text{ Fig. 5})$. $\dot{V}O_2$ was not different at any min during exercise or 280 at the end of exercise (73.1 ± 16.7 vs 75.8 ± 18.0 ml/min, p = 0.68, Fig. 6). The results of the 281 $\dot{V}O_2$ kinetics analysis are presented in Table 2 (n = 7).

282

283 85 %Ppeak exercise

The mean power for 85 % P_{peak} was 4.7 ± 0.2 W. Nitrate had no effect on T_{lim} compared to placebo (358 ± 29 vs 341 ± 34 s, p = 0.3, Fig. 7). \dot{Q}_{BA} was not different at rest or any time during exercise after nitrate supplementation. \dot{Q}_{BA} increased at exercise onset and attained end exercise values of 368 ± 42 and 353 ± 46 ml/min (p = 0.56, Fig. 3), for nitrate and placebo, respectively.

289 Deoxy-[Hb + Mb] was not different at rest and increased at exercise onset in both 290 conditions, with nitrate elevated over placebo for time points preceding end exercise (60 - 180 s,291 p < 0.05), but not 240 s (p = 0.08). At T_{lim}, nitrate and placebo were different (203 ± 26 vs 180 ± 292 19 μ M, p = 0.03, Fig. 5). Total-[Hb + Mb] was not different after nitrate supplementation, both 293 conditions showed a progressive increase toward the end exercise values (447 ± 30 vs 440 ± 31 294 μ M, p = 0.65). $\dot{V}O_2$ increased 897 ± 183% and 838 ± 191% (p = 0.83) from rest to T_{lim} for nitrate and placebo, respectively. There was no difference for end exercise $\dot{V}O_2$ after nitrate supplementation (112 ± 12 vs 107 ± 14 ml/min, p = 0.62, Fig. 6). The results of the $\dot{V}O_2$ kinetics analysis are presented in Table 2 (n = 7). Both supplemental conditions had significantly higher primary amplitudes during 85% P_{peak} compared to 40% P_{peak} (p < 0.05). Nitrate supplementation also increased the primary amplitude within 85% P_{peak} (p = 0.02) and reduced the time constant (τ ; p = 0.04) compared to placebo.

301

302 Discussion

303 The present study investigated the effects of acute nitrate supplementation on conduit 304 artery Q concurrently with local muscle microvascular oxygenation characteristics during 305 moderate and severe intensity handgrip exercise. The acute dosage utilized (~13 mmol nitrate), 306 elevated plasma [nitrite] more than 5-fold higher than that seen with placebo and was associated 307 with reductions in blood pressure at rest of 4-8%. Contrary to our first hypothesis, nitrate had no 308 effect on \dot{Q}_{BA} at rest or any time point during moderate or severe intensity handgrip exercise 309 compared to placebo. The primary novel finding of the present study, in agreement with our 310 second hypothesis, was that the $\dot{V}O_2$ primary amplitude was elevated and the kinetics were faster 311 after nitrate during severe intensity handgrip exercise consequent to an increased O₂ extraction 312 (deoxy-[Hb + Mb]). Additionally, nitrate had no effect on T_{lim} when exercise was performed in 313 the severe intensity domain.

314

315 Effect on control of blood flow

Ferguson and colleagues (23, 24) discovered that nitrate supplementation increased bulk hindlimb \dot{Q} in rats with the largest effect in muscles composed of a high percentage of type IIb

318 and IIx fibers (23, 24). To date, the previous studies (5, 12, 34) and the present investigation that 319 directly measured Q in young healthy humans during small muscle mass (handgrip) exercise, 320 have been unable to replicate the findings of Ferguson et al. (23, 24) or Cosby et al. (14). The 321 work of Kim and colleagues (34) had young healthy subjects perform rhythmic exercise under 322 both nitrate and placebo conditions; however the work done was performed at fairly low work rates. The greatest \dot{Q}_{BA} achieved in the work of Kim et al. (34) was approximately 200 ml/min 323 324 for both supplementations, which was lower than the \dot{Q}_{BA} measured in the present investigation 325 at 40 % P_{peak} (~260 ml/min). If dietary nitrate does in fact have preferential effects in high order 326 fiber types, it is likely that Kim et al. (34) and the lower work rate in the present study did not 327 recruit said fibers.

328 The other two studies (5, 12) and the present investigation performed higher intensity 329 exercise that increased the likelihood of recruiting higher order fibers. However, in agreement with the lower intensity data, nitrate supplementation had no effect on the steady state \dot{Q}_{BA} in 330 331 healthy young subjects. The present investigation advanced these previous studies by measuring 332 the dynamic response during the onset of exercise. While there was no difference in the speed of the \dot{Q}_{BA} adjustment to exercise, there was evidence of improved O₂ delivery within the 333 334 exercising muscle (see *Effect on tissue oxygenation and* \dot{VO}_2 *below*). Casey and colleagues (12) 335 attempted to maximize the stimulus for nitrite conversion to NO (and thus maximize the potential augmentation of \dot{Q}_{BA}) by putting their subjects in hypoxia, but there was still no 336 337 difference between nitrate and placebo. The study by Bentley and colleagues (5) used a 338 hydrostatic challenge to alter O_2 delivery and found no differences in the absolute Q_{BA} following nitrate supplementation. These authors did find there was less attenuation of \dot{Q}_{BA} induced by the 339 340 hydrostatic challenge following nitrate, which the authors attributed to an increased

341 compensatory vasodilation (5). The supine exercise model used in these aforementioned studies 342 (5, 12, 34) differed from the present investigation in that our subjects were seated upright with 343 the arms at heart level. It has been shown that the seated posture increases muscle sympathetic 344 activity and reduced central venous pressure compared to the supine posture (9). Since the 345 present findings are largely in agreement with these previous studies (5, 12, 34) and handgrip 346 exercise is not limited by cardiac output, these postural differences were likely inconsequential. 347 Nevertheless, a recent study found that acute nitrate supplementation increased peak 348 cardiac output and VO_2 in CHF patients with preserved ejection fraction during a supine peak 349 incremental exercise test (51). This study was not designed to resolve the spatial distribution of 350 the ~10% increase in cardiac output. If nitrate does favorably affect VC and \dot{Q} to type II fibers, 351 as suggested by Ferguson and colleagues (24), the increased reliance on type II fibers with CHF, 352 and other diseases (28, 46) supports the notion that nitrate supplementation may be more 353 effective in these populations with O₂ delivery challenges. The discovery that nitrate can increase 354 \dot{Q}_{BA} in older adults in hypoxia (12) further bolsters this hypothesis.

355

356 Effect on tissue oxygenation and $\dot{V}O_2$

Larsen and colleagues (38) were the first to show that a dietary nitrate salt supplement could reduce the $\dot{V}O_2$ associated with a given work rate. Subsequent studies utilizing beetroot supplementation have yielded mixed results across a variety of exercise modalities, with some showing ~3-5% reductions in $\dot{V}O_2$ (2, 4, 37, 42, 48, 50), and others no change (6, 13, 31, 32) after supplementation. NIRS-derived variables measured concurrently with $\dot{V}O_2$ paralleled the change in $\dot{V}O_2$ when it occurred (4, 6). 363 Given the above, attempting to interpret the present findings in the context of whole body 364 exercise is difficult. In the present investigation, deoxy-[Hb + Mb] was elevated after nitrate 365 supplementation throughout severe-, but not moderate-, intensity exercise. Moreover, total-[Hb + 366 Mb] was not impacted by nitrate supplementation during both exercise intensities. Changes in 367 total-[Hb + Mb] from rest to exercise are thought to reflect the change in microvascular 368 hematocrit (17). To the best of our knowledge, the current study is the first to observe an 369 increased primary amplitude, exercising level, and T_{lim} value for deoxy-[Hb + Mb] after nitrate 370 supplementation. It should be noted that Breese et al. (6) reported a higher value on beet root 371 juice across the transition from moderate to severe exercise due to faster kinetics, but no 372 differences in the amplitude or T_{lim} were observed (6). Increased deoxy-[Hb + Mb] relative to 373 unchanged total-[Hb + Mb] (and \dot{Q}_{BA}), suggests an increased fractional O₂ extraction. The 50% 374 duty-cycle used in the present investigation has been shown to mechanically constrain \dot{Q}_{BA} and 375 VO_2 during severe intensity handgrip exercise (7), such that the present changes should be 376 viewed as positive and suggest improvements in the microvascular distribution of O₂ rather than 377 a decrease in efficiency. Nitrate may facilitate the delivery of O₂ to regions that were otherwise 378 under perfused in this exercise model. This improved O₂ extraction was manifest in the kinetic 379 response (see discussion below); however, there was no difference in $\dot{V}O_2$ at the end of exercise 380 after nitrate supplementation in the present investigation. This implies that the efficiency of the 381 work was neither positively nor negatively impacted and the improvements in fractional O₂ 382 extraction were likely obscured by the mechanical limitations of the exercise. Future work could 383 usefully attempt to elucidate if there is a 'threshold' type effect of the duty cycle used for the 384 exercise (i.e., employing 20, 30, 40% duty cycles in the severe intensity domain).

385

386 Effect on $\dot{V}O_2$ kinetics parameters and tolerance to exercise

No differences in the end exercise amplitude of local muscle $\dot{V}O_2$ were found in the 387 present study, but kinetics analyses revealed that the initial amplitude of $\dot{V}O_2$ was increased 388 389 during exercise at 85% P_{peak} after nitrate supplementation. Nitrate supplementation resulted in a 390 faster τ and a substantially greater rate constant (amplitude/ τ ; ~73% increase). These findings are 391 in agreement with the speeding of pulmonary $\dot{V}O_2$ kinetics shown during whole body exercise in 392 instances of compromised O₂ delivery and/or recruitment of higher order Type II muscle fibers 393 (3, 6, 31, 32) and the equivalent microvascular PO₂ response in rats (23). The present 394 investigation is the first to show significant differences in kinetics parameters after an acute 395 dosage of nitrate, where other studies used chronic supplementation to see effects (3, 6, 31). 396 However, these improvements in $\dot{V}O_2$ amplitude and speed of adjustment did not lead to 397 improved T_{lim} in the present investigation. Previous work has found that speeding $\dot{V}O_2$ kinetics 398 during whole body exercise does not always result in improvements to T_{lim} (10, 36), indicating 399 that the relationship between these two variables is not a simple relationship. Indeed, interactions 400 between VO_2 kinetics and other physiological parameters are likely requisite to see 401 improvements in exercise tolerance. Had there been a summation of the improved kinetics across 402 multiple transitions (such as that seen during daily activity), a greater sparing of the O_2 deficit 403 could result in an accumulated improvement.

404

405 Limitations

406 We acknowledge that the method used to estimate $\dot{V}O_2$ herein utilizes several 407 assumptions (7) and likely overestimated the $\dot{V}O_2$. We contend that these assumptions, held 408 constant throughout, should not obscure an impact of nitrate on $\dot{V}O_2$. Additionally, our sample

409 size was small and as such could have resulted in the present investigation not being sufficiently 410 powered to detect differences in some variables (e.g., exercising MAP and VC). Finally, we did not measure plasma [nitrite] in all nine subjects (access to the NOA was precluded during later 411 412 data collection); however, the three subjects without this measurement exhibited similar 413 differences in blood pressure to the six with plasma [nitrite] measurements. It should be noted 414 that no pre-dose blood pressure measurements were made in the present investigation. However, 415 each subject served as their own control and thus had two separate days of blood pressure 416 measurements for each condition (i.e., nitrate and placebo), increasing our confidence that nitrate 417 influenced blood pressure herein.

418

419 Conclusions

420 The present study reaffirmed previous findings that an acute dose of nitrate is associated 421 with lower SBP, DBP, and MAP in healthy, young men. The acute dose was also an effective method to increase and speed the local muscle $\dot{V}O_2$ on-kinetics parameters during severe 422 intensity handgrip exercise, primarily through an increased fractional O₂ extraction rather than 423 424 increased blood flow. However, the ergogenic effects associated with nitrate supplementation 425 (i.e., improved tolerance to exercise) during large muscle mass exercise were not seen when the 426 exercise was performed in small muscle mass handgrip exercise. These findings warrant future 427 studies investigating the effects of nitrate supplementation during the dynamic adjustment at the 428 onset of exercise in populations at risk of O₂ delivery impairment and reduced NO 429 bioavailability.

430

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439	No conflicts of interest, financial or otherwise, are declared by the author(s).
440	

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- 603
- 604

U		
	Placebo	Nitrate
SBP (mmHg)	130 ± 4	121 ± 4 †
DBP (mmHg)	69 ± 4	66 ± 5 *
MAP (mmHg)	89 ± 4	84 ± 4 *

Table 1. Resting Blood Pressure After Acute Dietary Nitrate Supplementation

SBP, DBP, and MAP denote systolic blood pressure, diastolic blood pressure, and mean arterial pressure, respectively. Values are expressed as means \pm SE. \dagger significantly different from placebo (p < 0.01), * significantly different from placebo (p < 0.05)

605

606

Table 2. $\dot{V}O_2$ Kinetics Parameters for the Onset of Handgrip Exercise

Baseline (ml/min) 17 ± 3 16 ± 4 Amplitude (ml/min) 52 ± 13 64 ± 8 τ (s) 38 ± 5 34 ± 9 TD (s) 1 ± 1 3 ± 1 RC (ml/min/s) 1.6 ± 0.4 1.9 ± 0.6 85% P _{peak} $85\% P_{peak}$ Baseline (ml/min) 15 ± 4 17 ± 2 Amplitude (ml/min) $72 \pm 16 \dagger$ $99 \pm 22 * \dagger$ τ (s) 37 ± 8 $25 \pm 3 *$ TD (s) 4 ± 2 6 ± 2 RC (ml/min/s) 2.2 ± 0.4 $3.8 \pm 0.6 *$	40% P _{peak}	Placebo	Nitrate
τ (s) 38 ± 5 34 ± 9 TD (s) 1 ± 1 3 ± 1 RC (ml/min/s) 1.6 ± 0.4 1.9 ± 0.6 85% P _{peak} 85% P _{peak} 17 ± 2Baseline (ml/min) $72 \pm 16 \dagger$ $99 \pm 22 * \dagger$ τ (s) 37 ± 8 $25 \pm 3 *$ TD (s) 4 ± 2 6 ± 2	Baseline (ml/min)	17 ± 3	16 ± 4
TD (s) 1 ± 1 3 ± 1 RC (ml/min/s) 1.6 ± 0.4 1.9 ± 0.6 85% P_{peak}15 \pm 4 17 ± 2 Baseline (ml/min) $72 \pm 16 \dagger$ $99 \pm 22 * \dagger$ τ (s) 37 ± 8 $25 \pm 3 *$ TD (s) 4 ± 2 6 ± 2	Amplitude (ml/min)	52 ± 13	64 ± 8
RC (ml/min/s) 1.6 ± 0.4 1.9 ± 0.6 85% P_{peak}15 \pm 4 17 ± 2 Baseline (ml/min) $72 \pm 16 \ddagger$ $99 \pm 22 * \ddagger$ τ (s) 37 ± 8 $25 \pm 3 *$ TD (s) 4 ± 2 6 ± 2	τ (s)	38 ± 5	34 ± 9
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	TD (s)	1 ± 1	3 ± 1
Baseline (ml/min) 15 ± 4 17 ± 2 Amplitude (ml/min) $72 \pm 16 \ddagger$ $99 \pm 22 \ddagger$ τ (s) 37 ± 8 $25 \pm 3 \ddagger$ TD (s) 4 ± 2 6 ± 2	RC (ml/min/s)	1.6 ± 0.4	1.9 ± 0.6
Amplitude (ml/min) $72 \pm 16 \ddagger$ $99 \pm 22 * \ddagger$ τ (s) 37 ± 8 $25 \pm 3 *$ TD (s) 4 ± 2 6 ± 2	85% P _{peak}		
τ (s) 37 ± 8 $25 \pm 3 *$ TD (s) 4 ± 2 6 ± 2	Baseline (ml/min)	15 ± 4	17 ± 2
TD (s) 4 ± 2 6 ± 2	Amplitude (ml/min)	72 ± 16 †	99 ± 22 *†
	τ (s)	37 ± 8	25 ± 3 *
RC (ml/min/s) 2.2 ± 0.4 $3.8 \pm 0.6 *$	TD (s)	4 ± 2	6 ± 2
	RC (ml/min/s)	2.2 ± 0.4	3.8 ± 0.6 *

 τ , TD, and RC denote time constant, time delay, and rate constant, respectively.

Values are expressed as means \pm SE. * significantly different from placebo within work rate, † significantly different from 40% P_{peak} within supplement (both p < 0.05). Analysis completed on 7 of 9 subjects.

608 Figure Legends

609 Figure 1. Schematic representation of experimental protocol

- 610 Left: overall protocol showing the timing of the five laboratory visits in relation to one another.
- 611 **Right:** expansion of an individual testing day (in this case, Testing day #1; each subsequent
- testing session followed the same timeline). Each testing session was assigned randomly to the
- 613 supplemental condition (i.e., nitrate or placebo) and exercise intensity (i.e., 40 or 85 % P_{peak}). All
- 614 exercise tests began approximately 2.5 h after supplement consumption.
- 615

616 Figure 2. Resting nitrite concentrations

- 617 Plasma nitrite concentration ([nitrite]) for individual subjects (gray lines) and group means (both,
- n = 6; see text for discussion of reduced subject number). Plasma nitrate concentrations were
- 619 similar to [nitrite], these data are presented in text only. Error bars represent SE. * significantly
- 620 different from placebo (p < 0.001).
- 621

622 Figure 3. Brachial artery blood flow during exercise

- 623 A: Mean brachial artery blood flow (\dot{Q}_{BA}) at the end of each minute of 40 % P_{peak} exercise. B: 624 Mean \dot{Q}_{BA} at the end of each minute of 85 % P_{peak} exercise and the limit of exercise tolerance
- (T_{lim}) . In both graphs, filled circles represent placebo and open circles represent nitrate
- 626 supplementation (both, n = 9). Error bars represent SE.
- 627

628 Figure 4. Mean blood pressure and vascular conductance responses to 40 %P_{peak} exercise

629 A: Mean arterial pressure (MAP) taken every 120 s during exercise. B: Vascular conductance

630 (VC) calculated as the product of brachial artery blood flow and MAP every 120 s during

631 exercise. In both graphs, filled circles represent placebo and open circles represent nitrate

- 632 supplementation (both, n = 8). Error bars represent SE.
- 633

634 Figure 5. NIRS-derived muscle and microvascular oxygenation responses during exercise

635 Left: 40 %P_{peak} exercise A: Mean deoxygenated-[hemoglobin + myoglobin] (deoxy-[Hb +

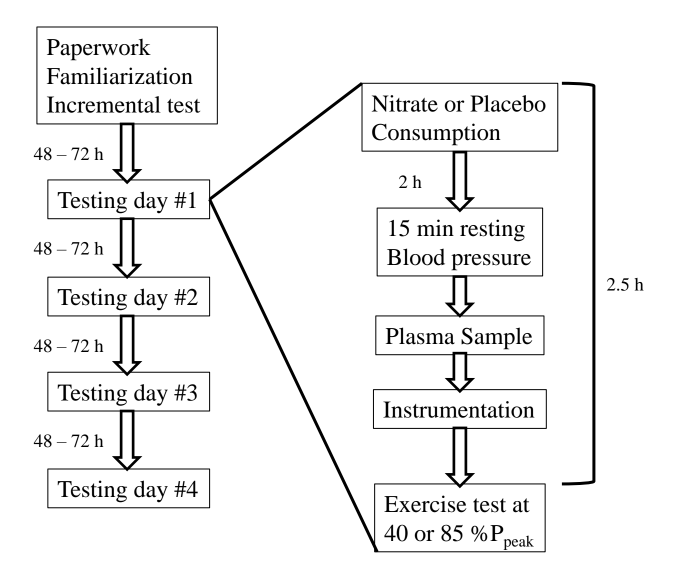
- Mb]) at the end of each minute of exercise. **B**: Mean total-[hemoglobin + myoglobin] (total-[Hb
- 637 + Mb]) at the end of each minute of exercise (both, n = 9). **Right: 85 %**P_{peak} exercise C: Mean
- 638 deoxy-[Hb + Mb] at the end of each minute of exercise and at the limit of exercise tolerance
- 639 (T_{lim}). **D**: Mean total-[Hb + Mb] at the end of each minute of exercise and at T_{lim} . In all graphs,
- 640 filled circles represent placebo and open circles represent nitrate supplementation (both, n = 9).
- Error bars represent SE. * significantly different from placebo (p < 0.05).
- 642

643 Figure 6. Estimated VO₂ during exercise

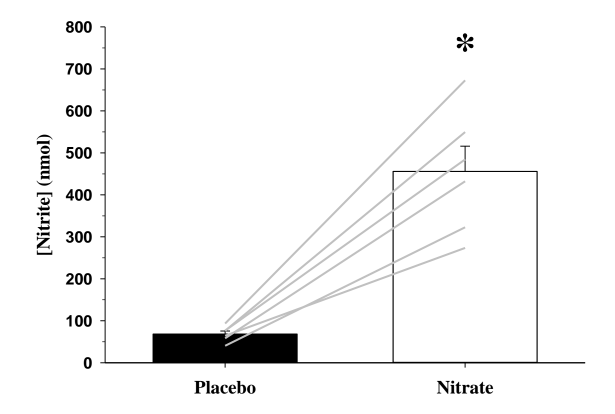
- 644 **A**: Mean estimated $\dot{V}O_2$ at the end of each minute of 40 % P_{peak} exercise. **B**: Mean estimated $\dot{V}O_2$
- at the end of each minute of 85 P_{peak} exercise and at the limit of exercise tolerance (T_{lim}). In

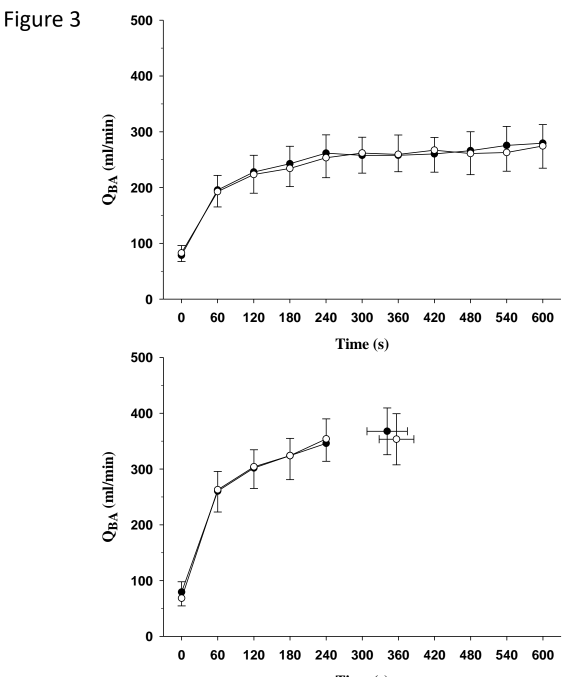
- both graphs, filled circles represent placebo and open circles represent nitrate supplementation
- 647 (both, n = 9). Error bars represent SE.
- 648
- 649 **Figure 7. Effect of supplementation on tolerance to exercise**
- 650 Individual (solid gray lines) and mean (n = 9) tolerance to exercise (T_{lim}) responses under both
- 651 supplementations during 85 % P_{peak} exercise. Error bars represent SE.
- 652

Figure 1

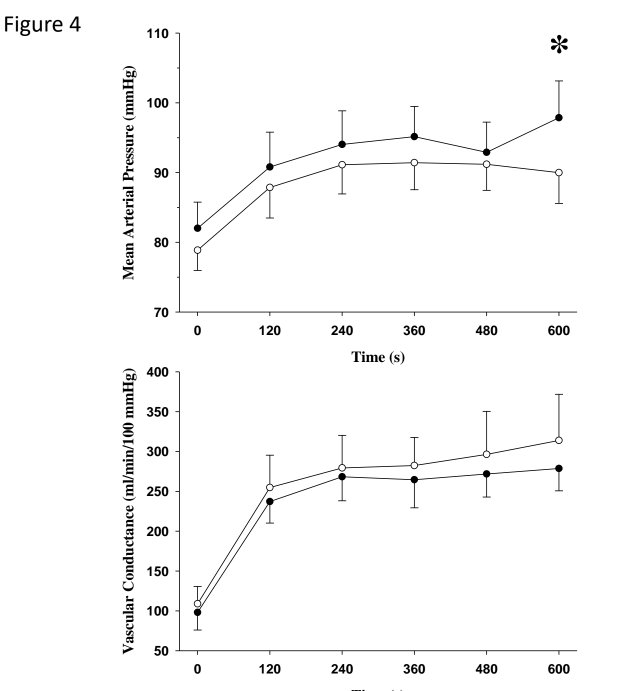




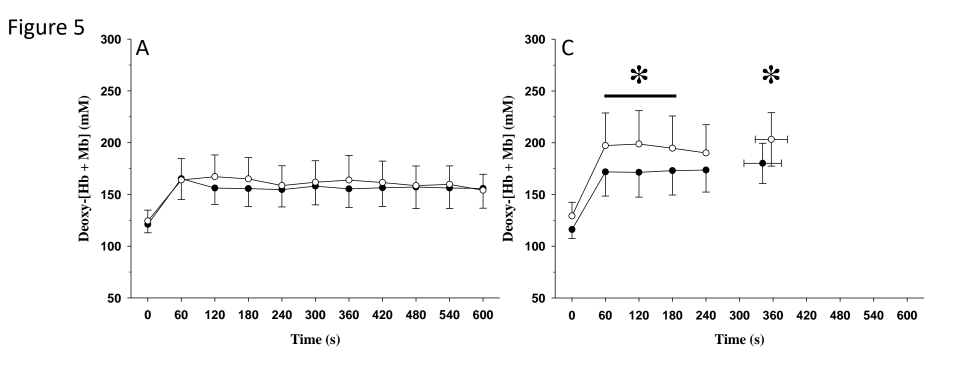


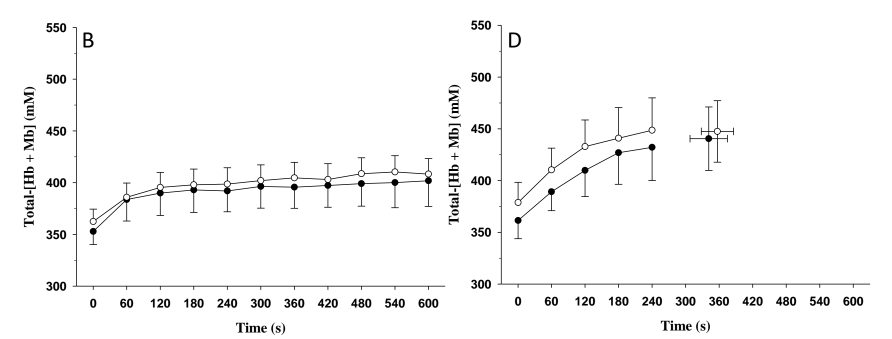


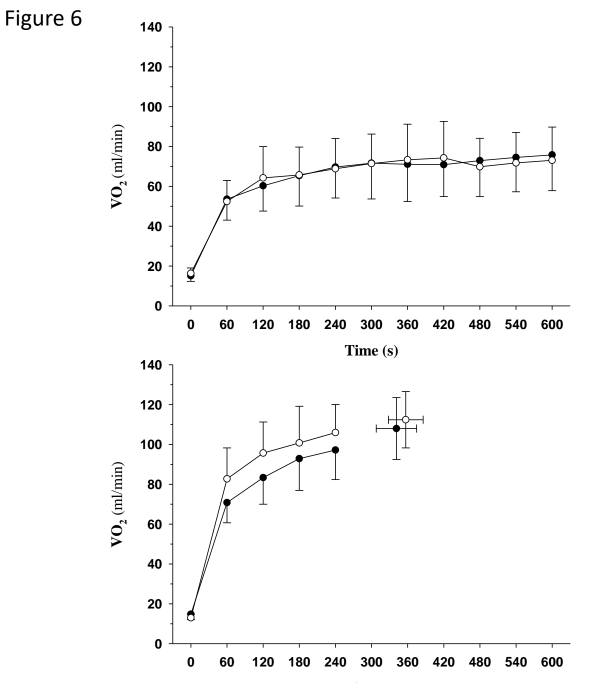
Time (s)



Time (s)







Time (s)

Figure 7

