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High dose Nitrate ingestion does not improve 40 km cycling time trial performance in trained cyclists

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1	High Dose Nitrate Ingestion Does Not Improve 40 km Cycling Time Trial Performance in
2	Trained Cyclists
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5 Abstract

6 This study evaluated the chronic effects of nitrate (NO_3) ingestion over three days, on 40 km 7 TT performance in 11trained cyclists (VO_{2max}: 60.8 \pm 7.4 ml.kg⁻¹.min⁻¹; age: 36 \pm 9 years; 8 height: 1.80 ± 0.06 m; body mass: 87.2 ± 12.0 kg). Utilising a double blind randomised cross-9 over design, participants completed three 40 km TT on a Velotron® ergometer following the ingestion of either a 140 ml of "BEET It sport®" NO₃- shot containing 12.8 mmol or 800 mg of 10 11 NO_3 , a placebo drink or nothing (control). Performance, oxygen consumption (VO₂), blood bicarbonate (HCO3-), pH and lactate (BLa) and ratings of perceived exertion (RPE) were 12 measured every 10 km throughout the TT. The present findings show that NO₃⁻ ingestion had 13 no effect on TT performance (NO₃: 4098.0 \pm 209.8 vs. Placebo: 4161.9 \pm 263.3 s, p = 0.296, 14 15 ES = 0.11), or VO₂ (p = 0.253, ES = 0.13). Similarly, blood lactate and RPE were also 16 unaffected by the experimental conditions (p = 0.522, ES = 0.06; p = 0.085, ES = 0.30) respectively. Therefore, these results suggest that a high dose of NO_3^{-1} over three days has 17 limited efficacy as an ergogenic aid for 40 km TT cycling performance in trained cyclists. 18

19 Introduction

20 Intake of exogenous dietary nitrates (NO_3) through supplementation and whole foods are 21 associated with beneficial cardiovascular health outcomes and are used as an ergogenic aid 22 for athletic performance (Clements et al., 2014). Green leafy vegetables and root vegetables 23 are examples of NO₃⁻ rich dietary sources, while commercially available beetroot supplements 24 provide a highly concentrated and practical method to ingest NO_3 prior to exercise. The 25 underlying action of NO_3 is derived from the enhanced bioavailability of nitric oxide (NO), 26 which can be synthesised via continuous reactions in the L-arginine-NO pathway (Moncada 27 and Higgs, 1993) or reduced from NO_{3} (Lundberg et al., 2004). The latter is prominent in 28 deriving NO from dietary beetroot supplementation, with NO_3^{-1} reduced to nitrite (NO_2) in the 29 oral cavity (Duncan et al., 1995). Thereafter it is reduced to NO during downstream reactions 30 and absorbed into circulating plasma (Lundberg et al., 2004). Nitric oxide consequently 31 diffuses into the vascular smooth muscle where it acts as a gaseous signalling molecule for numerous physiological processes. 32

33

34 Nitric oxide is predominantly cited as a potent vasodilator (Kelm and Schrader, 1990); but is 35 also central to the processes of muscle contraction (Reid, 2001), mitochondrial respiration (Brown, 1999) and immunoregulation (Bogdan, 2001). Indeed, the effect on these 36 37 physiological systems has led to NO_{3⁻} rich supplements demonstrating beneficial effects on 38 the oxygen cost of moderate intensity exercise (Larsen et al., 2010) and gross efficiency 39 (Larsen et al., 2007). Therefore, mediating an improvement in mitochondrial efficiency (Larsen et al., 2011) and reducing the ATP demand of muscular contractions during exercise (Bailey 40 et al., 2010). Cermak et al. (2012a) found a significantly improved 10 km TT performance 41 42 (after 60 min of submaximal exercise) and power output compared to the placebo condition. 43 Given these promising physiological responses mediated by NO₃, the ergogenicity of 44 concentrated beetroot supplementation is translated into improved time to exhaustion (TTE) 45 and time trial (TT) performance by some authors (Lansley et al., 2011; Bailey et al., 2009; 46 Larsen et al., 2010; Cermak et al., 2012a). This beneficial effect is most apparent during

47 shorter duration performance bouts, where an intensity equivalent to the severe intensity 48 domain is undertaken (Bailey et al., 2009; Kelly et al., 2014). Time trial events of short 49 durations (4 - 16 km), that require a relatively high exercise intensity, have also reported an 50 ergogenic benefit (Lansley et al., 2011 Cermak et al., 2012a); however, no studies have 51 investigated the effects of this dietary supplement on 40 km TT performance.

52

53 Despite the positive outcomes, the overall meta-analytic effect of NO₃ on TT performance is 54 considered to be unclear (standardised mean difference \pm confidence interval: -0.1 \pm 0.06; 55 McMahon et al., 2016). This differential effect could be attributed to the training status of 56 participants and/or the intensity of the bout not eliciting a sufficient physiological demand 57 (Jones, 2014). Indeed, no effects have been observed during prolonged (> 30 min) TT 58 performance from beetroot juice supplementation within highly trained individuals with VO_{2max} 59 of >60 ml·kg⁻¹·min⁻¹ (Cermak et al., 2012b; Wilkerson et al., 2012; Lane et al., 2013). Wilkerson 60 et al., (2012) administered 500 ml beetroot juice containing 6.2 mmol of NO_3^- prior to an 80 61 km TT, however the participants exhibited only a small 25% increase in plasma NO₂ compared 62 to the 139% increases observed in earlier studies with a similar participant cohort with an 63 identical dose (Lansley et al., 2011). Indeed, a strong negative correlation between plasma NO_2 increase and TT performance improvement was reported (r = -0.85; p = 0.01); thus, 64 65 highlighting the importance of a high bioavailable NO for performance improvements. 66 However, two separate doses 12 hrs apart, increasing total NO₃⁻ ingestion to 17.4 mmol prior 67 to exercise, also reported no ergogenic effects during a 43.8 km TT (Lane et al. 2013). These authors, however, utilised a superiorly trained participant cohort (VO_{2max}: 71.6 \pm 4.6), which 68 69 may explain the lack of improvement. Therefore, on the basis of previous work, the aim of this 70 study was to evaluate the chronic effects of nitrate (NO₃) ingestion over three days, on 40 km TT performance in trained cyclists. 71

72

73 Methods

Eleven trained cyclists (VO_{2max} $60.8 \pm 7.4 \text{ ml.kg}^{-1}$.min⁻¹; age: 36 ± 9 years; height: 1.80 ± 0.06 m; body mass: $87.2 \pm 12.0 \text{ kg}$) volunteered to participate in this study. The cyclists had a minimum of one year cycling experience and a minimum weekly training volume of (5 hr.wk⁻¹; and >150 km.wk⁻¹). The research was approved by the Departmental Ethics Committee and participants gave written informed consent prior to any data collection. The participants were categorised as trained competitive cyclists (De Pauw et al., 2013), who were fully familiar with similar cycling TT distances and laboratory testing procedures.

81

Participants were required to visit the laboratory to complete a preliminary VO_{2max} test which 82 83 used a ramp incremental exercise test on an electronically braked cycle ergometer (Lode 84 Excalibur Sport, Groningen, Netherlands). Participants were required to cycle at an initial 85 intensity of 75 W for 1 min, after which the intensity increased at a ramp rate of 30 W.min⁻¹ 86 until volitional fatigue. Breath-by-breath pulmonary gas exchange data was collected 87 continuously using a gas analysis system (K5, Cosmed, Italy) during the incremental test and 88 subsequently analysed using 5 s averaging. The VO_{2max} was calculated as the highest 15 s 89 mean value attained before the participants' volitional termination of the test. Following 90 completion of this test participants were familiarised with the Velotron[®] (RacerMate Inc, 91 Velotron, USA) in order to ensure that the specific frame geometry settings were recorded for 92 use in all subsequent TT laboratory visits.

93

94 Participants were then required to attend the laboratory on a further three separate occasions 95 at the same time of day. Prior to each of these subsequent visits, participants recorded dietary 96 intake and physical activity, which was replicated for the 24 h preceding each visit. They were also instructed to attend euhydrated, having abstained from caffeine and food ingestion a 97 98 minimum of 4 h prior to the start of exercise. Participants were also told to avoid the use of antibacterial mouthwash during the ingestion and testing periods. Using a randomised 99 crossover design, the experimental conditions of nitrate, placebo and control were 100 101 administered. This required participants to ingest either 140 ml of beetroot juice (BEET It sport 102 (B) UK) containing 12.8 mmol of NO_{3}^{-} (Larsen et al., 2010) or 140 ml of blackcurrant (P) sugar 103 free cordial (Shannon et al., 2016) once per day for three days. Experimental drinks were 104 consumed 3 h prior to the intended exercise time. In the control condition, participants 105 consumed no experimental supplement. A 48 h period between each ingestion strategy was 106 imposed to ensure adequate washout of supplementary NO_{3}^{-} (Lansley et al., 2011).

107

108 The exercise trials required participants to perform a self-selected warm-up for 5 min, which 109 was then replicated prior to all subsequent trials. Exercise performance was then assessed 110 using a 40 km TT at each of final three laboratory visits to assess the effects of the 111 experimental conditions. The TT's were performed on an electronically braked cycle 112 ergometer (Velotron, Racermate, Seattle, USA) which allowed participants to see an avatar 113 representation of their performance via software (Velotron 3D, Racermate, Seattle, USA), 114 which recorded time, distance, power, and speed. At 10 km intervals, measurements of heart 115 rate (T31, Polar, Finland), ratings of perceived exertion (Borg 1982), felt arousal (Svebak and 116 Murgatroyd, 1985), respiratory gases (VO₂) (K5, Cosmed, Italy) and fingertip capillary blood 117 samples were obtained. Blood samples were analysed to determine blood pH and bicarbonate 118 ion (HCO₃) concentrations (ABL800 basic, Radiometer, Denmark) and blood lactate (Lactate 119 Pro2 LT-1730, Arkray, Japan). During the TT all performance data were obscured from view 120 apart from distance completed and remaining. Participants received no verbal or temporal 121 feedback, or any encouragement during the TT.

122

123 Statistical Analysis

Data were assessed for normality using standard graphical procedures. All respiratory and blood biochemical responses, RPE, felt arousal, heart rate and mean performance indicators (time, speed and power output) were analysed using repeated measures ANOVA. Post-hoc pairwise comparisons were made using a Bonferroni procedure. Effect sizes were calculated using partial eta squared ($\Box p^2$), and were interpreted as: a small (< 0.01), medium (0.01 - 0.06) or a large (≥ 0.14) effect. Statistical significance was set at p < 0.05 and all procedures
were conducted using SPSS for Windows (Version 22, IBM®, Chicago, USA).

131

132 Results

133 Ingestion of the NO₃⁻ drink had no significant effect on overall TT performance time (figure 1 a and 1b), mean power, or mean speed (f = 1.27, p = 0.296, $\Box p^2 = 0.11$, f = 1.14, p = 0.339, 134 $\Box p^2 = 0.10$; and f = 1.15, p = 0.336, $\Box p^2 = 0.10$) respectively. There was also no effect on 135 either mean heart rate (f = 3.51, p = 0.054, $\Box p^2 = 0.31$) or VO₂ (f = 1.47, p = 0.253, $\Box p^2 =$ 136 0.13), but there was a significant increase in VO₂ (f = 25.68, p < 0.001, \Box p² = 0.72) during TT 137 138 performance in all conditions (figure 2). Similarly, lactate and RPE (Table 1) were unaffected 139 by the experimental conditions (f = 0.67, p = 0.522, $\Box p^2 = 0.06$; and f = 2.96, p = 0.085, $\Box p^2 =$ 140 0.30) respectively, but increases in these variables (Table 1) were observed during the course of the TT's (f = 41.98, p < 0.001, $\Box p^2$ = 0.80; and f = 46.55, p < 0.001, $\Box p^2$ = 0.87) for lactate 141 142 and RPE respectively. In the case of lactate, concentrations at 40 km where elevated 143 compared to all other distances (p < 0.001). Whereas RPE (Table 1) increased between 10-144 20 km (p = 0.001), 20-40 km (p = 0.001), between 10-40 km (p < 0.001), and in the last 10 km 145 of the TT (p = 0.014). Felt Arousal (Table 1) was unaffected by either the experimental 146 condition (f = 1.85, p = 0.184, $\Box p^2 = 0.16$) or during the TT (f = 1.66, p = 0.196, $\Box p^2 = 0.14$).

147

148 [Insert Figure 1 Near Here]

149 [Insert Figure 2 Near Here]

150

The blood acid-base response variable of pH and HCO₃ (Table 1) were also unaffected by the experimental conditions (f = 2.02, p = 0.158, $\Box p^2 = 0.17$) and (f = 1.54, p = 0.239, $\Box p^2 = 0.13$) respectively. There were however some perturbations in these variables during the course of the TT's (f = 37.60, p < 0.001, $\Box p^2 = 0.79$; f = 48.88, p < 0.001, $\Box p^2 = 0.83$), for pH and HCO₃ respectively. The pH response was unchanged between 10 and 20 km (p = 0.060), but it was significantly elevated between 10-30 km (p = 0.006) and then decreased in the final 10 km (p < 0.001). Between 10-30 km there was a significant increase in HCO₃ (p = 0.017), but this then decreased rapidly in the final 10 km (p < 0.001).

159

160 [Insert Table 1 Near Here]

161

162 Discussion

163 The aim of this study was to investigate the effects of a 12.8 mmol·day⁻¹ NO₃⁻ dose for three 164 days prior the performance of a 40 km TT in trained cyclists. The principle performance 165 findings suggest that NO₃, in the form of beetroot juice, elicited no significant improvements 166 in time to completion or mean power output. These findings support previous research which 167 have reported no change in performance in TT greater than 30 min in superiorly trained 168 cohorts (Cermak et al., 2012b; Wilkerson et al., 2012; Lane et al., 2013). Nevertheless, this 169 study is the first to utilise a chronic 3-day dosing strategy and therefore, suggesting that 170 increasing the concentration of NO₃ ingestion does not alter performance during a longer 171 duration 40 km TT in trained cyclists.

172

173 The 3-day supplementation protocol utilised in this study administered a total of 38.4 mmol of 174 NO_{3} prior to exercise, which is substantially greater than the 6 to 17.4 mmol administered 175 during previous investigations (Cermak et al., 2012b; Wilkerson et al., 2012; Lane et al., 2013). 176 These studies reported a plasma NO₂ increase by 25% (Wilkerson et al., 2012) and 96% 177 (Cermak et al., 2012b) from baseline to pre-exercise. While we did not measure plasma NO₂, 178 Kelly et al., (2014) utilised a similar 3-day supplementation approach providing 25.2 mmol 179 NO3⁻ and reported a substantial 242% and 557% mean increase from baseline, in a similarly trained population (VO_{2max}: 58.3 ± 6.3 ml·kg⁻¹·min⁻¹). Given a greater concentration of NO₂ was 180 supplemented in the current study it is fair to postulate a similar increase in plasma NO₂, which 181 is substantially larger then that previously reported during prolonged TT investigations 182 (Cermak et al., 2012b; Wilkerson et al., 2012). Consequently, the lack of change in 183 184 performance is unlikely to be caused by insufficient plasma NO₂ bioavailability.

185

186 A recent investigation by Shannon et al., (2017) highlighted the ergogenic effects of NO_3^- may be a manifestation of the exercise intensity and duration. Acute beetroot juice supplementation 187 188 significantly improved 1500 m TT performance but not 10000 m TT performance in trained 189 runners. The shorter distance requires a relatively higher intensity, thereby prompting an 190 increase in pulmonary VO_2 and HR demand, a greater perturbation to muscle metabolic milieu 191 (e.g. Hydrogen ion and phosphate accumulation) and an increased recruitment of type 2 192 muscle fibres. The underlying mechanism of NO₃ ergoencity are vast, but include improved 193 mitochondrial efficiency and reductions in muscle metabolic perturbations (Vanhatalo et al., 194 2011), particularly within type 2 fibres where blood flow and O₂ delivery appears to be 195 enhanced (Ferguson et al., 2015). However, during the current investigation, the 40 km TT 196 was performed at submaximal intensities and thus limiting type 2 fibre recruitment. The 197 differentiation in fibre recruitment may therefore, explain the lack of improvement in the current 198 study, in comparison to the ergogenic effects reported in higher intensity 4 km and 16 km 199 TT's (Lansley et al., 2011). Interestingly, Wilkerson et al., (2012) observed a significant 200 supplement effect during the final 16 km split of the 80.4 km TT, which was a higher intensity 201 than preceding splits and is characterised by the common increase in power output commonly 202 observed during the final part of TT performances (Jones et al., 2015). While this may suggest 203 NO_{3} may be beneficial during stochastic activities and the final part of a TT, this effect was 204 not observed in the current study; thereby questioning the application of NO_3^{-1} in this context. 205

In the present study there was no effect of NO_3^- on VO_2 , but in several studies reduced VO_2 has been reported (Bailey et al., 2012; Lansley et al., 2011). Typically, these studies have used shorter duration and therefore, higher intensity exercise protocols; however, Wilkerson et al., (2012) reported a significant increase in the power output: VO_2 ratio, suggesting improved efficiency in well trained cyclist with VO_{2max} values of $63 \pm 8 \text{ ml.kg}^{-1}$. This study also reported an increase in plasma NO_3^- , suggesting that the NO_3^- ingestion caused the established physiological adaptations of reduced ATP cost of muscle force production and 213 increased mitochondrial functioning. Despite this, along with improvement in the final guintile 214 of the TT, this was still not enough to elicit an overall improvement in TT performance following 215 NO₃⁻ ingestion. The present respiratory response data showed no differences between 216 conditions, which is in agreement with the work of Bourdillon et al., (2015) and Christensen et 217 al., (2013) who also reported no change in VO₂ and RER during 120 min submaximal cycling 218 or in a subsequent 400 kcal TT in elite cyclists. Christensen et al., (2013), have suggested 219 that training status, determined as a VO_{2max} of 50 mL.kg⁻¹.min⁻¹ in this case, may limit the 220 effectiveness of supplementation and therefore be a factor to explain the lack of difference 221 found in the present study.

222 Conclusion

The aim of this research was to investigate a high dose of NO_3^- supplementation over three days ingestion on trained cyclists performing a 40km TT performance. This represents a novel ingestion strategy in a highly ecologically valid TT distance. However, the ingestion of dietary NO_3^- had no significant effect on performance, failed to improve respiratory efficiency, lower blood acidity or affect psychological and physiological fatigue indicators. Therefore, the use of an ergogenic aid containing 12.8 mmol (800 mg) of NO_3^- each day for three days prior to a 40 km cycling TT, is unlikely to provide a beneficial effect on performance in trained cyclists.

230

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240	Conflict of Interest
241	The authors declare that they have no conflicts of interests.
242	
243	

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- 339

341 Figure Legends

Figure 1. Mean (± SD) overall 40 km performance time for each experimental condition (a)
and individual performance responses (b).

Figure 2. Mean (\pm SD) VO₂ responses during each experimental condition during the 40 km time trials. (*) Denotes a significant increase in VO₂ from the other distances in all conditions (p < 0.01).

Table 1. Mean (±SD) Blood metabolite and subjective rating scale responses to nitrate and placebo ingestion during 40 km cycle time trials. (*) Represents a significantly increase from all other distances (p < 0.001); (•) represents a significant change over the previous 30 km (p < 0.01); (∇) Denotes a significant decrease over the previous 10 km (p < 0.05); and (Δ) represents a significant increase over the previous 10 km (p < 0.05).

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	Distance (km)				
	10	20	30	40	
Lactate (mmol.l ⁻¹)					
Control	5.15 ± 2.04	5.04 ± 2.47	6.56 ± 3.70	9.15 ± 3.47*●	
Placebo	5.05 ± 2.14	4.95 ± 2.93	5.09 ± 3.03	8.75 ± 3.38*●	
Nitrate	4.33 ± 2.05	3.97 ± 2.00	3.99 ± 2.06	7.79 ± 2.00*●	
рН					
Control	7.37 ± 0.05	7.39 ± 0.06	7.40 ± 0.05	$7.34 \pm 0.06\nabla$	
Placebo	7.40 ± 0.04	7.41 ± 0.05	7.42 ± 0.05	$7.34 \pm 0.06\nabla$	
Nitrate	7.39 ± 0.05	7.40 ± 0.06	7.41 ± 0.06	7.34 ± 0.07∇	
HCO ₃ - (mmol.l ⁻¹)					
Control	20.81 ± 2.86	20.79 ± 3.13	21.20 ± 2.56	18.33 ± 2.43∇	
Placebo	21.43 ± 2.17	21.88 ± 2.41	22.25 ± 2.52	18.58 ± 2.46∇	
Nitrate	21.26 ± 2.37	21.43 ± 2.60	21.94 ± 2.91	17.79 ± 2.20∇	
RPE					
Control	14.0 ± 1.31	$14.9 \pm 2.0\Delta$	15.8 ± 2.6	19.4 ± 1.1∙	
Placebo	14.1 ± 1.7	$15.3 \pm 2.0\Delta$	16.0 ± 2.2	19.0 ± 1.2∙	
Nitrate	14.5 ± 2.3	15.8 ± 1.6∆	16.5 ± 1.9	19.4 ± 1.2∙	
Felt Arousal					
Control	4.18 ± 0.75	4.45 ± 1.04	4.64 ± 0.67	4.00 ± 0.77	
Placebo	4.00 ± 0.77	4.00 ± 1.10	4.00 ± 1.34	4.18 ± 1.33	
Nitrate	4.27 ± 1.01	4.45 ± 0.82	4.45 ± 0.69	4.64 ± 0.81	





