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1 **Title:** Effect of lactate supplementation and sodium bicarbonate on 40 km cycling time trial  
2 performance

3

4 **Running head:** Sodium bicarbonate and lactate time-trial

5

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38 ABSTRACT

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40 The use of nutritional supplements to improve sporting performance and increase training  
41 adaptations is commonplace amongst athletes and is an expanding market in terms of product  
42 choice and availability. The purpose of this study was to examine the effects of two ergogenic  
43 aids with extracellular blood buffering potential, namely sodium bicarbonate ( $\text{NaHCO}_3$ ) and a  
44 lactate supplement, during a 40 km cycling time trial. Seven recreationally active males (age,  
45  $22.3 \pm 3.3$  years; height,  $182.5 \pm 6.5$  cm; body mass,  $79.2 \pm 6.3$  kg) completed five 40 km  
46 cycling time trials, including a familiarization trial in a randomized blind double placebo  
47 design. Subjects ingested either (a) 300 mg per kg body mass  $\text{NaHCO}_3$  (BICARB), (b) 45 mg  
48 per kg sodium chloride (PL-BICARB) as the placebo for the  $\text{NaHCO}_3$  trial, (c) 1115 mg  
49 lactate (LACTATE), or (d) plain flour as the placebo for the lactate trial (PL-LACTATE) 60  
50 minutes before exercise. There was no significant difference in performance between the four  
51 conditions ( $p > 0.05$ ). Whilst  $\text{NaHCO}_3$  ingestion induced significant changes in all the acid-  
52 base variables (all  $p < 0.05$ ), no significant change was seen following lactate ingestion ( $p >$   
53  $0.05$ ). Subjects in the LACTATE condition did have a significantly higher heart rate ( $p <$   
54  $0.05$ ) without experiencing any greater perceived exertion ( $p > 0.05$ ) than the other three  
55 conditions. Neither  $\text{NaHCO}_3$  nor lactate supplementation appear to improve 40 km cycling  
56 time trial performance. However the potential benefits following LACTATE regarding  
57 perceived exertion require further research.

58

59 **Key Words:** buffering, alkalosis, ergogenic aid,  $\text{NaHCO}_3$ , acid-base

60

61 INTRODUCTION

62

63 The use of nutritional ergogenic supplements are commonplace within sport as both  
64 recreational and professional athletes aim to improve performance and increase training  
65 adaptations(19). Previous research into the benefits of induced metabolic alkalosis on both  
66 prolonged continuous and intermittent high-intensity exercise has proved to be equivocal(22).  
67 The majority of research has found no significant improvement in endurance performance  
68 following induced alkalosis in cycling(1, 32, 36). The exception to these however was  
69 McNaughton, Dalton and Palmer(21) who reported a 14% increase in work capacity during  
70 60 minutes of high-intensity cycling following the ingestion of sodium bicarbonate  
71 ( $\text{NaHCO}_3$ ). If the results from such fixed time duration studies could be replicated in a more  
72 practical setting such as time trial cycling involving set distance, then  $\text{NaHCO}_3$  could prove  
73 to be an inexpensive ergogenic aid. A negative side effect of  $\text{NaHCO}_3$  however is the  
74 possibility of gastrointestinal (GI) distress(6, 34) which ultimately may offset any possible  
75 positive benefits to be gained.

76

77 Decreases in intramuscular pH have previously been reported to inhibit the contractile  
78 processes by either a) restricting myofilament function through reducing  $\text{Ca}^{2+}$  sensitivity(7,  
79 11) or b) effecting the excitation-contraction process relating to the uptake and release of  $\text{Ca}^{2+}$   
80 by the sarcoplasmic reticulum(16, 33). By ingesting  $\text{NaHCO}_3$  prior to exercise, extracellular  
81 bicarbonate ( $\text{HCO}_3^-$ ) reserves are supplemented, resulting in an increased plasma pH and an  
82 induced state of metabolic alkalosis(30). The extracellular to intracellular pH gradient  
83 therefore increases as  $\text{HCO}_3^-$  is impermeable to cellular membranes(22) resulting in a greater  
84 efflux of  $\text{H}^+$  and lactate from active muscles(26). This occurs via either simple diffusion or by

85 the lactate/H<sup>+</sup> co-transporters(17) and has been demonstrated by the higher lactate  
86 concentrations post-exercise following NaHCO<sub>3</sub> ingestion(1, 28). Increases in plasma HCO<sub>3</sub><sup>-</sup>  
87 have also been reported following the ingestion of lactate with no reported GI distress(24,  
88 39), showing potential for it to be utilized as an alternative exogenous buffer to NaHCO<sub>3</sub>.

89

90 Within exercise metabolism the role of lactate and in particular its production, has been a  
91 source of much dispute(4, 11, 15). Debate remains whether the presence of lactate acts as a  
92 limiting factor during exercise by inducing acidosis or actually attenuates the onset of fatigue  
93 by consuming the excess H<sup>+</sup> responsible for acidosis(4). Many of the studies associating  
94 lactate with the development of fatigue tend to be based upon correlational data(4) therefore a  
95 cause and effect relationship cannot be ascertained. Furthermore, lactate has the potential to  
96 serve as a source of glucose generated from within the body as a substrate for  
97 gluconeogenesis(3). Based upon the lactate shuttle theory(2), exogenous lactate  
98 supplementation therefore has the potential to increase plasma glucose supplied via  
99 gluconeogenesis thus sparing glycogen stores<sup>20</sup>. However to date, research(3, 27, 38) has  
100 failed to support this theory.

101

102 The purpose of this study therefore was to determine whether either NaHCO<sub>3</sub> or lactate  
103 supplementation had any ergogenic potential to improve 40 km time trial performance.  
104 Additionally it was designed to establish whether any improvement in performance was  
105 associated with changes in acid-base status and buffering capacity. It was hypothesized that  
106 the use of either NaHCO<sub>3</sub> or a lactate supplement would improve the performance of a 40 km  
107 cycling time trial. Furthermore, it was hypothesized that lactate supplementation would

108 increase both plasma lactate levels and buffering capacity whilst causing less GI distress than  
109 is associated with NaHCO<sub>3</sub> consumption.

110

## 111 METHODS

112

### 113 EXPERIMENTAL APPROACH TO THE PROBLEM

114

115 Using a randomized, double placebo-controlled design; subjects were required to complete a  
116 total of five trials, one familiarization trial and four experimental trials. Subjects were  
117 instructed to arrive for testing in a rested state having refrained from strenuous exercise and  
118 alcohol in the 24 hour prior to testing and had no history of either NaHCO<sub>3</sub> or lactate  
119 supplementation. Subjects were asked to ingest a minimum of 500 ml of water before  
120 arriving at the laboratory to ensure they arrived in a well hydrated state, avoiding caffeine in  
121 the 12 hours prior to each trial. They were also asked to consume the same standardized  
122 breakfast a minimum of 1 hour prior to arriving for each trial. Subjects performed each of the  
123 five trials at the same time of day to control for circadian variation in performance(9).  
124 Additionally, each trial was separated by a period of 6 to 9 days in order to ensure an  
125 adequate recovery period was attained whilst limiting the opportunity for any improvement  
126 being the result of training.

127

128 The four experimental conditions were (a) 300 mg per kg body mass NaHCO<sub>3</sub> (BICARB),  
129 (b) 45 mg per kg sodium chloride (PL-BICARB) as the placebo for the NaHCO<sub>3</sub> trial, (c)

130 1115 mg lactate from a combination of calcium lactate pentahydrate and magnesium lactate  
131 dihydrate, equivalent to a mean of 14.1 mg.kg<sup>-1</sup> body mass per participant based on mean  
132 body weight (Sport Specifics, Inc., Chagrin Falls, OH, USA) (LACTATE), and (d) plain  
133 flour as the placebo for the lactate trial (PL-LACTATE). All supplements were ingested  
134 within gelatine capsules with 500 ml low calorie cordial over a 10 minute period, 60 minutes  
135 prior to exercise. Due to the disparity between the NaHCO<sub>3</sub> and lactate trials in terms of  
136 capsules required, a double placebo design was chosen to improve validity. The use of 300  
137 mg per kg body mass NaHCO<sub>3</sub> has been established as the optimal dose for enhanced  
138 buffering capacity(22) and has previously been used in a number of studies into NaHCO<sub>3</sub>  
139 supplementation(5, 25, 33). Furthermore, peak HCO<sub>3</sub><sup>-</sup> levels are typically achieved 60  
140 minutes following ingestion(34). The lactate supplement dosage used was as per the  
141 manufacturer's instructions. It was felt that this dosage of the supplement should be chosen as  
142 consumers who purchase this supplement are unlikely to exceed the recommended dosage.

143

144 In terms of performance, the dependent variables of interest were overall performance time,  
145 split performance time, heart rate and rate of perceived exertion (RPE). For changes in acid-  
146 base status the dependent variables were pH, base excess (BE), HCO<sub>3</sub>, lactate and H<sup>+</sup>.  
147 Changes in overall performance time represent an accurate comparison between trials whilst  
148 ultimately being the key variable of interest for competitive cyclists. The use of split times  
149 allowed for changes during individual stages to also be identified. The use of pH, BE, HCO<sub>3</sub>  
150 and H<sup>+</sup> in research looking at changes in buffering capacity following supplementation is  
151 well established (5, 34, 35). Furthermore, as one of the supplements contained exogenous  
152 lactate, it was important to establish whether any changes in plasma lactate occurred  
153 following its ingestion.



154

155 SUBJECTS

156

157 Seven recreationally active non-smoking male subjects (mean  $\pm$  SD: age,  $22.3 \pm 3.3$  years;  
158 height,  $182.5 \pm 6.5$  cm; body mass,  $79.2 \pm 6.3$  kg) with no previous history of supplementing  
159 their diets with ergogenic agents volunteered to participate in this study. The subjects  
160 consisted of four cyclists, two footballers and a long distance runner, all whom were in a  
161 period of regular training at the time of testing. They were all completing a minimum of four  
162 hours ( $6.3 \pm 3.3$  hours) training per week and all were free from any known cardiac or  
163 metabolic diseases. All subjects provided written informed consent, and the study was  
164 approved by the Departmental Human Ethics Committee and following the principles  
165 outlined in the Declaration of Helsinki.

166

167 PROCEDURES

168

169 On arrival at the laboratory, the subject had a capillary blood sample taken to establish basal  
170 acid-base measurements (pH, BE,  $\text{HCO}_3^-$ , lactate and  $\text{H}^+$ ) before ingesting the relevant  
171 supplement. During the 60 minute post-ingestion period, further capillary blood samples were  
172 taken at 10, 20, 30, 45 and 60 minutes post ingestion to evaluate any induced changes in the  
173 acid-base variables. All blood samples were collected using 100  $\mu\text{l}$  balanced heparin blood  
174 capillary tubes (Radiometer, West Sussex, UK) and immediately analyzed (Radiometer,  
175 ABL800, Copenhagen, Denmark).

176

177 During the ingestion period, subjects were asked to rate any GI discomfort experienced every  
178 15 minutes using a visual analogue scale (VAS) until the exercise commenced. The potential  
179 symptoms listed were: nausea, flatulence, stomach cramping, stomach bloating, stomach-  
180 ache, belching, vomiting, bowel urgency and diarrhoea. The VAS scale consisted of nine  
181 separate 100 mm scales, anchored at each end with either 'no symptom' or 'severe symptom'  
182 and subjects indicated with a vertical mark the severity of each symptom during the ingestion  
183 period(5, 34). None of the subjects reported any instances of GI disturbance during the 60  
184 minute pre-exercise period as a result of ingesting BICARB, LACTATE or either placebo.

185

186 Following the 60 minute post ingestion capillary blood sample, the subject completed a ten  
187 minute warm up at an intensity of 75 watts prior to beginning the 40 km time trial. The time  
188 trial was conducted using a Wattbike cycle ergometer (Wattbike Ltd, Nottingham, UK) with  
189 heart rate (Polar FS1 HRM, Polar Electro, OY, Finland) and rate of perceived exertion (RPE)  
190 recorded at five minute intervals. RPE was monitored using a modified version of Foster et  
191 al.(12) perceived exertion scale. Subjects were permitted to drink water ad libitum. During  
192 each trial, subjects were blinded to all performance data except the distance countdown. The  
193 purpose for this was to minimize any learning effect to be gained from previous trials.  
194 Additional capillary blood samples were collected at 20 km, 40 km and at 15 minutes post-  
195 exercise.

196

197 STATISTICAL ANALYSES

198

199 Statistical analyses were completed using IBM PASW statistics 18 (SPSS inc., Chicago, IL).  
200 Central tendency and dispersion of all data are displayed as mean  $\pm$  standard deviation (S.D).  
201 Performance time data was compared using a one way analysis of variance (ANOVA) with  
202 repeated measures whilst changes in acid-base status, heart rate and RPE were investigated  
203 using two way ANOVA with repeated measures. Sidak-adjusted  $p$  values were used for  
204 subsequent pairwise comparisons to establish the significant paired differences when  
205 significant  $F$  ratios were found by the respective ANOVA. Statistical significance was  
206 accepted as  $p < 0.05$  with effect size reported according to partial eta squared.

207

## 208 RESULTS

209

### 210 PERFORMANCE DATA

211

212 The mean times for 10 km, 20 km, 30 km and 40 km along with the individual split times for  
213 each 10 km interval are displayed in Table 1. Whilst overall performance time for LACTATE  
214 was between 1-3% faster than the other three conditions, the difference was not significant ( $p$   
215  $>0.05$ ,  $\eta_p^2 = 0.20$ ). Furthermore, the individual split times for each 10 km stage of the time  
216 trial were not significantly different between the four conditions ( $p >0.05$ ). Individual  
217 responses to the supplements were varied with 3 subjects performing their fastest trial in the  
218 LACTATE condition, whilst 2 performed fastest in PL-LACTATE, and 1 subject completing  
219 the time trial fastest in each of the BICARB and PL-BICARB conditions (Figure 1).

220

221

INSERT TABLE 1

222

223

INSERT FIGURE 1

224

225 Average heart rate during the LACTATE condition ( $169 \pm 9$  bpm) was significantly higher  
226 than in the other three conditions (BICARB:  $160 \pm 16$  bpm; PL-BICARB:  $158 \pm 13$  bpm; PL-  
227 LACTATE:  $160 \pm 14$  bpm respectively;  $p < 0.05$ ,  $\eta_p^2 = 0.49$ ) throughout the duration of the  
228 time trials. No significant difference was seen however between the other three conditions.  
229 Both heart rate and RPE (both  $p < 0.05$ ,  $\eta_p^2 = 0.71$  and  $\eta_p^2 = 0.85$  respectively) was seen to  
230 increase progressively with each 10 km stage of the time trial (Table 2). No significant main  
231 effect for condition or interaction effect between condition and stage (both  $p > 0.05$ ,  $\eta_p^2 =$   
232  $0.18$  and  $\eta_p^2 = 0.11$  respectively) was seen for RPE between the four conditions.

233

234

INSERT TABLE 2

235

236 ACID-BASE BALANCE

237

238 Changes in pH, BE,  $\text{HCO}_3^-$ , lactate and  $\text{H}^+$  for the four conditions across the study are  
239 displayed in Table 3. There was no significant difference between pre-ingestion levels for any  
240 of the blood variables between the four conditions. During the BICARB condition, blood pH  
241 was significantly higher at 45 and 60 minutes post ingestion than seen at pre-ingestion ( $p <$   
242  $0.05$ ,  $\eta_p^2 = 0.84$ ), whilst  $\text{H}^+$  levels were significantly lower at the same time points ( $p < 0.05$ ,

243  $\eta_p^2 = 0.67$ ). By 60 minutes post-BICARB ingestion, BE had increased by in excess of 5  
244 mEq/L ( $p < 0.05$ ,  $\eta_p^2 = 0.85$ ) and plasma  $\text{HCO}_3^-$  by approximately 4.5 mmol/L compared to  
245 the pre-ingestion levels ( $p < 0.05$ ,  $\eta_p^2 = 0.84$ ) and significantly increased compared to  
246 LACTATE, PL-BICARB & PL-LACTATE (all  $p < 0.05$ ) at the same time point. No  
247 significant differences were seen within the other three experimental conditions during the  
248 pre-ingestion to 60 minute post-ingestion period. Furthermore, no significant difference were  
249 seen for lactate concentration between pre-ingestion and 60 minutes post-ingestion ( $p > 0.05$ ,  
250  $\eta_p^2 = 0.11$ ) for any of the four conditions.

251

252

INSERT TABLE 3

253

254 After 20 km and 40 km, pH, BE and  $\text{HCO}_3^-$  remained elevated and  $\text{H}^+$  was lower for the  
255 BICARB condition compared to the other three conditions (Table 3). All the differences  
256 between the BICARB condition and the other three conditions were significant except for pH  
257 and  $\text{H}^+$  at 40 km compared to PL-LACTATE (both  $p > 0.05$ ) and BE at 40 km compared to  
258 the LACTATE condition ( $p > 0.05$ ). Whilst at the end of the time trial, plasma lactate was  
259 between 2-3 mmol/L higher for the BICARB condition, the difference was only significant  
260 compared to PL-LACTATE ( $p < 0.05$ ,  $\eta_p^2 = 0.48$ ). No significant difference was seen for  
261 lactate between the LACTATE, PL-LACTATE and PL-BICARB.

262

263 DISCUSSION

264

265 Whilst mean performance time following the ingestion of the lactate supplement was over 30  
266 seconds faster than the next nearest condition (Table 1) the performance effect was not  
267 significant. This mean difference was influenced by subject 2 whose individual time during  
268 the supplement trial was around 3 minutes faster than the other three conditions (Figure 1).  
269 Additionally ingestion  $\text{NaHCO}_3$  did not provide any significant ergogenic effect on 40 km  
270 time trial performance. Using the same lactate supplement, Peveler and Palmer(27) also  
271 found no significant effect on performance of 20 km time trial cycling, heart rate or mean  
272 power with the lactate condition actually marginally slower than placebo by ~17.4 seconds on  
273 average. They did however fail to show the individual performance times for each condition  
274 making it difficult to establish if there was an individual specific response from any of their  
275 subjects. Additionally, other forms of lactate supplementation focusing on lactate as a  
276 gluconeogenic precursor for endurance exercise have also been previously used  
277 unsuccessfully. Both Bryner et al.(3) and Swensen et al.(38) combined lactate and  
278 carbohydrate to examine its effect on time to exhaustion (TTE). Bryner et al.(3) found no  
279 significant effect on either TTE or peak power using a protocol that involved cycling at 10  
280 beats below target heart rate and ended with a Wingate power test during the last 30 seconds  
281 of the trial. Swensen et al.(38) also found no effect on TTE between when cycling at 70%  
282  $\text{VO}_{2\text{max}}$  until exhaustion.

283

284 Whilst the current study also supports the majority of research in finding that  $\text{NaHCO}_3$  did  
285 not improve prolonged exercise performance(1, 28, 36) one exception remains(21).  
286 McNaughton et al.(21) reported an increase in both overall work (in kilojoules) and average  
287 power following the ingestion of  $\text{NaHCO}_3$  over a 60 minute period of maximal cycling. In  
288 this study the 40-km time trial was chosen as it represented a similar duration to that seen in

289 McNaughton et al.(21) however the use of a set distance as opposed to set time duration  
290 provided a greater reflection of competitive cycling.

291

292 The increase in buffering capacity achieved by ingesting  $\text{NaHCO}_3$  prior to exercise has  
293 previously been well documented(6, 22, 35) although the benefits are typically associated  
294 with events lasting between 30 seconds to 3 minutes(22, 30, 34). Lactate levels during the  
295 current study were higher following  $\text{NaHCO}_3$  ingestion than that of the other three conditions  
296 at both 20 km and 40 km although only to a significant level at 40 km over PL-LACTATE  
297 (Table 3). It has been suggested that by increasing extracellular levels of  $\text{HCO}_3^-$ , the efflux of  
298 lactate and  $\text{H}^+$  from within the muscle is facilitated(13, 21) with similar results having  
299 previously been found by Price et al.(28). In theory this could have improved performance by  
300 maintaining pH closer to the homeostatic levels(21). McNaughton et al.(21) attributed their  
301 significant increase in work during their 60 minute cycling study to the maintenance of pH  
302 nearer to resting levels allowing greater contractile performance. Interestingly though,  
303 McNaughton et al.(21) also reported plasma lactate levels lower than those of their control  
304 (no supplement) and placebo (sodium chloride) trials. Whilst conflicting with the expected  
305 higher lactate levels seen in the current study, a difference in protocol may account for the  
306 disparity between studies.

307

308 Despite the improved acid-base status prior to and during exercise following  $\text{NaHCO}_3$   
309 ingestion in the current study, the lack of improvement in performance would appear to  
310 indicate an alternative factor separate from acidosis was the predominant cause of fatigue.  
311 Although using an alternative buffer in the form of sodium citrate, Schabert et al.(32)  
312 supported this as during the trial with the highest pH, lactate concentrations were not the

313 highest showing other factors contributed to the fatigue. Whilst allowing the subjects to  
314 consume the same standardized breakfast before each trial was intended to attenuate the  
315 effect of glycogen depletion on fatigue its effects cannot be ruled out. Furthermore the  
316 accumulation of inorganic phosphate rather than  $H^+$  has also been associated with restricting  
317 the contractile processes(40) however as these were not measured in the current study, its role  
318 cannot be determined.

319

320 Lactate supplementation has also been suggested as an alternative acid-base buffer to  
321  $NaHCO_3$ (24) however the results from this study fail to support this. Previous research has  
322 reported increases in plasma  $HCO_3^-$  following lactate ingestion(10, 24, 39) however a lack of  
323 reported pre-ingestion  $HCO_3^-$  levels mean that the level of increase is difficult to  
324 quantify(10, 39). Morris et al.(24) reported increases in plasma  $HCO_3^-$  levels of  
325 approximately 3 mmol/L between pre-ingestion levels and 80 minutes post-ingestion. In the  
326 present study, four of the seven subjects experienced an increase in  $HCO_3^-$  following  
327 ingestion of the lactate supplement although the largest increase seen was just 1 mmol/L  
328 between pre-ingestion and 60 minutes post ingestion compared to an average increase of 4.6  
329 mmol/L for the  $NaHCO_3$  condition. However, the concentration of lactate supplement in this  
330 study was considerably less than the 120 mg/kg body mass of lactate used by Morris et  
331 al.(24) or the 320 mg/kg body mass of lactate used by Van Montfoort et al.(39). The reduced  
332 dosage in the current study was used as it followed the manufacturers' guidelines and is  
333 similar to that previously used by Peveler and Palmer(27).

334

335 In the current study, the expected increase in plasma lactate failed to be observed following  
336 the ingestion of the lactate supplement. The lactate shuttle theory by which exogenous lactate



337 supplementation is thought to increase gluconeogenesis and improve performance however is  
338 highly disputed(27). An increase in plasma lactate is thought to promote increased plasma  
339 lactate oxidation whilst inhibiting intramuscular lactate production(14). Miller et al.(23)  
340 regulated lactate plasma levels to 4 mmol/L during exercise of moderate intensity via  
341 intravenous infusion. As a result of this, the contribution of glycogenolysis in supplying  
342 plasma glucose decreased as increased lactate oxidation compensated potentially sparing  
343 glycogen stores. However, in the current study an increase in plasma lactate levels was not  
344 observed following ingestion. Neither Morris et al.(24) nor Van Montfoort et al.(39) reported  
345 significant increases in plasma lactate despite using considerably larger quantities of lactate.  
346 Whilst a change in the rate of lactate oxidation potentially may account for a rise in plasma  
347 lactate not being shown, other explanations may exist. It is possible that the oral consumption  
348 of lactate was either too small to elicit a change or it failed to increase the lactate availability  
349 due to degradation by stomach acid and/or lack of absorption, unlike the direct intravenous  
350 method used by Miller et al.(23).

351 .

352 Given athletes are likely to follow manufacturers recommendations when using  
353 supplementations due to safety and overall cost issues(27), both the small change in acid-base  
354 status and the absence of an increase in plasma lactate following lactate supplementation,  
355 suggests the dosage used does not represent a viable alternative to  $\text{NaHCO}_3$  for increasing  
356 buffering capacity. The use of a chronic dosing of lactate over a number of days may be an  
357 option in the future as it has been previously shown to increase acid-base status when using  
358  $\text{NaHCO}_3$ (8, 20). However, given the negligible increases in acid-base status in this current  
359 study, each individual dose would possibly need to be increased from current study for any  
360 effect to be seen. Future investigations into alternative dosing strategies are therefore  
361 warranted

362

363 An increase in RPE over time was observed but this was not different between conditions  
364 (Table 2). However average heart rate was higher during the LACTATE condition compared  
365 to the other three conditions. Although performance differences were not significant overall  
366 in this group, the increased heart rate during the LACTATE condition may therefore have  
367 contributed for the faster performance time, although heart rate measurements were only  
368 recorded every 5 minutes meaning heart rate for each 10 km stage is based on a total of two  
369 or three measurements, which may have affected the results gained. Whilst the increased  
370 heart rate in the LACTATE condition occurred without altering perceived exertion, the  
371 subjective nature of RPE measure makes it difficult to conclude if the difference was related  
372 to the supplement. In a similar study, Peveler and Palmer(27) reported reduced RPE  
373 following the ingestion of a lactate supplement although this may have been accounted for by  
374 the slower performance time compared to the placebo condition. The effect of induced  
375 alkalosis upon the RPE following  $\text{NaHCO}_3$  and lactate ingestion has been equivocal to date  
376 with both positive(18, 31, 37) and negative(13) effects reported. This variety in results  
377 however can probably be accounted for by the variety of different exercise protocols,  
378 ingestion strategies and subject training statuses used throughout the previous literature(10,  
379 13, 18, 31, 37).

380

## 381 PRACTICAL APPLICATIONS

382

383 The pursuit of legal ergogenic aids continues at both a recreational and professional level(21).  
384 Whilst the majority of  $\text{NaHCO}_3$  supplementation research has focused on either single or

385 multiple bouts of short duration maximal intensity exercise(22, 30, 34), the research  
386 conducted into prolonged continuous and intermittent exercise has proved equivocal(1, 21,  
387 28). The current study has demonstrated there is little ergogenic benefit to be gained by  
388 inducing metabolic alkalosis via NaHCO<sub>3</sub> supplementation prior to prolonged cycling.  
389 Although not significant the LACTATE condition was fastest for three of the seven subjects  
390 and was on average approximately 30 seconds faster than the nearest condition. Whilst this  
391 figure was influenced by an individual performance of around 3 minutes faster during the  
392 lactate than the other three conditions, it raises the possibility that the ergogenic effect is  
393 individual specific. Considering the tight winning margins typically associated with time trial  
394 competition, any legal supplement that could provide such performance gains obviously  
395 would prove beneficial.

396

397 Using the dosages seen in the current study, lactate supplementation did not offer a viable  
398 alternative to NaHCO<sub>3</sub> in terms of improving blood buffering capacity. However, given  
399 NaHCO<sub>3</sub> ingestion is associated with GI distress(5, 6, 34) which may reduce any ergogenic  
400 benefits that may be achieved(34), research into alternative buffering agents is warranted. In  
401 this study no GI distress for either supplement was reported, suggesting that lactate  
402 supplementation is not associated with GI distress at this concentration and that the response  
403 to NaHCO<sub>3</sub> is individual specific, as recently alluded to by Price and Simons(29). Future  
404 work on lactate supplementation should therefore focus of dosing strategies in order to  
405 maximize the potential for an ergogenic effect to be seen on performance. Given any  
406 ergogenic effect of lactate supplementation appears to be individual specific, experimentation  
407 of the supplement prior to prolonged use is essential to assess the cost-benefit analysis to the  
408 individual.

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540 Strength and Conditioning Association.

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## FIGURES

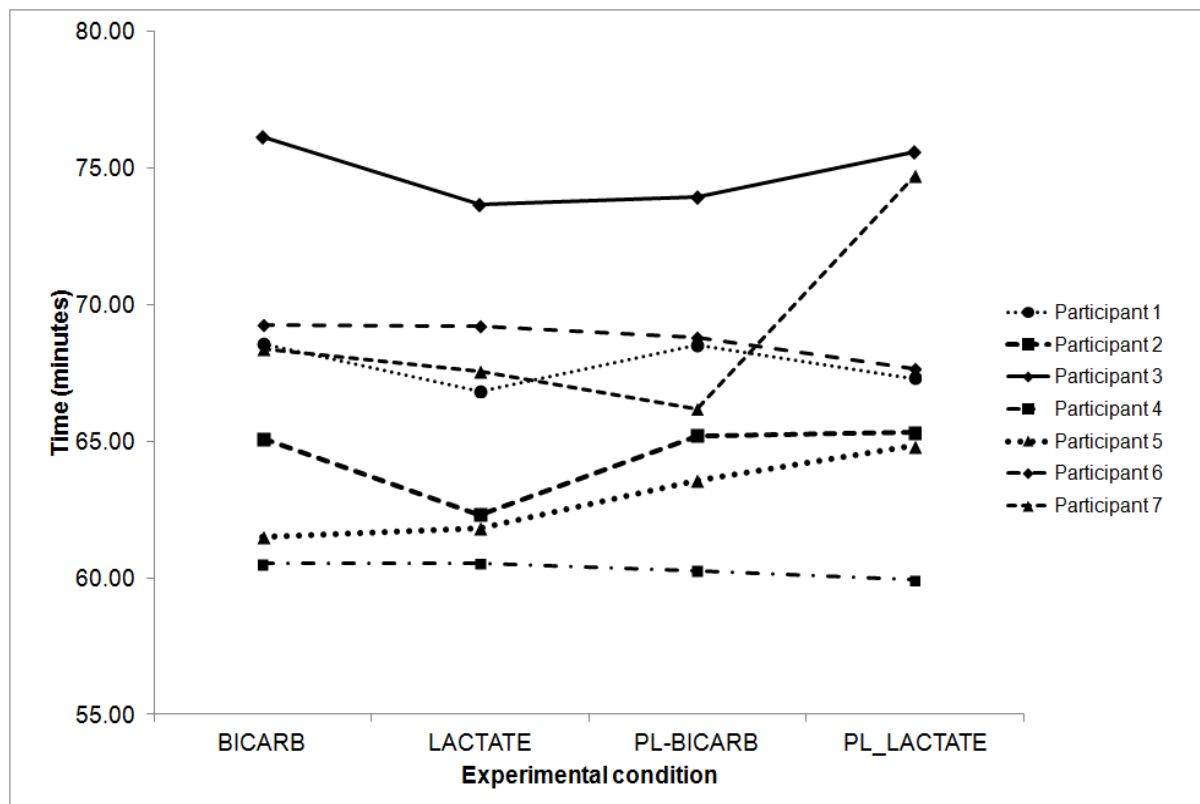


Figure 1. Individual performance times for each of the four experimental conditions

BICARB = Sodium Bicarbonate; LACTATE = Lactate supplement; PL-BICARB = Sodium Chloride; PL-LACTATE = Flour



## TABLES

Table 1. Mean 40 km Cycling time trial performance times (minutes) including split times for the four experimental conditions (mean  $\pm$  S.D) (n=7)

	10 km	20 km	30 km	40 km
BICARB	17.07 $\pm$ 1.45	34.06 $\pm$ 3.15 (17.00 $\pm$ 1.32)	50.56 $\pm$ 4.29 (16.50 $\pm$ 1.15)	67.08 $\pm$ 5.04 (16.11 $\pm$ 1.00)
LACTATE	16.19 $\pm$ 1.14	32.55 $\pm$ 2.26 (16.36 $\pm$ 1.15)	49.47 $\pm$ 3.40 (16.52 $\pm$ 1.16)	66.02 $\pm$ 4.44 (16.15 $\pm$ 1.06)
PL-BICARB	16.48 $\pm$ 1.21	33.31 $\pm$ 2.24 (16.55 $\pm$ 1.12)	50.41 $\pm$ 3.23 (16.59 $\pm$ 1.09)	66.41 $\pm$ 4.04 (15.59 $\pm$ 0.50)
PL-LACTATE	16.57 $\pm$ 1.38	33.55 $\pm$ 2.59 (16.58 $\pm$ 1.27)	51.05 $\pm$ 4.02 (17.10 $\pm$ 1.08)	67.54 $\pm$ 5.34 (16.49 $\pm$ 1.53)

Split times displayed in brackets

BICARB = Sodium Bicarbonate; LACTATE = Lactate supplement; PL-BICARB = Sodium Chloride;

PL-LACTATE = Flour

Table 2. Mean heart rate and rate of perceived exertion for the four conditions during each 10 km stage of the time trial (mean  $\pm$  S.D) (n=7)

	Stage (km)			
	0-10 km	10-20 km	20-30 km	30-40 km
<b>Heart rate* (BPM)</b>				
BICARB	149.3 $\pm$ 17.6	157.0 $\pm$ 15.0	161.4 $\pm$ 13.8	170.3 $\pm$ 10.6
LACTATE	162.9 $\pm$ 9.5	169.3 $\pm$ 6.1	169.7 $\pm$ 6.3	176.1 $\pm$ 4.3
PL-BICARB	150.9 $\pm$ 11.1	155.6 $\pm$ 10.5	158.7 $\pm$ 12.9	168.0 $\pm$ 12.7
PL-LACTATE	152.6 $\pm$ 17.6	158.0 $\pm$ 13.8	162.7 $\pm$ 10.1	168.7 $\pm$ 8.1
<b>RPE*</b>				
BICARB	4.4 $\pm$ 1.1	5.0 $\pm$ 1.0	5.8 $\pm$ 0.7	6.7 $\pm$ 0.8
LACTATE	4.7 $\pm$ 0.5	5.5 $\pm$ 0.9	6.1 $\pm$ 1.2	7.6 $\pm$ 1.2
PL-BICARB	4.9 $\pm$ 1.3	5.6 $\pm$ 1.2	6.1 $\pm$ 0.5	7.3 $\pm$ 1.1
PL-LACTATE	3.9 $\pm$ 1.3	4.7 $\pm$ 0.9	5.5 $\pm$ 0.9	7.1 $\pm$ 0.9

\*Significant main effect for stage,  $p < 0.05$ ; BICARB = Sodium Bicarbonate; LACTATE = Lactate supplement; PL-BICARB = Sodium Chloride; PL-LACTATE = Flour

Table 3 Mean acid-base variables at different time points pre- and post- ingestion for the four conditions (mean  $\pm$  S.D) (n=7)

		Pre- ingestion	10 min post- ingestion	20 min post- ingestion	30 min post- ingestion	45 min post- ingestion	60 min post- ingestion	20 km	40 km	15 min post- exercise
pH	BICARB	7.402	7.416	7.418	7.433	7.443* <sup>+</sup>	7.450** <sup>+++</sup>	7.446* <sup>o</sup>	7.385 <sup>o</sup>	7.460* <sup>*</sup>
	LACTATE	7.399	7.403	7.401	7.402	7.398	7.399	7.366	7.332	7.388
	PL-BICARB	7.397	7.396	7.405	7.404	7.411	7.392	7.374	7.312	7.375
	PL-LACTATE	7.406	7.400	7.398	7.398	7.397	7.393	7.381	7.333	7.377
BE	BICARB	0.2	1.0	2.6	3.6	4.6* <sup>+</sup>	5.7** <sup>+++</sup>	1.3* <sup>*</sup>	-3.9 <sup>#</sup>	2.3** <sup>**</sup>
(mEq/L)	LACTATE	0.6	0.7	1.0	0.9	0.7	0.7	-6.0	-9.0	-3.5

	PL-BICARB	1.0	0.5	0.8	0.5	0.8	0.6	-4.6	-9.2	-4.4
	PL-LACTATE	0.6	0.3	0.2	0.6	0.7	0.7	-4.3	-7.7	-3.2
HCO <sub>3</sub>	BICARB	24.6	25.3	26.4	27.4	28.3* <sup>†</sup>	29.2*** <sup>††</sup>	26.0*** <sup>††</sup>	21.6*	26.7**
(mmol/L)	LACTATE	24.8	24.9	25.1	25.0	24.8	24.8	20.0	17.8	21.8
	PL-BICARB	25.0	24.7	25.0	24.8	25.1	24.7	20.9	17.4	21.1
	PL-LACTATE	24.8	24.6	24.5	24.8	24.7	24.7	21.3	18.6	22.0
H <sup>+</sup>	BICARB	39.7	38.4	38.3	36.9	36.0*	35.5**	35.9**	41.4 <sup>#</sup>	34.6*
(mmol/L)	LACTATE	39.9	39.6	39.7	39.7	40.0	40.0	43.1	46.6	41.0

	PL-BICARB	40.1	40.2	39.4	39.5	38.9	40.5	42.4	48.8	42.2
	PL-LACTATE	39.3	39.8	40.0	40.0	40.1	40.5	41.8	46.5	41.6
La	BICARB	2.2	1.9	2.0	1.9	1.8	1.6	8.6	12.8 <sup>^</sup>	7.5
(mmol/L)	LACTATE	1.9	2.1	1.7	1.8	1.7	1.8	7.8	10.7	5.3
	PL-BICARB	1.8	1.8	1.7	1.8	1.6	1.6	6.6	10.6	5.4
	PL-LACTATE	1.6	1.7	1.8	1.6	1.7	1.7	6.9	9.7	5.0

BICARB = Sodium Bicarbonate; LACTATE = Lactate supplement; PL-BICARB = Sodium Chloride; PL-LACTATE = Flour; BE = Base excess; HCO<sub>3</sub> = Bicarbonate; H<sup>+</sup> = Hydrogen ion; La = Lactate

\*Significant difference between BICARB and LACTATE/PL-BICARB/PL-LACTATE,  $p < 0.05$ , \*\*Significant difference between BICARB and LACTATE/PL-BICARB/PL-LACTATE,  $p < 0.01$ , ° Significant difference between BICARB and LACTATE/PL-BICARB,  $p < 0.05$ , # Significant difference between BICARB and PL-BICARB/PL-LACTATE,  $p < 0.05$ , ^ Significant difference between BICARB and PL-LACTATE,  $p < 0.05$ , + Significantly different to pre-ingestion levels,  $p < 0.05$ , ++ Significantly different to pre-ingestion levels,  $p < 0.01$ .

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