Expectancy of ergogenicity from sodium bicarbonate ingestion increases high-intensity cycling capacity.

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Title: Expectancy of ergogenicity from sodium bicarbonate ingestion increases high-intensity cycling capacity.

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Abstract

This study examined whether expectancy of ergogenicity of a commonly used nutritional supplement (sodium bicarbonate; NaHCO₃) influenced subsequent high-intensity cycling capacity. Eight recreationally active males (age: 21±1 years, body mass: 75±8 kg, height: 178±4 cm, W_{PEAK}: 204±23 W) performed a graded incremental test to assess peak power output (W_{PEAK}), one familiarisation trial and two experimental trials. Experimental trials consisted of cycling at 100% W_{PEAK} to volitional exhaustion (T_{LIM}) 60 min after ingesting either a placebo (PLA; 0.1 g.kg⁻¹ sodium chloride (NaCl), 4 ml.kg⁻¹ tap water and 1 ml.kg⁻¹ squash) or a sham placebo (SHAM; 0.1 g.kg⁻¹ NaCl, 4 ml.kg⁻¹ carbonated water and 1 ml.kg⁻¹ squash). SHAM aimed to replicate the previously reported symptoms of gut fullness (GF) and abdominal discomfort (AD) associated with NaHCO₃ ingestion. Treatments were administered double blind and accompanied by written scripts designed to remain neutral (PLA) or induce expectancy of ergogenicity (SHAM). After SHAM mean T_{LIM} increased by 9.5% compared to PLA (461±148 s vs. 421±150 s; P=0.048, d=0.3). Ratings of GF and AD were mild but ~ 1 unit higher post ingestion for SHAM. After 3 min T_{LIM} RPEₜ was 1.4±1.3 units lower for SHAM compared to PLA (P=0.020, d=0.6). There were no differences between treatments for blood lactate, blood glucose or heart rate. In summary, ergogenicity after NaHCO₃ ingestion might be influenced by expectancy which mediates perception of effort during subsequent exercise. The observed ergogenicity with SHAM did not affect our measures of cardiorespiratory physiology or metabolic flux.

Keywords: Psychobiology, endurance, fatigue, RPE, perceived exertion
Introduction

A significant amount of research has examined the efficacy of sodium bicarbonate (NaHCO₃) as an ergogenic aid (Cameron et al. 2010; Krstrup et al. 2015; Lavender and Bird 1989; Price and Simons 2010; Tan, et al. 2010). Cameron et al. (2010) reported that ingestion of 0.3 g.kg⁻¹ NaHCO₃ 60 min prior to exercise had no effect on multiple indices of repeated sprint performance in elite rugby players. Cameron et al. (2010) suggested the most likely reason for the lack of performance enhancement was due to the higher frequency and greater severity of gastrointestinal (GI) symptoms after NaHCO₃ ingestion. However, the lack of performance improvement might also have been influenced by the trained nature of participants with Peart et al. (2012) reporting that untrained individuals are significantly more likely to experience ergogenic benefit with NaHCO₃. Indeed it is now well established that individuals who undertake high-intensity training have elevated levels of muscle carnosine compared with endurance trained and untrained individuals (Parkhouse and McKenzie 1984; Parkhouse et al. 1985). As carnosine is thought to play an important role in the homeostasis of muscle cells during high-intensity exercise (as an intracellular buffer) greater basal levels might ‘offset’ any potential ergogenic contribution from augmented extracellular NaHCO₃ (Aschenbach et al. 2000; Derave et al. 2010).

In support of this, Higgins et al. (2013) reported that recreationally active healthy males who ingested 0.3 g.kg⁻¹ NaHCO₃ 60 min prior to cycling at 100% WPEAK to volitional exhaustion (T_LIM) had on average 17% greater
exercise capacity versus a taste matched placebo. Furthermore, the individual who had the highest increase in $T_{\text{LIM}} (+38\%)$ also reported moderate-high GI symptoms 30 min prior and immediately prior to exercise. The authors suggested that GI symptoms might not necessarily negatively affect exercise performance, something that is supported by previous research (Price and Simons 2010).

According to Clark et al. (2000; pg 1642), the placebo effect is “...a favourable outcome arising purely from belief that one has received a beneficial treatment...” Expectancy is closely associated and in some instances assumed to have a direct relationship with the placebo effect (Kirsch 1985). It is suggested by enhancing the degree of expectancy, subsequently ones belief in the effectiveness of the placebo might increase (Evans 2003). The effects of expectancy on sport and exercise performance has been evaluated in a variety of different experimental models (Broatch et al. 2014; Maganaris et al. 2000; McClung and Collins 2007; Ross et al. 2015). For example, Maganaris et al., (2000) administered a saccharine placebo to eleven national level power lifters but provided false information to enhance expectancy of performance benefit. Participants were led to believe that they had consumed anabolic steroids which resulted in changes from baseline of 10±2 kg, 11±2 kg and 12±1 kg, for maximal bench press, dead lift and squat performance, respectively. However, when approximately half of the group were told the true nature of the placebo these performance gains almost completely dissipated although remained for the group who believed they had consumed anabolic steroids (Maganaris, et al. 2000).
To the best of our knowledge, only one study has previously evaluated the effects of expectancy on exercise performance with respect of NaHCO$_3$ ingestion (McClung and Collins 2007). Sixteen trained endurance athletes ran 5 x 1000 m time trials after ingesting either sodium bicarbonate (NaHCO$_3$) or a placebo in combination with being told correctly or deceived regarding what treatment had been administered (i.e. told NaHCO$_3$/placebo, given/not given NaHCO$_3$/placebo). The told NaHCO$_3$/given placebo treatment performed the time trial significantly faster (-3.4 s / -1.8 %) than the told placebo/given NaHCO$_3$ treatment demonstrating the effects of expectancy on exercise performance were greater than the purely pharmacological effects of NaHCO$_3$.

As there has been limited research on the potential psychobiological impact of NaHCO$_3$ administration on exercise performance (McClung and Collins 2007) and based on previous data that suggests GI distress does not necessarily negatively influence exercise performance (Higgins et al. 2013; Price and Simons 2010) we speculated that replicating side effects of NaHCO$_3$ ingestion might initiate a degree of expectancy of ergogenicity which might influence exercise performance. Therefore, the aim of this study was to evaluate the effect of expectancy associated with NaHCO$_3$ ingestion on high intensity cycling capacity. We hypothesised that by increasing the expectancy of ergogenicity a relatively inert compound (NaCl) mimicked as closely as possible and described as NaHCO$_3$ would induce a significant increase in cycling capacity.
Materials and methods

Participants

Eight healthy and active male participants volunteered to take part in this study (age: 21±1 years, body mass: 75±8 kg, height: 178±4 cm) which received university ethics committee approval. All participants were recreationally active although not specifically cycle trained ($W_{PEAK}$: 204±23 W). In order to maximise the integrity of the treatment deception all participants were pre-screened to ensure they had not used alkalotic buffers for a minimum of 3 months. Furthermore, no participants had ever previously consumed NaHCO$_3$ and only one had ever heard of its use with respect of sport and exercise performance.

Pre-experimental procedures

Participants were requested to avoid alcohol and strenuous exercise 12 and 24 hours prior to exercise, respectively. Participants were also requested to adopt the same balanced diet for 24 hours prior to each trial with adherence monitored via self-reported food diaries. Participants were specifically requested to avoid low carbohydrate intake which can induce mild metabolic acidosis and subsequently negatively affect exercise capacity (Greenhaff et al. 1988). The aforementioned information was included in the participant information sheet (PIS) and confirmed verbally before written consent was provided. Each participant completed physical activity readiness
(PAR-Q) and blood screening questionnaires prior to the commencement of the study. Specifically, the blood screening questionnaire asked participants to report any serious infections, jaundice, haemophilia or anything that might be hazardous to the health of the participant or researcher. Individuals were only permitted to participate if they were deemed healthy and able to do so. Participants attended the laboratory at the same time of the day for each visit to minimise the influence of circadian changes on exercise performance (Fernandes et al. 2014). Trials were carried out 3 to 7 days apart.

Protocol

On the first visit to the laboratory, participants performed a graded incremental test to assess peak power output ($W_{\text{PEAK}}$) on a cycle ergometer (Ergomedic 884E 1 – Sprint Cycle, Monark, Sweden). Participants selected the seat and pedal strap positions that felt most comfortable with these settings being adopted for all subsequent trials. After 5 min seated rest a finger prick capillary blood sample was taken and analysed for blood lactate concentration ([BLa]) (Biosen C_line, EKF Diagnostic, Magdeburg, Germany). All participants’ blood samples were analysed within ~ 10 minutes of the participant leaving the laboratory.

Cycling commenced on the unloaded ergometer (70 W) at a cadence of 70 rev.min$^{-1}$ with an increase of 35 W every 3 minutes until volitional exhaustion. Further blood samples were taken at the end of exercise and 5 min post-exercise. Exhaustion was assumed via voluntary stoppage or the
cadence dropping below 70 rev.min\(^{-1}\) for longer than 5 seconds. Peak power output \(W_{\text{PEAK}}\) was calculated as the mean power achieved during the final minute of the test (Lamberts et al. 2012). If exhaustion occurred less than one minute into a stage the appropriate duration undertaken at each power output was used to calculate a pro-rata \(W_{\text{PEAK}}\) (Higgins et al. 2013).

The second trial was a \(T_{\text{LIM}}\) familiarisation at 100% \(W_{\text{PEAK}}\). After 5 min seated rest ratings of gut fullness (GF) and abdominal discomfort (AD) were recorded against an 11 point (0 to 10) scale with 0 representing ‘empty’ and ‘completely comfortable’, respectively and 10 representing ‘bloated’ and ‘unbearable pain’, respectively (Higgins et al. 2013). A fingerpick capillary blood sample was then taken and analysed for [BLa] and blood glucose concentration [BG] (Biosen C_line, EKF Diagnostic, Magdeburg, Germany). Participants then ingested a control solution consisting of 1 ml.kg\(^{-1}\) double strength no added sugar orange squash with sweeteners (Cooperative Group, Manchester, UK) mixed with 4 ml.kg\(^{-1}\) tap water in the first 5 min of a 60 min seated rest period with GF and AD recorded at the end of this 5 min period. Prior to commencing exercise (i.e. at end of 60 min rest), heart rate (HR; F1 Polar Heart Rate Monitor, Polar, Finland), GF and AD were recorded. Participants then warmed up for 3 minutes on the unloaded ergometer (70 W) at a cadence of 70 rev.min\(^{-1}\). After a brief rest (~ 60 sec) participants commenced exercise from a stationary start position and were counted down from 5 seconds (Wittekind et al. 2011). Exhaustion was assumed as previously described. Measures of AD and GF were recorded again at the end of exercise and 5 min post-exercise. Heart rate, ratings of
perceived exertion specific to cardiovascular strain (RPE\(_{C}\)) and specific to the leg musculature (RPE\(_{L}\)) were recorded at 1 min intervals during exercise and at the end of exercise. The 15 point (6-20 category-ratio scale (CR-15) was used to assess RPE\(_{C}\) which is deemed to have a linear relationship with work load, oxygen consumption and heart rate (Borg 1982). In contrast, the 10 point (1-10) category-ratio scale (CR-10) was used to assess RPE\(_{L}\). This approach has been shown to correlate better with sensations associated with lactate accumulation (McClung and Collins 2007). Further blood samples for [BLa] and [BG] were taken at the end of exercise and 5 min post-exercise. After final measurements had been collected and after completing a self-selected warm down participants left the laboratory.

On the third and fourth visits to the laboratory participants completed experimental trials which were identical to the aforementioned exercise protocol with the following amendments. Prior to treatment ingestion participants were provided with a written manuscript which detailed the proposed effects of the subsequent treatment. For PLA the manuscript suggested that NaCl was relatively inert and would have little impact on exercise performance, whereas for SHAM the ‘proven’ ergogenicity of NaHCO\(_{3}\) was presented and in line with the AD and GF scales adopted the potential for associated feelings of GI distress such as build of gas and bloating were outlined. Participants then ingested either 0.1 g.kg\(^{-1}\) sodium chloride (NaCl) mixed with orange squash as previously described and either 4 ml.kg\(^{-1}\) tap water (placebo; PLA), or 4 ml.kg\(^{-1}\) carbonated water (SHAM placebo; SHAM). Carbonated water was used for SHAM to differentiate from
PLA and to attempt to induce mild GI symptoms often associated with NaHCO$_3$ ingestion such as gas/bloating. Control solutions demonstrated that the osmolality of PLA and SHAM were almost identical (510 ± 0 mOsmol/kg$^{-1}$ and 512 ± 10 mOsmol/kg$^{-1}$, respectively). Solutions were prepared by a technician not involved in data collection and both treatments and manuscripts were administered double blind. Trials were counterbalanced to minimise any potential order effects.

Statistical analysis

Statistical analysis was completed using SPSS (IBM v21, Chicago, USA). For all data normality (Shapiro-Wilk) and homogeneity of variance/sphericity (Mauchly) were checked prior to choosing the appropriate statistical tests. For 2-way repeated measures ANOVAs ([BLa], [BG]) Bonferroni corrections were applied. In some instances an ANOVA was chosen despite the majority of data not being normally distributed (i.e. AD and GF). This was decided so as to minimise the potential for type-I error due to multiple individual (non-parametric) comparisons and because of the general robustness of the univariate ANOVA test to violations of normality (Schmider et al. 2010). Additionally although RPE$_O$, RPE$_L$ and HR were collected during each minute of T$_{LIM}$, due to the variation in performance and small samples sizes at some time points analysis has only been completed between treatments for 1 to 7 minutes and at the end of exercise where at least 50% of data points were available for both treatments (minimum of n=4 comparison). With the exception of after 4 min T$_{LIM}$, where a Mann-Whitney
U test was used (for unequal sample size comparisons), RPE₀, RPE₅ and HR were analysed by paired t-tests or Wilcoxon tests. Exercise capacity (T_LIM) was analysed using a paired t-test.

Data were analysed and quantified using a mixture of effect sizes (ES), P values (minimum P ≤ 0.05) and, where appropriate, 95% confidence intervals (Watt et al. 2002). Unless otherwise stated data is presented as mean ± standard deviation. For ANOVA main effects and interactions the ES is reported as the partial η² value. Otherwise, for normally distributed data the ES (d) was calculated using the difference in means divided by the pooled SD of the compared trials (Nakagawa and Cuthill 2007).
Results

Exercise capacity

Mean $T_{\text{LIM}}$ increased by 9.5% (40±47 s; 95%CI: 0.4 to 80 s) for SHAM vs. PLA (461±148 s vs. 421±150 s; $P=0.048$, $d=0.3$; Figure 1). No order effect was found between trials (448±130 s vs. 435±168 s; $P=0.574$, $d=0.1$).

****Figure 1 near here****

Perceptual variables

There was no time and treatment interaction for GF ($P=0.276$, $\eta^2=0.2$). Although GF increased by ~ 1 unit at all time points post SHAM ingestion compared with PLA ingestion only a tendency towards a main effect for treatment was observed ($P=0.080$; $\eta^2=0.4$). In contrast, there was a main effect for time ($P<0.001$; $\eta^2=0.7$). More specifically, GF post-ingestion (1.7±1.1) was greater than rest (0.0±0.0; $P=0.002$; $d=2.1$), end of exercise (0.4±1.1, $P=0.031$; $d=1.2$) and 5 min post-exercise (0.3±0.9, $P=0.012$; $d=1.4$). Pre-exercise, GF (0.7±0.9) was also greater than 5 min post-exercise ($P=0.025$; $d=0.4$; Figure 2). There was no time and treatment interaction for AD ($P=0.451$, $\eta^2=0.1$). Similar to GF, although AD increased by ~ 1 unit at all time points post SHAM ingestion compared with PLA ingestion, only a tendency towards a main effect for treatment was observed ($P=0.096$;
Unlike GF there was no main effect for time for AD (P=0.299, $\eta^2=0.2$).

****Figure 2 near here****

After 3 min $T_{\text{LIM}}$ RPE$_O$ was $1.4\pm1.3$ units lower for SHAM ($12.9\pm2.5$) units compared to PLA ($14.3\pm2.5$; P=0.020, d=0.6). Although none of the other comparisons between treatments were significantly different, RPE$_O$ was also lower for SHAM vs. PLA after 2, 5, 6 and 7 min and at the end of exercise (Figure 3). There were no differences between treatments for RPE$_L$ although values were lower for SHAM vs. PLA after 3, 5, 6 and 7 min and at the end of exercise.

****Figure 3 near here****

Blood and cardiorespiratory variables

There was no time and treatment interaction (P=0.149, $\eta^2=0.2$) or main effect for treatment for [BLa] (P=0.552, $\eta^2=0.1$). However, there was a main effect for time (P<0.001, $\eta^2=0.9$). At the end of exercise $8.91\pm1.70$ mmol.l$^{-1}$ and 5 min post-exercise ($6.57\pm1.10$ mmol.l$^{-1}$) [BLa] was greater than at rest ($1.27\pm0.20$ mmol.l$^{-1}$; both P<0.001 and d>6). Additionally, [BLa] was lower 5 min post-exercise compared to the end of exercise (P=0.004, d=1.6). There was no time and treatment interaction (P=0.210, $\eta^2=0.2$) or main effect for treatment for [BG] (P=0.629, $\eta^2<0.1$). However, there was a
main effect for time ($P<0.001$, $\eta^2=0.9$). At the end of exercise (3.68±0.22 mmol.l$^{-1}$) and 5 min post-exercise (3.54±0.26 mmol.l$^{-1}$) [BG] was lower than at rest (4.64±0.40 mmol.l$^{-1}$; both $P\leq0.001$; $d>3$). There was no difference between treatments for HR although HR increased linearly over time. At the end of exercise HR for SHAM (174±9 bpm$^{-1}$) and PLA (173±13 bpm$^{-1}$) was greater than at rest (77±9 bpm$^{-1}$ and 79±7 bpm$^{-1}$, respectively, both $P<0.001$, $d>9$).
Discussion

This study evaluated the effect of providing pre-exercise information designed to increase expectancy associated with NaHCO$_3$ ingestion on high intensity cycling capacity in recreationally active males. In support of our original hypothesis mean $T_{\text{Lim}}$ for SHAM was 9.5% longer (40 s) than PLA. In terms of contextualising the magnitude of this change, this is considered a moderate improvement. More simply the effect size of 0.3 sits between the 0.2 (small) and 0.5 (medium) benchmarks as originally suggested by Cohen (1988). Interestingly, this magnitude of improvement is in line with the range of effect sizes (mean: 0.69; 95%CI: -0.07-1.63) for untrained individuals who have actually ingested NaHCO$_3$ prior to exercise using a time to volitional fatigue ($T_{\text{Lim}}$) protocol (Peart et al. 2012).

Beedie et al. (2007) suggest that placebo effects in sports performance are likely when individuals believe that they have ingested an ergogenic substance. Similar to McClung and Collins (2007), manipulative scripts were used in the present study to influence participant perception as to what they would be ingesting. Additionally, SHAM was mixed with carbonated water to try and replicate the reported side effects of NaHCO$_3$ ingestion. Although symptoms were mild, in the present study gut fullness increased by ~ 1 unit for all trials post SHAM ingestion compared with PLA suggesting this strategy might have contributed to any increase in expectancy experienced by participants. It is important to highlight that we purposefully chose to not replicate the experimental design as adopted by
McClung and Collins (2007). Due to the well acknowledged variety and sometimes severe GI responses (Cameron et al. 2010; Carr et al. 2011) it was felt that actually administering NaHCO$_3$ to participants could render the SHAM treatment ineffective in enhancing expectancy due to participants associating more severe GI symptoms with ‘genuine’ NaHCO$_3$ ingestion (Cameron et al. 2010). As such by comparing only PLA and SHAM, whose only differences were the information given pre-exercise and whether the water was tap or carbonated, we have been able to isolate the effects of pre-exercise information designed to increased expectancy of ergogenicity with NaHCO$_3$ ingestion on exercise capacity. It appears that this strategy has contributed to the observed increase in exercise capacity, at least in some individuals.

Compared with PLA RPE$_O$ was 1.4±1.3 units lower for SHAM after 3 min T$_{LIM}$ with the associated effect size (0.6) equivalent to a medium to large magnitude of change (Cohen 1988). Moreover, both RPE$_O$ and RPE$_L$ were generally lower for SHAM versus PLA throughout T$_{LIM}$. These results suggest that the information given to participants prior to exercise might have led to reduced perception of effort during exercise for SHAM which is likely to have contributed to the increase in T$_{LIM}$. Our results are supported by McClung and Collins (2007) who reported that RPE (based on perceptions of lactate accumulation and therefore most similar to RPE$_L$ in the present study) was 1.5 units lower for the told NaHCO$_3$/given placebo trial compared to the told placebo/given placebo trial with the told NaHCO$_3$/given placebo trial performing the 1 km time trial -2.8 s / -1.5% faster. Interestingly, both results
are similar to Higgins et al. (2013) who reported RPE \(_L\) was 1.5 units lower after both 1 and 2 min of T\(_{\text{lim}}\) cycling at 100% \(W_{\text{peak}}\) after NaHCO\(_3\) ingestion. Moreover, the authors reported significant correlations between T\(_{\text{lim}}\) and RPE\(_L\) at the same time points. Therefore, it is plausible that reductions in RPE and any impact on exercise performance are not solely due to NaHCO\(_3\) induced mediation of acid-base balance (i.e. the pharmacological effects). These results are in accordance with the psychobiological model which suggests that reduced perception of effort can increase exercise performance (Ross et al. 2015; Smirmaul, et al. 2013) with perception of effort postulated as a key determinant of exercise tolerance (Marcora and Staiano 2010). The reduction in perception of effort might be linked to the mediation of endogenous opioids via expectancy orientated placebo effects (Benedetti et al. 2005).

In the present study the mean 9.5% (+40 s) increase in exercise capacity after SHAM is substantially larger than the -1.5 % (-2.8 s) improvement for the placebo/told NaHCO\(_3\) trial compared with the given placebo/told placebo trial reported by McClung and Collins (2007). This is likely due to a number of methodological differences between studies such as exercise protocol (capacity test vs. time trial), exercise modality (cycling ergometry vs. track running), participant training status (recreationally active vs. trained), placebo dosage (0.1 g.kg\(^{-1}\) vs. “pinch” of NaCl), volume (5 ml.kg\(^{-1}\) vs. standardised 750 ml) and choice of solvent (carbonated water vs. (presumably) tap water) used to mix experimental solutions. However, it seems clear from both studies that deceptive information given pre-exercise
with regards to NaHCO₃ ingestion has subsequently improved performance when compared with placebo.

Despite the expected increases in [BLa] and HR and decreases in [BG] from rest to post-exercise, respectively, in the present study there were no interactions or main effects for treatment for any of the cardiorespiratory or haematological data collected. The lack of difference between treatments for [BLa] after exercise in the present study (SHAM: 8.84 ± 1.72 mmol.l⁻¹, PLA: 8.99 ± 1.79 mmol.l⁻¹) is in accordance with McClung and Collins (2007). Taken together, these data suggest that these physiological variables did not contribute to the observed differences in Tₘₜₐₓ.

To the best of our knowledge the daily variation for Tₘₜₐₓ at 100% Wₚₑᵃᵏ in recreationally active participants is currently unknown. Therefore, we have compared participant’s individual data to the daily variation of ~11% / 24 s reported for participants of a similar training status who engaged in two bouts of Tₘₜₐₓ cycling at 110% Wₚₑᵃᵏ (Higgins et al. 2014). In the present study five of the eight participants increased Tₘₜₐₓ for SHAM above daily variation (range: 12% to 32%; 53 to 104 s) with two participants showing no difference (range: -2% to 3%; -10 to 14 s) with only one recording lower Tₘₜₐₓ for SHAM (-9%; -39 s). It is unclear why expectancy appears to have influenced some but not all individuals but it is plausible that individuals might respond differently to expectancy at different exercise intensities.
It is important to acknowledge that there are some limitations to our study. Firstly, the present study examined the effects of expectancy related to an oral bolus solution ingested 60 min prior to exercise. This method was chosen because NaHCO$_3$ based solutions are widely used in contemporary research (Higgins et al. 2013; McClung and Collins 2007; Price and Simons 2010). In contrast NaHCO$_3$ is also often administered via capsules and/or ingestion is staggered over multiple dosages (Carr et al. 2011; Krstrup et al. 2015). Further research is warranted to examine whether these ingestion methods report results similar to the present study. Secondly, although limited anecdotal evidence was recorded (e.g. participant one left the lab stating “that stuff definitely works!”; analysis after the study revealed this was for a SHAM trial which translated into a 65 s / 13% increase compared with PLA) we did not carry out post-hoc qualitative analysis which could have provided important information to verbally confirm the presence or not of expectancy at an individual level. Finally, as the placebo effect is a psychobiological phenomenon that is linked to different mechanisms including Pavlovian conditioning and expectation of improvement, it is important to acknowledge that there is no singular placebo effect. As such, different mechanisms might be responsible in different conditions (Benedetti et al. 2005). That said, according to McClung and Collins (2007) the vast majority of placebo effects are due to expectancy and in the context of our study design it seems likely expectancy has played at least some role in facilitating improvements in $T_{LIM}$. 

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From a practical application perspective, the results from this study suggest that athletes and coaches should consider psychobiological strategies which could positively impact physical training and/or performance to a similar extent as reported pharmacological agents. This might be especially important for individuals who have empirical evidence that \( \text{NaHCO}_3 \) induced GI distress negatively impacts their performance. Indeed, the development of psychological skills can be as important as physiological adaptations when looking to improve sport and exercise performance (Beedie and Foad 2009).

In conclusion, providing pre-exercise information designed to increase expectancy associated with \( \text{NaHCO}_3 \) ingestion, administered via an oral bolus solution, improved mean cycling capacity at 100% \( \text{W}_\text{PEAK} \) by \( \sim 10\% \). In conjunction with the psychobiological model of exercise performance the increase in exercise capacity was likely facilitated by reduced perception of overall effort, possibly due to mediation of endogenous opioids via expectancy orientated placebo effects. The observed ergogenicity with SHAM did not affect our measures of cardiorespiratory physiology or metabolic flux.
Conflict of interest statement

The authors declare that there is no conflict of interest associated with this study.
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Figure Legends

Figure 1 – Individual time until volitional exhaustion ($T_{\text{LIM}}$; s) at 100% $W_{\text{PEAK}}$ for SHAM vs. PLA. SHAM > PLA ($P=0.048$, $d=0.3$).

Figure 2 – Perceptual ratings of gut fullness (GF) over time for SHAM vs. PLA. Error bars represent 1 SD ($^a=$SHAM, $^b=$PLA). * Post-ingestion > Rest ($P=0.002$), $^#$ Post-ingestion > end of exercise ($P=0.031$), $^\$ Post-ingestion > 5 min post-exercise ($P=0.012$), ** Pre-exercise > 5 min post-exercise ($P=0.025$).

Figure 3 – Ratings of perceived exertion representative of cardiovascular strain ($RPE_0$) over time for SHAM vs. PLA. Error bars represent 1 SD. * SHAM < PLA ($P=0.020$).