



Dietary nitrate supplementation to enhance exercise capacity in hypoxic COPD: EDEN-OX, a double-blind, placebo-controlled, randomised cross-over study

Matthew J Pavitt,¹ Adam Lewis ¹, Sara C Buttery,¹ Bernadette O Fernandez,² Monika Mikus-Lelinska,² Winston A S Banya,¹ Martin Feelisch,^{3,4} Michael I Polkey,^{1,5} Nicholas S Hopkinson ¹

► Additional supplemental material is published online only. To view, please visit the journal online (<http://dx.doi.org/10.1136/thoraxjnl-2021-217147>).

¹National Heart and Lung Institute, Imperial College London, London, UK

²Clinical and Experimental Sciences, Southampton Hospital, Southampton, UK

³Faculty of Medicine, Clinical and Experimental Sciences, University of Southampton, Southampton, UK

⁴Southampton NIHR Respiratory Biomedical Research Unit, Southampton General Hospital, Southampton, UK

⁵Respiratory Medicine, Royal Brompton Hospital, London, UK

Correspondence to

Professor Nicholas S Hopkinson, National Heart and Lung Institute, Imperial College London, London SW7 2BX, UK; n.hopkinson@ic.ac.uk

Received 23 February 2021
Accepted 4 November 2021
Published Online First
1 December 2021



© Author(s) (or their employer(s)) 2022. No commercial re-use. See rights and permissions. Published by BMJ.

To cite: Pavitt MJ, Lewis A, Buttery SC, et al. *Thorax* 2022;**77**:968–975.

ABSTRACT

Rationale Dietary nitrate supplementation improves skeletal muscle oxygen utilisation and vascular endothelial function. We hypothesised that these effects might be sufficient to improve exercise performance in patients with COPD and hypoxia severe enough to require supplemental oxygen.

Methods We conducted a single-centre, double-blind, placebo-controlled, cross-over study, enrolling adults with COPD who were established users of long-term oxygen therapy. Participants performed an endurance shuttle walk test, using their prescribed oxygen, 3 hours after consuming either 140 mL of nitrate-rich beetroot juice (BRJ) (12.9 mmol nitrate) or placebo (nitrate-depleted BRJ). Treatment order was allocated (1:1) by computer-generated block randomisation.

Measurements The primary outcome was endurance shuttle walk test time. The secondary outcomes included area under the curve to isotime for fingertip oxygen saturation and heart rate parameters during the test, blood pressure, and endothelial function assessed using flow-mediated dilatation. Plasma nitrate and nitrite levels as well as FE_{NO} were also measured.

Main results 20 participants were recruited and all completed the study. Nitrate-rich BRJ supplementation prolonged exercise endurance time in all participants as compared with placebo: median (IQR) 194.6 (147.5–411.7) s vs 159.1 (121.9–298.5) s, estimated treatment effect 62 (33–106) s ($p<0.0001$). Supplementation also improved endothelial function: NR-BRJ group +4.1% (–1.1% to 14.8%) vs placebo BRJ group –5.0% (–10.6% to –0.6%) ($p=0.0003$).

Conclusion Acute dietary nitrate supplementation increases exercise endurance in patients with COPD who require supplemental oxygen.

Trial registration number ISRCTN14888729.

INTRODUCTION

People with COPD may develop hypoxaemia as the condition becomes more severe, impacting on their ability to perform day-to-day activities. Mechanisms include ventilation perfusion mismatch, reduced cardiac output due to hyperinflation and pulmonary vascular limitation, as well as reduced muscle efficiency.^{1–5} In individuals who are sufficiently hypoxaemic, long-term oxygen therapy (LTOT) improves survival, and in many individuals

Key messages

What is the key question?

⇒ Can dietary nitrate supplementation enhance exercise performance in individuals with a hypoxic COPD phenotype?

What is the bottom line?

⇒ In a double-blind, placebo-controlled, randomised cross-over study, an acute dose of dietary nitrate increased endurance shuttle walk time in individuals with a hypoxic COPD phenotype.

Why read on?

⇒ As COPD becomes more severe, hypoxaemia may develop which impacts on the ability to perform day-to-day activities.
⇒ Interventions which improve endothelial function, as demonstrated here, and increase the efficiency of oxygen use may help to address this.

ambulatory oxygen therapy (AOT) improves exercise performance.⁶

Nitric oxide (NO) has potential as a modulator of exercise performance. A ubiquitous signalling molecule, NO is involved in a number of processes at a tissue and cellular level, including mitochondrial and cellular respiration,^{7–8} glucose uptake into the skeletal muscle,⁹ skeletal muscle contraction,^{10–11} neurotransmission¹² and fatigue development.¹³ NO is produced both by oxygen-dependent NO synthases catalysing its production from L-arginine and an alternative nitrate (NO_3^-)–nitrite (NO_2^-)–NO pathway.¹⁴ The latter can be influenced by supplementation with exogenous dietary NO_3^- and is enhanced in conditions of hypoxia and low pH as found in exercising skeletal muscle.¹⁵

Dietary NO_3^- supplementation has been shown to reduce the oxygen cost of exercise in healthy individuals in normoxic conditions^{16–17} and in conditions of hypoxaemia.^{18–23} Recently our research group has shown that it augments improvement in exercise capacity seen in people with COPD following pulmonary rehabilitation.^{24–25} We have also previously shown that dietary NO_3^- supplementation reduces the oxygen cost of exercise during



endurance cycle ergometry in COPD.²⁶ However, that study, which excluded patients who required supplemental oxygen, did not demonstrate an improvement in exercise capacity.

The aim of the present study was therefore to assess the acute effect of dietary supplementation in the form of nitrate-rich beetroot juice (NR-BRJ) on exercise performance in individuals with COPD who require supplemental oxygen on exertion, hypothesising that this would increase exercise capacity, measured as endurance shuttle walk time (ESWT), as well as improve endothelial function in people with this specific phenotype.

MATERIALS AND METHODS

Study design

The EDEN-OX (Effect of Dietary Nitrate Supplementation on Exercise Performance in Hypoxia) study was a single-centre, double-blind, placebo-controlled, randomised cross-over trial comparing the effects of dietary NO_3^- supplementation with a matched placebo in individuals with COPD who require LTOT and use AOT during exercise. All participants provided informed consent. The study was registered prospectively in a publicly accessible database. The data presented here relate only to the planned COPD cohort in that study.

People with Global Initiative for Chronic Obstructive Lung Disease grade II–IV COPD²⁷ who were established users of LTOT, in accordance with the NICE guidelines,²⁸ were recruited from the outpatient clinical services of Royal Brompton and Harefield NHS Foundation Trust (North West London) between 4 November 2016 and 8 August 2017, with the last participant's final visit completed on 15 January 2018.

Exclusion criteria for the study included clinical instability (ie, less than 1 month after an exacerbation), significant comorbidity limiting exercise tolerance, significant renal impairment (estimated glomerular filtration rate <50 mL/min), hypotension (systolic blood pressure <100 mm Hg), pregnancy, use of NO_3^- -based medicine or phosphodiesterase V inhibitors, or presence of other conditions that might be influenced by NO_3^- supplementation (ie, ischaemic heart disease or peripheral vascular disease). These conditions were assessed at the screening visit through review of clinical history and assessment of relevant clinical data.

Methods

Interventions

The intervention was a commercially available concentrated NO_3^- -rich BRJ (NR-BRJ) (98%) drink cut with organic lemon juice (2%) containing 0.8 g, 12.9 mmol NO_3^- (140 mL Beet-It Sport Shot, James White Drinks, Ipswich, UK). The placebo beetroot juice (PL-BRJ), produced by James White, was 140 mL of the same beverage in which NO_3^- was removed by a standardised method of passing the juice, prior to pasteurisation, through an ion exchange column, containing Purolite A520E, which exchanges NO_3^- against chloride.²⁹ The PL-BRJ is identical in appearance, packaging, taste and smell, and also causes beeturia (orange to red discolouration of urine).

Study conduct

At an initial baseline visit, COPD Assessment Test, Hospital Anxiety and Depression Scale, and Medical Research Council Dyspnoea Scale scores were recorded. Body composition was measured by bioelectrical impedance analysis using a Bodystat 4000 device (Bodystat, Isle of Man, UK). Participants then performed two incremental shuttle walk tests to determine the walking speed to be used for the ESWT³⁰ and then a practice

ESWT. All walking tests throughout the study were performed on the participant's usual AOT flow rate, and the method for carrying the AOT was recorded to ensure the same method was always used (online supplemental appendix figure E1).

Prior to the two subsequent intervention visits and throughout the study period, participants were asked to avoid the use of antimicrobial mouthwash and chewing gum, as these have been shown to reduce the oral facultative bacteria whose NO_3^- reductase activity is essential for the metabolism of an oral NO_3^- load.³¹ They were asked to consume the same meal on the morning of each study assessment. This was to create as standardised conditions as possible, reducing differing levels of dietary NO_3^- consumption as a source of variation within individuals, while not altering their usual diet greatly. They were also asked to match caffeine consumption to standardise any ergogenic effect arising from it³² and to avoid strenuous exercise in the 24-hour period prior to the intervention visits.

The two intervention visits began at the same time of day (± 2 hours), with a minimum of 7-day washout period and a maximum 1-month gap between them. Participants were randomly assigned to the order in which they received NR-BRJ or PL-BRJ using a computer-generated block randomisation list, with a block size of 10, produced by an independent statistician. The researchers responsible for enrolment and outcome measurements remained blinded throughout the study and during data analysis. Following their arrival, after a 10 min rest period, participants were observed consuming either the NR-BRJ or the PL-BRJ, and empty bottles were collected and recorded. All outcome measures were undertaken 3 hours after ingestion of either NR-BRJ or PL-BRJ.

Outcomes

Exercise capacity

The primary outcome was ESWT compared between treatment conditions. Given the cross-over design and taking 65 s (95% CI 45 to 85) to be the minimal clinically important difference (MCID) in ESWT³⁰ and a pooled mean difference within individuals of 26 s for repeat testing, to have an 80% statistical power, with a significance level of 0.05, 16 participants would be required to reject the null hypothesis that the active intervention was not superior to placebo. To allow for a 25% withdrawal rate, a sample size of 20 was chosen.

Plasma nitrate/nitrite levels and markers of oxidative stress

Plasma NO_3^- and NO_2^- levels were used as a combined biomarker of NO_3^- ingestion, metabolism and NO availability.^{33 34} Plasma samples were obtained on arrival and 3 hours after consumption of NR-BRJ or PL-BRJ (see online supplemental appendix for full details).

Oxidative stress biomarkers were assessed in plasma samples by a combination of three distinct readouts, including antioxidant potential, that is, measurement of the ferric-reducing ability of plasma (FRAP),³⁵ lipid oxidation products by thiobarbituric acid reactive substances (TBARS)³⁶ and total free thiols with normalisation for protein³⁷ (see online supplemental appendix for full details).

Fractional exhaled nitric oxide

FE_{NO} was measured as a steady exhalation rate of 50 mL/s with a NIOX Mino (Aerocrine Systems, Solna, Sweden) at the screening visit and then at the intervention visits at baseline prior to NR-BRJ/PL-BRJ consumption and then at six further intervals (30, 60, 90, 120, 150 and 180 min). Both the study participant

and the researchers were blinded to the results, and an independent researcher, not directly involved with the trial, uploaded the data into a password-protected database. These data were only available to the researchers following unblinding of the study.

Endothelial function

Endothelial function was assessed by flow-mediated dilatation (FMD) of the brachial artery 3 hours after NR-BRJ/PL-BRJ consumption³⁸ using a high-resolution Doppler ultrasound to measure at baseline and sequentially over a period of 120 s after release of circulatory arrest of the upper arm.³⁹ All measurements were performed by a single trained operator (see online supplemental appendix for full details).

Continuous oxygen saturation and heart rate analysis

For each ESWT performed, pulse oximetry values were recorded (Pulsox 300i Pulse Oximeter, Konica Minolta, Tokyo, Japan) throughout until the participant had recovered (recovery was defined by return of Borg Dyspnoea Scale to that recorded prior to the ESWT). To maintain blinding, the pulse oximeter display was covered throughout the testing and the data were downloaded by an independent researcher, not directly involved with the trial, who uploaded the data to a password-protected database. These data were only available to the researchers following unblinding of the trial.

Statistical analysis

Data are presented as mean (SD), or if not normally distributed as median and IQR. Differences in response between treatment conditions were assessed using a paired t-test or a Wilcoxon signed-rank test as appropriate. Treatment effect was estimated using the Hodges-Lehmann estimate of shift parameters. The process of determining the Hodges-Lehmann estimator entails estimating the average difference in outcomes (x-y) for every possible $n(n+1)/2$ pair

and then deriving the overall median of all averages (the Hodges-Lehmann estimator). A distribution-free CI is estimated using large-sample approximation. Analysis was performed using SPSS V.24 for Windows and Stata V.16.1 for Windows.

To compare continuous oxygen saturations (SpO_2) and heart rate (HR) between the two treatment conditions, individual ESWT data periods were subjected to a 30 s rolling average using MATLAB (MATLAB and Statistics Toolbox Release V.2017a, The MathWorks, Natick, Massachusetts, USA) and then expressed as percentages of isotime (defined as the duration of the shortest of the two ESWT). These individual responses were then grouped to allow analysis of HR and SpO_2 against the percentage of isotime (plotted at the midpoint of each 10th percentile of isotime). The area under the curve (AUC) was assessed for each individual participant and the two treatment conditions compared using Wilcoxon signed-rank test. Figures were prepared using GraphPad Prism V.6.0 for Windows (GraphPad Software, San Diego, California, USA). A p value of <0.05 was considered statistically significant.

RESULTS

We screened 67 people for eligibility (figure 1); 31 declined to participate, 7 had a comorbidity precluding participation and 9 were not using supplementary oxygen. Of the 20 participants enrolled in the study, 10 were randomised to receive PL-BRJ first and 10 NR-BRJ first. All participants completed the study. Table 1 shows their baseline characteristics, which were well matched between the two order allocation groups. There were no serious adverse effects reported, although all participants reported beeturia. The average time between each intervention visit was 7 days.

Exercise outcomes

Exercise endurance time was longer for all study participants after NR-BRJ compared with PL-BRJ (figure 2): median (IQR) ESWT: NR-BRJ 194.6 (147.5–411.7) s vs PL-BRJ 159.1

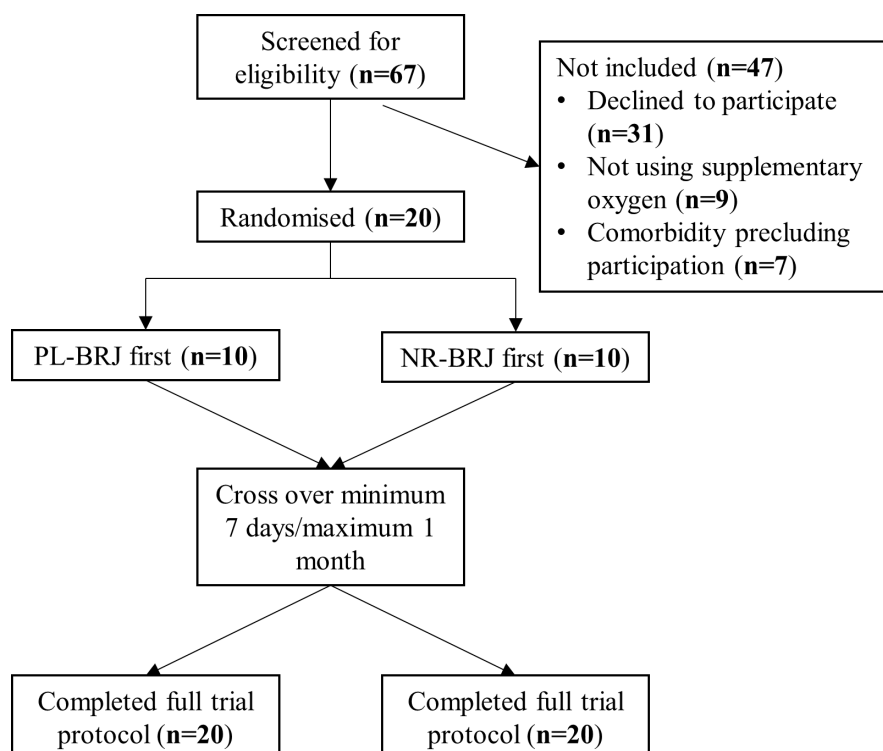


Figure 1 CONSORT diagram for recruitment and trial completion. CONSORT, Consolidated Standards of Reporting Trials; NR-BRJ, nitrate-rich beetroot juice; PL-BRJ, placebo beetroot juice.

Table 1 Characteristics of cross-over allocation groups: NR-BRJ or PL-BRJ received first

Measurement	NR-BRJ first (n=10)	PL-BRJ first (n=10)	P value	Whole group (N=20)
Sex (% female: male)	30:70	50:50	1.0	40:60
Age (years)	68 (62–73)	67 (64–76)	0.7	67.6 (8.5)
Caucasian (%)	100	100	1	100
Smoking (pack years)	28 (14–50)	64 (57–96)	0.006	52 (21.6)
BMI (kg/m ²)	24.3 (20.9–29.0)	26.2 (21.4–30.1)	0.5	25.2 (4.7)
FFMI (kg/m ²)	18.4 (15.8–20.1)	18.0 (14.0–20.2)	0.8	18.1 (15.8–19.9)
Inhaled medications				
SABA (%)	91	100	0.4	95
LABA-ICS (%)	91	78	0.4	95
LAMA (%)	91	100	0.4	85
Baseline resting oxygen saturation FiO ₂ 0.21 (%)	92 (89–94)	92 (89–93)	0.8	92
Pre-LTOT prescription baseline pO ₂ (kPa)	6.9 (6.0–7.3)	6.6 (6.4–7.2)	0.6	6.8
Oxygen prescription (L/min)	4 (2–6)	2 (2–4)	0.2	3.0 (2.0–6.0)
CAT score	20 (18–29)	19 (15–28)	0.6	21 (8.0)
MRC Dyspnoea Score	4 (4–4)	4 (4–4)	1.0	4 (4–4)
HADS-A score	4 (3–7)	7 (2–10)	0.3	4.0 (2.3–8.8)
HADS-D score	4 (4–5)	5 (4–7)	0.7	4.5 (4.0–5.6)
Systolic BP (mm Hg)	139 (123–149)	135 (115–139)	0.1	137 (121–143)
Diastolic BP (mm Hg)	76 (66–83)	70 (65–80)	0.4	73 (65–82)
MAP (mm Hg)	94 (91–104)	91 (85–94)	0.2	92 (80–100)
Lung function				
FEV ₁ (L)	0.7 (0.6–1.0)	0.7 (0.3–1.0)	0.3	0.7 (0.6–1.0)
FVC (L)	2.7 (1.9–3.1)	1.6 (1.4–3.2)	0.3	2.7 (1.6–3.1)
FEV ₁ :FVC ratio	0.3 (0.3–0.3)	0.3 (0.2–0.4)	1.0	0.3 (0.3–0.3)
RV %predicted	211 (181–235)	212 (188–233)	0.8	212 (186–233)
TL _{co} %predicted	33 (19–45)	36 (28–44)	0.9	32 (19–44)
GOLD stage				
III (%)	22	45	1.0	35
IV (%)	78	55	1.0	65
ISWT distance (m)	300 (280–360)	370 (220–280)	0.04	279 (70)
ESWT (s)	172 (137–267)	181 (158–193)	0.6	179 (152–193)

Data in order of intervention, either NR-BRJ or PL-BRJ first.
 Data shown are median (IQR), mean (SD) or percentage (%).
 P value is for independent t-test or Mann-Whitney test comparing groups.
 BMI, body mass index; BP, blood pressure; CAT, COPD Assessment Test; ESWT, endurance shuttle walk test; FFMI, fat-free mass index; FiO₂, fraction of inspired oxygen; GOLD, Global Initiative for Chronic Obstructive Lung Disease; HADS-A, Hospital Anxiety Depression Scale-Anxiety; HADS-D, Hospital Anxiety Depression Scale-Depression; ICS, inhaled corticosteroid; ISWT, incremental shuttle walk test; LABA, long-acting beta agonist; LAMA, long-acting muscarinic agonist; LTOT, long-term oxygen therapy; MAP, mean arterial pressure; MRC, Medical Research Council; NR-BRJ, nitrate-rich beetroot juice; PL-BRJ, placebo beetroot juice; RV, residual volume; SABA, short-acting beta agonist; TL_{co}, transfer Factor for carbon monoxide corrected for haemoglobin.

(121.9–298.5) s, estimated treatment effect 62.5 (95% CI 33 to 106) s ($p=0.000089$). There was no evidence of an intervention order effect (online supplemental appendix figure E2). There was one individual who was a clear outlier for exercise endurance response. However, a sensitivity analysis removing their data led to a slight change in primary study outcome but not the overall statistical significance: median (IQR) ESWT: NR-BRJ 193.8 (145–389.6) s vs PL-BRJ 158.2 (121.6–236.6) s, estimated treatment effect 56.5 (95% CI 30 to 88) s, indicating a significant increase in ESWT with NR-BRJ ($p=0.0001$) (online supplemental appendix results E1).

Pulse oximetry data were available for only 18 participants because recording failed for 2 of them. The average AUC for SpO₂ was higher in the NR-BRJ group compared with the

PL-BRJ group. These differences were more apparent at isotime and peak exercise, with no difference at rest, during warm-up or recovery (figure 3A and online supplemental appendix table E1). The estimated treatment effect was also statistically significant: 43.69 (29.09–58.28) ($p<0.0001$). The AUC for HR response to NR-BRJ or PL-BRJ did not show any difference. The estimated treatment effect was also not statistically significant (–41.17 (–116.74 to 34.40), $p=0.27$) (figure 3B and online supplemental appendix table E1).

Endothelial function and blood pressure

Two participants declined the FMD assessment; therefore, data were available for 18 participants. At 180 min following dosing,

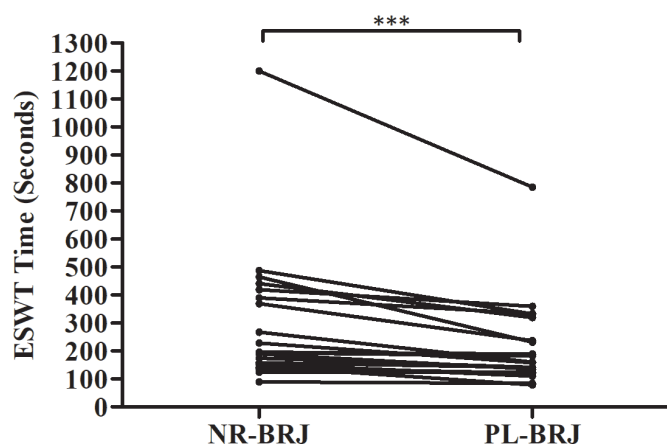


Figure 2 Effect of dietary nitrate supplementation on ESWT (in seconds) for PL-BRJ and NR-BRJ dosing conditions. Data presented as individual ESWT (in seconds) in both dosing conditions. Wilcoxon signed-rank test was used to compare ESWT between the different dosing conditions: NR-BRJ 194.6 (147.5–411.7) s vs PL-BRJ 159.1 (121.9–298.5) s, estimated treatment effect 62 s (95% CI 33 to 106). *** $p < 0.0001$. ESWT, endurance shuttle walk test; NR-BRJ, nitrate-rich beetroot juice; PL-BRJ, placebo beetroot juice.

FMD increased: NR-BRJ 4.1% (–1.1% to 14.8%) compared with placebo –5.0% (–10.6% to –0.6%), estimated treatment effect –11.9% (95% CI –18.9 to –7.15) ($p = 0.0003$) (figure 4).

There was no statistically significant difference in change in blood pressure parameters from predosing levels between NR-BRJ and PL-BRJ (figure 5): median (IQR) Δ systolicBP: NR-BRJ –1.5 (15.0–10.8) mm Hg vs PL-BRJ –0.5 (–10.5 to 6.8) mm Hg, estimated treatment effect 1 mm Hg (95% CI –5.5 to 7.0) ($p = 1.0$); Δ diastolicBP: NR-BRJ –4.0 (–14.0 to 7.0) mm Hg vs PL-BRJ –1.0 (–9.3 to 5.0) mm Hg, estimated treatment effect 1 mm Hg (95% CI –3 to 5) ($p = 0.481$); mean arterial pressure: NR-BRJ –5.0 (–15.3 to 6.0) mm Hg vs PL-BRJ –2.5 (–13.5 to 7) mm Hg, estimated treatment effect 1.5 mm Hg (95% CI –3.5 to 5) ($p = 0.359$).

Plasma nitrate and nitrite levels and oxidative stress markers

Paired data on plasma NO_2^- and NO_3^- concentrations were available for 19 participants, as 1 individual declined sampling. Following supplementation with NR-BRJ, there was an 84% increase in plasma NO_2^- and an 887% increase in plasma NO_3^- at 180 min post supplementation, but no change with placebo (figure 6 and online supplemental appendix table E2). Both the NR-BRJ and PL-BRJ supplements were analysed for NO_2^- and NO_3^- content as well (online supplemental appendix table E3). The change in plasma NO_3^- and NO_2^- from baseline to 180 s was calculated and used to estimate the treatment effect of NR-BRJ. The treatment effect of NO_3^- was 550 (461–639) μM . The results suggest that this was higher for NR-BRJ than for PL-BRJ and this change was statistically significant ($p = 0.0003$). The treatment effect of NO_2^- was 0.248 (0.138–0.408) μM . The results suggest that this was higher for NR-BRJ than for PL-BRJ and this change was statistically significant ($p = 0.0011$).

There was no statistically significant difference in measures of oxidative stress following acute consumption of either supplement: FRAP: NR-BRJ 1018 (853.0–1125) μM vs PL-BRJ 930.2 (836.8–1073) μM ($p = 1.0$); TBARS: NR-BRJ 1.499 (0.855–3.209) mM vs PL-BRJ 0.971 (0.766–1.614) mM ($p = 0.4$); total free thiol per protein: NR-BRJ 7.079 (5.961–8.115) $\mu\text{mol/g}$

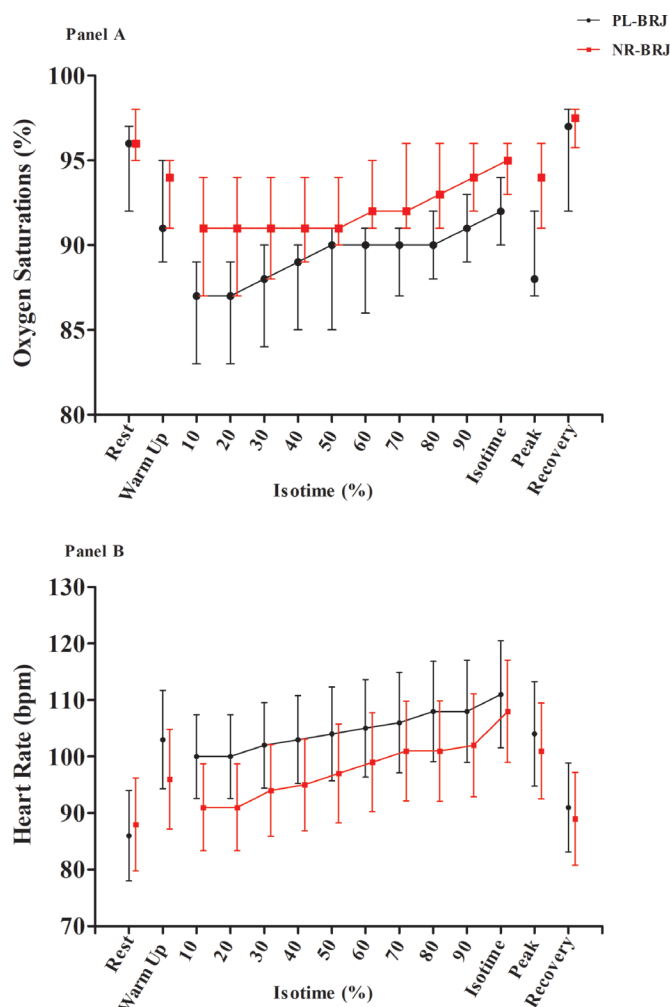


Figure 3 Effect of dietary nitrate supplementation on isotime oxygen saturation and heart rate during endurance shuttle walk test. (A) Oxygen saturation analysis in the NR-BRJ (red) and PL-BRJ (black) dosing conditions at the 10th percentile of isotime and at rest, warm-up, peak exercise and recovery. Data presented as median and IQR. The area under the curve for each treatment group was estimated and reported as mean (SD). Saturation for when the subjects were on PL-BRJ was 1161.85 (47.59) and when the subjects were on NR-BRJ was 1205.54 (46.39). The treatment effect was estimated to be 43.69 (29.09–58.28) ($p < 0.0001$). The results suggest that on average the area under the curve for saturations was higher when on NR-BRJ than when on PL-BRJ. These differences tended to show more during the isotime and peak periods. (B) Heart rate analysis in the NR-BRJ (red) and PL-BRJ (black) dosing conditions at the 10th percentile of isotime and at rest, warm-up, peak exercise and recovery. Data presented as median and IQR. The area under the curve for each treatment group was estimated and reported as mean (SD). The mean (SD) area under the curve for the heart rate data when the subjects were on PL-BRJ was 1299.93 (186.05) and for when the subjects were on NR-BRJ was 1258.76 (174.01). The estimated treatment effect was –41.17 (–116.74 to 34.40) ($p = 0.27$). The results show that while at individual time points the heart rate was higher for when the subjects were on PL-BRJ, there was no statistically significant difference in the area under the curve. bpm, beats per minute; NR-BRJ, nitrate-rich beetroot juice; PL-BRJ, placebo beetroot juice.

protein vs PL-BRJ 6.942 (5.768–8.026) $\mu\text{mol/g}$ protein ($p = 0.5$) (online supplemental appendix figure E3).

Paired measures of FE_{NO} were available for 16 participants at all seven time points (figure 7 and online supplemental appendix

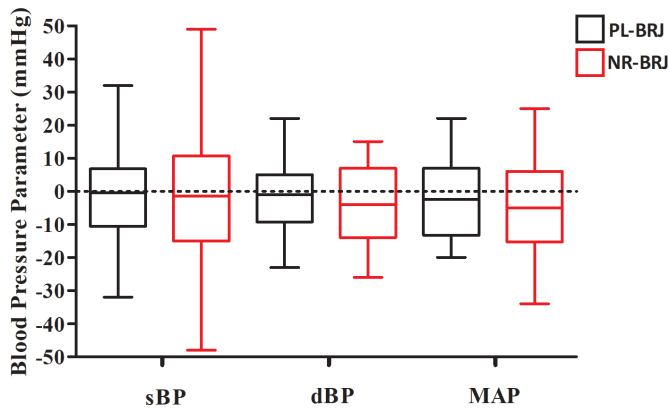


Figure 4 Effect of dietary nitrate supplementation on blood pressure parameters. Change in blood pressure parameters (sBP, dBP and MAP) relative to baseline blood pressure 3 hours prior to dosing with either NR-BRJ or PL-BRJ. Data presented as 25th–75th percentile with the solid line representing the median value and the whiskers the minimum to maximum values. Wilcoxon signed-rank test was used to compare blood pressure parameters. Median (IQR) change in sBP: NR-BRJ -1.5 (15.0 – 10.8) mm Hg vs PL-BRJ -0.5 (-10.5 to 6.8) mm Hg ($p=1.0$). Median (IQR) in dBP: NR-BRJ 4.0 (-14.0 to 7.0) mm Hg vs PL-BRJ -1.0 (-9.3 to 5.0) mm Hg ($p=0.481$). Median (IQR) change in MAP: NR-BRJ -5.0 (-15.3 to 6) mm Hg vs PL-BRJ -2.5 (-13.5 to 7) mm Hg ($p=0.359$). dBP, diastolic blood pressure; MAP, mean arterial pressure; NR-BRJ, nitrate-rich beetroot juice; PL-BRJ, placebo beetroot juice; sBP, systolic blood pressure.

table E4). For four participants, there was device failure resulting in no data being recorded. The median (IQR) AUC for when the subjects were on PL-BRJ was 3622.5 (3181.9–4796.9) and the corresponding result for when the subjects were on NR-BRJ was 9440.6 (6273.8–11 831.3), and the treatment effect with

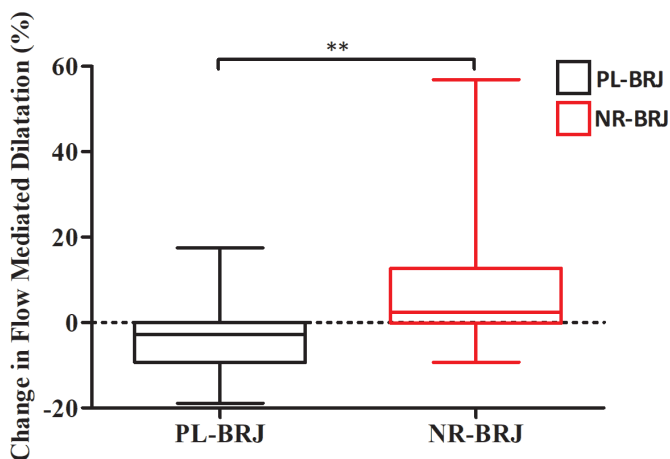


Figure 5 Effect of dietary nitrate supplementation on endothelial function. Percentage change in FMD from baseline and 180 min after supplementation with NR-BRJ or PL-BRJ. Data presented at the 25th and 75th percentile boxes with the solid line representing the median value and the whiskers the minimum and maximum values. Wilcoxon signed-rank test was used to compare the percentage change in FMD in the NR-BRJ (red) and PL-BRJ (black) dosing conditions. There was a statistically significant difference in the FMD percentage change with an increase in the NR-BRJ group (4.1 , -1.1 to 14.8) versus a reduction in the PL-BRJ group (-5.0 , -10.6 to -0.6). $**p=0.0003$. FMD, flow-mediated dilatation; NR-BRJ, nitrate-rich beetroot juice; PL-BRJ, placebo beetroot juice.

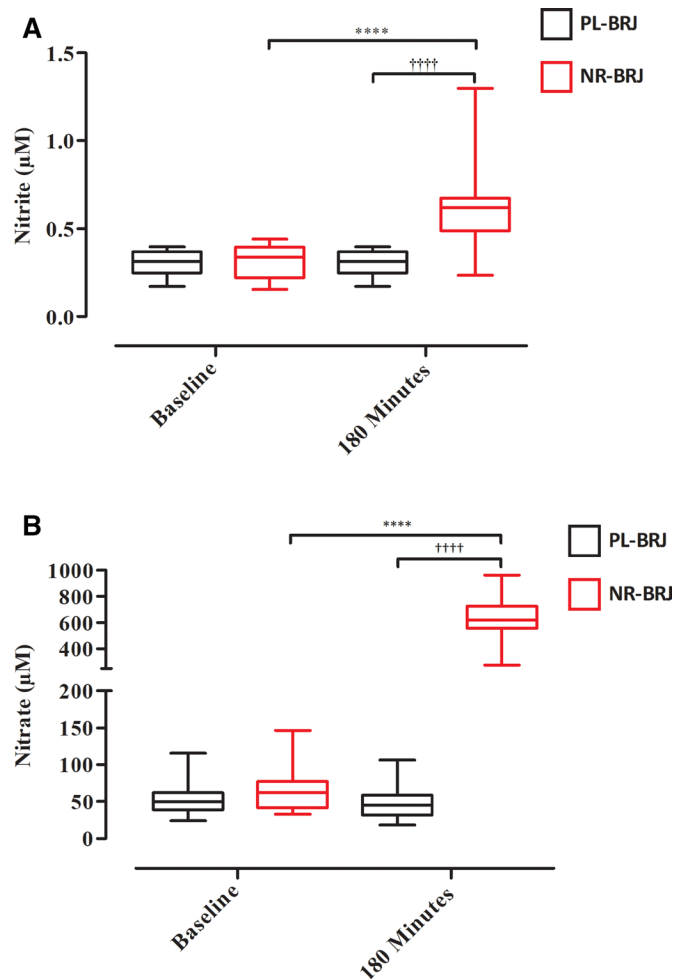


Figure 6 Plasma nitrite and nitrate levels. Data presented are median (IQR) with whiskers representing minimum to maximum values. Plasma NO_2^- and NO_3^- concentrations were measured at baseline (0 min) and 180 min after dosing with the interventions. Wilcoxon signed-rank test was used to compare change in plasma NO_2^- and NO_3^- concentrations between the intervention groups. Mann-Whitney U test was used to compare change in plasma NO_2^- and NO_3^- concentrations between the treatment conditions. (A) Changes in plasma NO_2^- concentrations. There was a statistically significant difference between baseline plasma NO_2^- concentration and postdosing with NR-BRJ for plasma NO_2^- : predosing plasma NO_2^- concentration 0.306 (0.227 – 0.402) μM vs postdosing 0.620 (0.488 – 0.673) μM ; $****p=0.000076$. There was also a statistically significant difference between postdosing plasma NO_2^- concentration between NR-BRJ and PL-BRJ dosing conditions: postdose of NR-BRJ NO_2^- concentration 0.620 (0.488 – 0.673) μM vs postdose of PL-BRJ NO_2^- concentration 0.306 (0.227 – 0.402) μM ; $++++p=0.000009$. (B) Changes in plasma NO_3^- levels. There was a statistically significant difference between baseline plasma NO_3^- concentration and postdosing with NR-BRJ for plasma NO_3^- : predosing plasma NO_3^- concentration 62.59 (41.68 – 77.29) μM vs postdosing 617 (556.25 – 725.88) μM ; $****p=0.00004$. There was also a statistically significant difference between postdosing plasma NO_3^- concentration between NR-BRJ and PL-BRJ dosing conditions: postdose NR-BRJ NO_3^- plasma concentration 617.71 (556.25 – 725.88) μM vs PL-BRJ plasma NO_3^- concentration 45.31 (31.39 – 58.84) μM ; $++++p=5.66 \times 10^{-11}$). NO_2^- , nitrite; NO_3^- , nitrate; NR-BRJ, nitrate-rich beetroot juice; PL-BRJ, placebo beetroot juice.

its 95% CI was 5407 (3096 to 7576) ($p=0.0011$). The results suggest that the FE_{NO} levels while the subjects were on NR-BRJ were significantly higher than when they were on PL-BRJ.

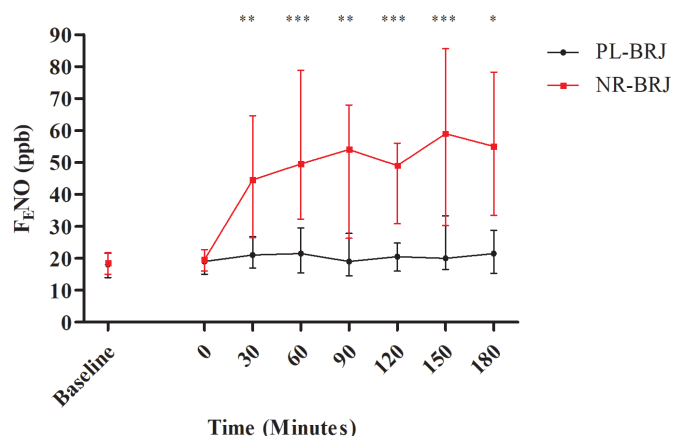


Figure 7 Exhaled nitric oxide. Data presented are median (dot) with whiskers representing IQR and with red dot and line representing NR-BRJ and black dot and line representing PL-BRJ. FE_{NO} was measured at baseline (study visit 1) and subsequently at intervention visits at seven time points (0, 30, 60, 90, 120, 150 and 180 min) dosing with either NR-BRJ or PL-BRJ. Kruskal-Wallis H test was used to assess the effect of either NR-BRJ or PL-BRJ on FE_{NO} . In both intervention groups there was no statistical difference between FE_{NO} at baseline (measured at study visit 1) and time point 0 min (measured at intervention visits prior to supplementation with intervention beverage). There was a statistically significant difference between measured FE_{NO} at all subsequent time points postintervention consumption. 30 min $p=0.0011$, 60 min $p=0.0001$, 90 min $p=0.0006$, 120 min $p=0.0002$, 150 min $p=0.0002$, 180 min $p=0.0024$; ** $p\leq 0.01$, *** $p\leq 0.001$ (see online supplemental appendix Table E4). NR-BRJ, nitrate-rich beetroot juice; PL-BRJ, placebo beetroot juice; ppb, parts per billion.

Postacute (0 min) supplementation with NR-BRJ FE_{NO} increased by 184% at 180 min post supplementation.

DISCUSSION

The major finding of this study was that, in people with COPD who are hypoxic to the extent that they meet the criteria for LTOT, dietary NO_3^- supplementation improves exercise capacity compared with placebo. The improvement in ESWT that we observed was accompanied by less desaturation during exercise. Supplementation also improved endothelial function assessed using FMD. In line with previous observations,^{24 25} blood pressure was numerically lower as well but the difference in response was not statistically significant.

Significance of findings

Although studies have previously considered dietary NO_3^- supplementation in COPD with inconsistent results,^{24–26 40} this is the first stratified medicine approach focusing on the specific phenotype of individuals with COPD with hypoxaemia requiring LTOT. Our previous study, in non-hypoxaemic patients with COPD, found that there was a reduction in the oxygen cost of exercise during cycle ergometry yet no improvement in exercise capacity.²⁶ In conditions of hypoxia, the L-arginine–nitric oxide synthase pathway is compromised, while the NO_3^- – NO_2^- –NO pathway is facilitated due to a lesser inhibition of NO_2^- bioactivation by oxygen.^{14 20 22} As such, dietary NO_3^- supplementation could be expected to have more impact in hypoxic rather than normoxic individuals, both through effects on the skeletal muscle and impacts on the pulmonary vasculature.

The mechanism by which ESWT lengthened is likely to involve multiple synergistic pathways. The finding of relatively preserved

SpO_2 during exercise in the NO_3^- -supplemented condition could reflect more efficient oxygen utilisation peripherally, a beneficial impact on central haemodynamics associated with/reduced hypoxia-induced pulmonary vasoconstriction, or a combination of the two. Despite each participant using their prescribed oxygen for each walk test, there was an observed desaturation in the placebo arm. This could mean that their oxygen prescription may have been insufficient and that a higher flow rate might also have increased exercise capacity. This finding of the attenuation of desaturation by NR-BRJ may well be explained by the enhancement of the NO_3^- – NO_2^- –NO pathway in conditions of hypoxia.

The observation that NO_3^- supplementation was associated with improvements in endothelial function assessed using FMD is likely to be relevant to the acute mechanism of benefit from NO_3^- supplementation, but also raises the possibility that longer-term dosing might reduce the risk of vascular events which are common in COPD. The effects seen are almost certainly not COPD-specific and work is needed to investigate possible benefits in other long-term lung conditions associated with hypoxia, including interstitial lung diseases and the various categories of pulmonary hypertension.

The estimated treatment effect of dietary NO_3^- supplementation on ESWT found in this study was 62.5 s, which falls fractionally short of the MCID defined in pharmacotherapy trials as 65 s.³⁰ However, in pharmacological trials where the ESWT is the outcome, interventions are typically administered over weeks or months. The demonstration of an effect of similar magnitude in a single-dose study is therefore encouraging, although further studies of longer-term use will be needed before any clinical recommendations can be made.

Study limitations

The use of a robust placebo strengthens the reliability of the findings, as does the fact that the improvement in walking time was accompanied by an appropriate physiological response (lower HR and higher SpO_2). An additional strength was the use of a walking rather than a cycling test, which is of clinical relevance to patients as it reflects most individuals' main form of exercise and daily physical activity. This was a single-dose study and therefore questions remain as to the impact that regular dosing might have and whether this would translate into meaningful clinical effects. The dose used was selected based on previous studies, but future work should investigate whether there is a dose response or ceiling effect. We have also shown that the NR-BRJ does indeed contain a higher quantity of NO_3^- and provide independent confirmation that NO_3^- is only present at very low levels in the placebo juice used in our study.

CONCLUSION

BRJ is cheap and readily accessible and has the potential to be used widely as a dietary supplement if effective in specific patient groups. Its beneficial effects appear to be mediated by inorganic NO_3^- without affecting plasma redox status, on acute administration. Further mechanistic work is needed to work out the relative impact of the possible mechanisms, in particular the impact of muscle versus pulmonary or cardiac/systemic circulation effects, and longer-term studies will be needed to establish if the effects on exercise performance and endothelial function observed here translate into clinically meaningful benefits.

Twitter Matthew J Pavitt @DrMattPav, Adam Lewis @apl104 and Nicholas S Hopkinson @COPDdoc

Acknowledgements The authors would like to thank all the participants who took part in this study.

Contributors NSH and MJP developed the original idea for the research study. NSH and MJP designed and wrote the study protocol. WASB designed the statistical

analysis plan. MJP, AL and SCB undertook patient visits and collected trial data. MF, BOF and MM-L undertook plasma analysis. MJP analysed the data and wrote the first draft of the manuscript. All authors edited and contributed to the final manuscript. NSH is the guarantor.

Funding The study was funded by a grant from Moulton Charitable Foundation.

Competing interests None declared.

Patient consent for publication Not required.

Ethics approval The study was approved by the London - Chelsea Research and Ethics Committee (ref: 15/LO/0975) and conducted in line with the principles of the Declaration of Helsinki.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available upon reasonable request. Individual participant data that underlie the results in the article after de-identification (text, tables, figures and appendices) will be made available from the corresponding author upon request. The study protocol and statistical analysis plan will also be available. Data will be available indefinitely.

ORCID iDs

Adam Lewis <http://orcid.org/0000-0002-0576-8823>

Nicholas S Hopkinson <http://orcid.org/0000-0003-3235-0454>

REFERENCES

- Hopkinson NS, Dayer MJ, Moxham J, *et al.* Abdominal muscle fatigue following exercise in chronic obstructive pulmonary disease. *Respir Res* 2010;11:15.
- Hopkinson NS, Sharshar T, Ross ET, *et al.* Corticospinal control of respiratory muscles in chronic obstructive pulmonary disease. *Respir Physiol Neurobiol* 2004;141:1–12.
- Jackson AS, Shrikrishna D, Kelly JL, *et al.* Vitamin D and skeletal muscle strength and endurance in COPD. *Eur Respir J* 2013;41:309–16.
- Maddocks M, Shrikrishna D, Vitoriano S, *et al.* Skeletal muscle adiposity is associated with physical activity, exercise capacity and fibre shift in COPD. *Eur Respir J* 2014;44:1188–98.
- Nataneek SA, Gosker HR, Slot IGM, *et al.* Heterogeneity of quadriceps muscle phenotype in chronic obstructive pulmonary disease (COPD); implications for stratified medicine? *Muscle Nerve* 2013;48:488–97.
- Sadaka AS, Montgomery AJ, Mourad SM, *et al.* Exercise response to oxygen supplementation is not associated with survival in hypoxemic patients with obstructive lung disease. *Int J Chron Obstruct Pulmon Dis* 2018;13:1607–12.
- Brown GC, Cooper CE. Nanomolar concentrations of nitric oxide reversibly inhibit synaptosomal respiration by competing with oxygen at cytochrome oxidase. *FEBS Lett* 1994;356:295–8.
- Umbrello M, Dyson A, Feelisch M, *et al.* The key role of nitric oxide in hypoxia: hypoxic vasodilation and energy supply-demand matching. *Antioxid Redox Signal* 2013;19:1690–710.
- Merry TL, Lynch GS, McConnell GK. Downstream mechanisms of nitric oxide-mediated skeletal muscle glucose uptake during contraction. *Am J Physiol Regul Integr Comp Physiol* 2010;299:R1656–65.
- Stamler JS, Meissner G. Physiology of nitric oxide in skeletal muscle. *Physiol Rev* 2001;81:209–37.
- Joyner MJ, Tschakovsky ME. Nitric oxide and physiologic vasodilation in human limbs: where do we go from here? *Can J Appl Physiol* 2003;28:475–90.
- Garthwaite J. Concepts of neural nitric oxide-mediated transmission. *Eur J Neurosci* 2008;27:2783–802.
- Percival JM, Anderson KNE, Huang P, *et al.* Golgi and sarcolemmal neuronal NOS differentially regulate contraction-induced fatigue and vasoconstriction in exercising mouse skeletal muscle. *J Clin Invest* 2010;120:816–26.
- Feelisch M, Fernandez BO, Bryan NS, *et al.* Tissue processing of nitrite in hypoxia: an intricate interplay of nitric oxide-generating and -scavenging systems. *J Biol Chem* 2008;283:33927–34.
- Lundberg JO, Weitzberg E. No generation from inorganic nitrate and nitrite: role in physiology, nutrition and therapeutics. *Arch Pharm Res* 2009;32:1119–26.
- Bailey SJ, Fulford J, Vanhatalo A, *et al.* Dietary nitrate supplementation enhances muscle contractile efficiency during knee-extensor exercise in humans. *J Appl Physiol* 2010;109:135–48.
- Larsen FJ, Weitzberg E, Lundberg JO, *et al.* Effects of dietary nitrate on oxygen cost during exercise. *Acta Physiol* 2007;191:59–66.
- Carriker CR, Mermier CM, Van Dusseldorp TA, *et al.* Effect of acute dietary nitrate consumption on oxygen consumption during submaximal exercise in hypobaric hypoxia. *Int J Sport Nutr Exerc Metab* 2016;26:315–22.
- Kelly J, Vanhatalo A, Bailey SJ, *et al.* Dietary nitrate supplementation: effects on plasma nitrite and pulmonary O₂ uptake dynamics during exercise in hypoxia and normoxia. *Am J Physiol Regul Integr Comp Physiol* 2014;307:R920–30.
- Masschelein E, Van Thienen R, Wang X, *et al.* Dietary nitrate improves muscle but not cerebral oxygenation status during exercise in hypoxia. *J Appl Physiol* 2012;113:736–45.
- Shannon OM, Duckworth L, Barlow MJ, *et al.* Dietary nitrate supplementation enhances high-intensity running performance in moderate normobaric hypoxia, independent of aerobic fitness. *Nitric Oxide* 2016;59:63–70.
- Vanhatalo A, Fulford J, Bailey SJ, *et al.* Dietary nitrate reduces muscle metabolic perturbation and improves exercise tolerance in hypoxia. *J Physiol* 2011;589:5517–28.
- Vanhatalo A, Jones AM, Blackwell JR, *et al.* Dietary nitrate accelerates postexercise muscle metabolic recovery and O₂ delivery in hypoxia. *J Appl Physiol* 2014;117:1460–70.
- Pavitt MJ, Tanner RJ, Lewis A, *et al.* Oral nitrate supplementation to enhance pulmonary rehabilitation in COPD: ON-EPIC a multicentre, double-blind, placebo-controlled, randomised parallel group study. *Thorax* 2020;75:547–55.
- Alsulayyim AS, Alasmari AM, Alghamdi SM, *et al.* Impact of dietary nitrate supplementation on exercise capacity and cardiovascular parameters in chronic respiratory disease: a systematic review and meta-analysis. *BMJ Open Respir Res* 2021;8:e000948.
- Curtis KJ, O'Brien KA, Tanner RJ, *et al.* Acute dietary nitrate supplementation and exercise performance in COPD: a double-blind, placebo-controlled, randomised controlled pilot study. *PLoS One* 2015;10:e0144504.
- Vogelmeier CF, Criner GJ, Martinez FJ, *et al.* Global strategy for the diagnosis, management, and prevention of chronic obstructive lung disease 2017 report. gold executive summary. *Am J Respir Crit Care Med* 2017;195:557–82.
- Hopkinson NS, Molyneux A, Pink J, *et al.* Chronic obstructive pulmonary disease: diagnosis and management: summary of updated NICE guidance. *BMJ* 2019;366:l4486.
- Lansley KE, Winyard PG, Fulford J, *et al.* Dietary nitrate supplementation reduces the O₂ cost of walking and running: a placebo-controlled study. *J Appl Physiol* 2011;110:591–600.
- Singh SJ, Puhan MA, Andrianopoulos V, *et al.* An official systematic review of the European respiratory Society/American thoracic Society: measurement properties of field walking tests in chronic respiratory disease. *Eur Respir J* 2014;44:1447–78.
- Webb AJ, Patel N, Loukogeorgakis S, *et al.* Acute blood pressure lowering, vasoprotective, and antiplatelet properties of dietary nitrate via bioconversion to nitrite. *Hypertension* 2008;51:784–90.
- Warren GL, Park ND, Maresca RD, *et al.* Effect of caffeine ingestion on muscular strength and endurance: a meta-analysis. *Med Sci Sports Exerc* 2010;42:1375–87.
- Cumpstey AF, Hennis PJ, Gilbert-Kawai ET, *et al.* Effects of dietary nitrate on respiratory physiology at high altitude - Results from the Xtreme Alps study. *Nitric Oxide* 2017;71:57–68.
- Kleinbongard P, Dejam A, Lauer T, *et al.* Plasma nitrite reflects constitutive nitric oxide synthase activity in mammals. *Free Radic Biol Med* 2003;35:790–6.
- Benzie IF, Strain JJ. The ferric reducing ability of plasma (FRAP) as a measure of "antioxidant power": the FRAP assay. *Anal Biochem* 1996;239:70–6.
- Kasielski M, Nowak D. Long-Term administration of N-acetylcysteine decreases hydrogen peroxide exhalation in subjects with chronic obstructive pulmonary disease. *Respir Med* 2001;95:448–56.
- Koning AM, Meijers WC, Pasch A, *et al.* Serum free thiols in chronic heart failure. *Pharmacol Res* 2016;111:452–8.
- Thijssen DHJ, Black MA, Pyke KE, *et al.* Assessment of flow-mediated dilation in humans: a methodological and physiological guideline. *Am J Physiol Heart Circ Physiol* 2011;300:H2–12.
- Rodriguez-Miguelez P, Seigler N, Harris RA. Ultrasound assessment of endothelial function: a technical guideline of the flow-mediated dilation test. *J Vis Exp* 2016. doi:10.3791/54011. [Epub ahead of print: 27 04 2016].
- Beijers RJHCG, Huysmans SMD, van de Boel C, *et al.* The effect of acute and 7-days dietary nitrate on mechanical efficiency, exercise performance and cardiac biomarkers in patients with chronic obstructive pulmonary disease. *Clin Nutr* 2018;37:1852–61.

SUPPLEMENTARY APPENDIX

Dietary nitrate supplementation to improve exercise capacity in hypoxic COPD

Matthew J. Pavitt¹, Adam Lewis¹, Sara C. Buttery², Bernadette O. Fernandez³, Monika Mikus-Lelinska³, Winston Banya¹, Martin Feelisch³, Michael I. Polkey¹, Nicholas S. Hopkinson¹

¹National Heart and Lung Institute, Imperial College, London, Royal Brompton Campus

²South London Healthcare NHS Trust

³Faculty of Medicine, Clinical & Experimental Sciences, University of Southampton and Southampton NIHR Respiratory Biomedical Research Unit, Southampton Hospital, Southampton, UK

Corresponding Author

Name: Professor Nicholas S. Hopkinson

Address: NHLI, Imperial College, Royal Brompton Hospital Campus, Fulham Road, London, SW3 6NP

Email Address: n.hopkinson@ic.ac.uk

Telephone: 0207 349 7775

Twitter: @COPDdoc

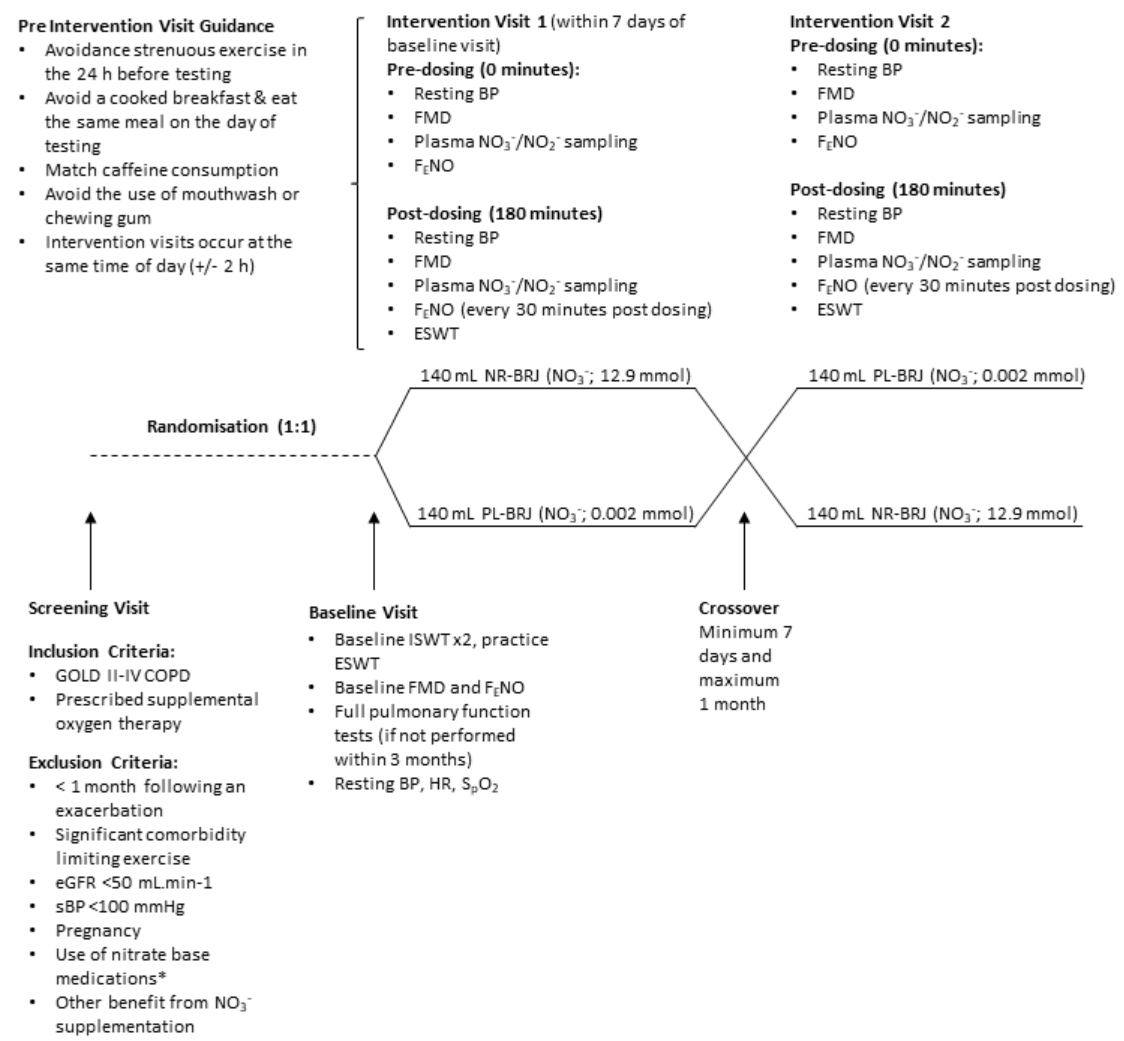


Figure E1. Study flow diagram

Abbreviations: NO_3^- - Nitrate; BP - Blood Pressure; FMD - Flow Mediated Dilatation; NO_2^- - Nitrite; F_tNO - Fractional Exhaled Nitric Oxide; ESWT - Endurance Shuttle Walk Test; BRJ - Beetroot Juice; PL - Placebo; GOLD - Global Initiative for Chronic Obstructive Lung Disease; COPD - Chronic Obstructive Pulmonary Disease; eGFR - Estimated Glomerular Filtration Rate; sBP - Systolic Blood Pressure; ISWT - Incremental Shuttle Walk Test; HR - Heart Rate; S_pO_2 - Oxygen Saturations

SUPPLEMENTARY METHODS

Plasma nitrate/nitrite levels - additional methods

Plasma NO_3^- and NO_2^- levels were used as a combined biomarker of NO_3^- ingestion, metabolism and nitric oxide availability [1, 2]. Plasma samples were obtained on arrival and three hours after consumption of either NR-BRJ or PL-BRJ. Samples were obtained by venesection of 6 mL of venous blood into lithium heparin tubes. Within five minutes of collection the vials were split into 3 mL aliquots, with one mixed with 300 μL of 100 mM stock of N-ethylmaleimide (NEM) solution (final concentration 10 mM). The samples were then centrifuged at 1,000 g for eight minutes at room temperature. Subsequently, 1 mL of the supernatant was aliquoted into 2 mL polypropylene cryotubes, snap frozen with liquid nitrogen and stored at -80°C . Plasma nitrate and nitrite concentrations were measured following protein precipitation with methanol (1:1 v/v) by a dedicated high-performance liquid chromatography (HPLC) system equipped with an anion-exchange column, an in-line Cd/Cu reduction column and a post-column diazo coupling reactor coil (Griess reaction) (Eicom NOx analyser, ENO-20, San Diego, USA) [3].

Oxidative stress – additional methods

Ferric-reducing ability of plasma (FRAP)

The FRAP assay is a measure of the antioxidant potential in the extracellular compartment [4]. The same plasma samples used for nitrate/nitrite measurement were also used to process this assay. Briefly, 150 μL of FRAP reagent (containing 300 mM acetate buffer at pH 3.6, 10 mM TPTZ [2,4,6-Tris(2-pyridyl)s-triazine], 20 mM FeCl_3 at a ratio of 10:1:1 (v:v:v)) was added to 5 μL of diluted plasma (1:3, v:v) into a 96-well plate containing 15 μL of MQ water in each well. The plate was incubated at 37°C for 30 minutes. The absorbance at 593 nm was taken immediately after incubation using a microplate reader (Spectramax M5, Molecular Devices, California USA). FRAP values for the samples were obtained by comparing the absorbance at 593 nm with the known concentrations in the standards ($\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$).

Thiobarbituric acid-reactive substance (TBARS)

TBARS is a measure of lipid oxidation and is measured using a TBARS assays [5]. The same plasma samples used for nitrate/nitrite measurement were also used to process this assay. In

brief the TBARS assay incorporated the use of an malodialdehyde (MDA) source such as 1,1,3,3 Tetramethoxypropane after hydrolysis as standard, 0.6N trichloroacetic acid as the acid reagent and thiobarbituric acid (0.26g in 50 mL glacial acetic acid) as colour reagent. Prior to analysis, samples were deproteinised by acid precipitation by taking 300 uL samples and adding an equal volume of acid reagent, mixed and incubated for 15 min at room temperature. The supernatant was isolated by 4 min centrifugation at $> 12,000 \times g$. The resulting supernatant was further treated with colour reagent (2:1, v:v), incubated for 1h at 100°C and immediately cooled on ice for 10 min. Treated samples were plated into 96-well microplates and absorbance readings were read at 532 nm using a microplate reader (Spectramax M5, Molecular Devices, California USA). TBARS values for the samples were obtained by comparing the absorbance with known concentrations of MDA standards.

Total free thiols per protein

Systemic oxidative stress can be measured as the depletion of the free thiol pool in plasma [6]. The same plasma samples used for nitrate/nitrite measurement were also used to process this assay. Thiol groups were measured as previously described [7, 8]. In brief, 75 µl plasma samples were diluted 1:4 (v:v) with a 0.1 M Tris buffer (pH 8.2) and transferred 90 uL of diluted sample to a 96-well microplate. Using a microplate reader (Molecular Devices Spectramax M5, California, USA), background absorption was measured at 412 nm with a reading at 630 nm for baseline correction. Subsequently, 20 µl 1.9 mM 5,5-Dithio-bis(2-nitrobenzoic acid) [DTNB] in 0.1 M phosphate buffer (pH 7) was added to the samples and standards. Following 20 minutes of incubation at room temperature while mixing, absorption was remeasured at 412 and 630 nm. The concentration of total free thiols in the samples was determined by comparing their absorbance reading to that of an L-cysteine standard before and after addition of DTNB to samples/standards.

Endothelial function - additional methods

Endothelial function was assessed using flow mediated dilatation (FMD) of the brachial artery [9] using a high-resolution doppler ultrasound (GE Logiq 3, GE Medical Systems, Milwaukee, Wisconsin, USA) and a 10 MHz multi-frequency linear array probe were used in B-mode. Brachial artery diameter was measured at baseline and sequentially after release of circulatory arrest of the upper arm over a period of 120 seconds [10], three hours after NR-BRJ/PL-BRJ

consumption. All measurements were performed by a single trained operator. Circulatory arrest was generated via a rapid cuff inflation system (Hokanson, Bellevue, WA, USA), which was positioned proximal to the brachial artery and rapidly inflated to 250 mmHg for five minutes. Data were saved for off-line analysis using ImageJ2 software [11].

SUPPLEMENTARY RESULTS

Table E1. Exercise oxygen saturations and heart rate analysis

Measure	PL-BRJ (n=18)	NR-BRJ (n=18)
Saturations (%)		
Rest	96 (90, 97)	96 (92, 97)
Warm-up	91 (89, 95)	94 (90, 95)
Isotime	92 (89, 94)	96 (93, 97)
Peak	88 (86, 92)	94 (91, 96)
Recovery	97 (92, 98)	98 (96, 98)
Heart Rate (bpm)		
Rest	86 (74, 88)	88 (78, 91)
Warm-up	103 (88, 108)	96 (88, 102)
Isotime	111 (103, 123)	109 (96, 116)
Peak	104 (96, 111)	101 (112)
Recovery	91 (79, 101)	89 (81, 98)

The area under the curve for each treatment group was estimated and reported as mean (SD). The results for Saturations for when the subjects were on placebo beetroot juice were 1161.85 (47.59) and the results for when the subjects on Nitrate-rich beetroot juice were 1205.54 (46.39). The treatment effect was estimated to be 43.69 (29.09 to 58.28) $p < 0.0001$. The results suggest that on the average the area under the curve for saturations was higher when on Nitrate-rich beetroot juice than when on placebo. These differences tended to show more during the Isotime and peak periods.

The mean (SD) area under the curve for the HR data when the subjects were on placebo beetroot juice was 1299.93 (186.05) for when the subjects were on Nitrate-rich beetroot juice results was 1258.76 (174.01). The estimated treatment effect was -41.17 (-116.74 to 34.40), $p=0.27$. The results show that while at individual time points the HR was higher for when the subjects were on Placebo, there was no statistically significant difference in the area under the curve.

Abbreviations; bpm – Beats Per Minute; PL-BRJ – Placebo Beetroot Juice; Nitrate-rich Beetroot Juice

Table E2. Between intervention analysis of plasma nitrite and nitrate

Measurement	PL-BRJ (n=19)	NR-BRJ (n=19)
Baseline Nitrite (µM)	0.32 (0.25, 0.37)	0.34 (0.22, 0.39)
Baseline Nitrate (µM)	51.89 (38.98, 62.28)	62.59 (41.68, 77.29)
180 Minute Nitrite (µM)	0.31 (0.23, 0.47)	0.60 (0.48, 0.67)
180 Minute Nitrate (µM)	45.31 (31.39, 58.84)	617.71 (508.6, 725.88)
Difference in Nitrite (µM) from baseline to 180 minutes	0.023 (-0.044, 0.079)	0.276 (0.144, 0.463)
Difference in Nitrate (µM) from baseline to 180 minutes	-4.61 (-9.63, 6.23)	543.25 (441.78, 674.23)

Results are reported as median (IQR).

The treatment effect of Nitrate was estimated with the Hodges-Lehman estimate and it was 550 (461 to 639) µM. The results suggest that there was a higher for when the subjects were on Nitrate-rich beetroot juice than when they were on placebo and this change was statistically significant, $p=0.0003$.

The treatment effect of Nitrite was estimated with the Hodges-Lehman estimate and it was 0.248 (0.138 to 0.408) µM. The results suggest that there was a higher for when the subjects were on Nitrate-rich beetroot juice than when they were on placebo and this change was statistically significant, $p=0.0011$.

Abbreviations: PL BRJ – Placebo Beetroot Juice; NR-BRJ – Nitrate-rich Beetroot Juice

Table E3. Nitrate levels in active and placebo juice and analytic materials

Samples	Nitrite				Nitrate			
	Mean (μM)	SD	SEM	% CV	Mean (μM)	SD	SEM	% CV
NEM Stock Solution	0.07	0.03	0.02	45.41	22.37	2.33	1.34	10.40
Cryotube	0.09	0.02	0.01	17.15	2.44	0.53	0.31	22.33
PL-BRJ	195.86	2.12	1.22	1.08	55.05	0.68	0.39	1.24
NR-BRJ	10.75	0.20	0.11	1.83	120411.03	5267.10	3040.96	4.37

Concentration of NO_2^- and NO_3^- in NEM Stock Solution, Cryotubes, Cryotubes with 0.9% Sodium Chloride, PL-BRJ and NR-BRJ.

Abbreviations: NEM – N-Ethylmaleimide; SD – Standard Deviation; SEM – Standard Error Mean; %CV – Percentage Coefficient of Variation; PL-BRJ – Placebo Beetroot Juice; NR-BRJ – Nitrate Rich Beetroot Juice; μM – micromole.

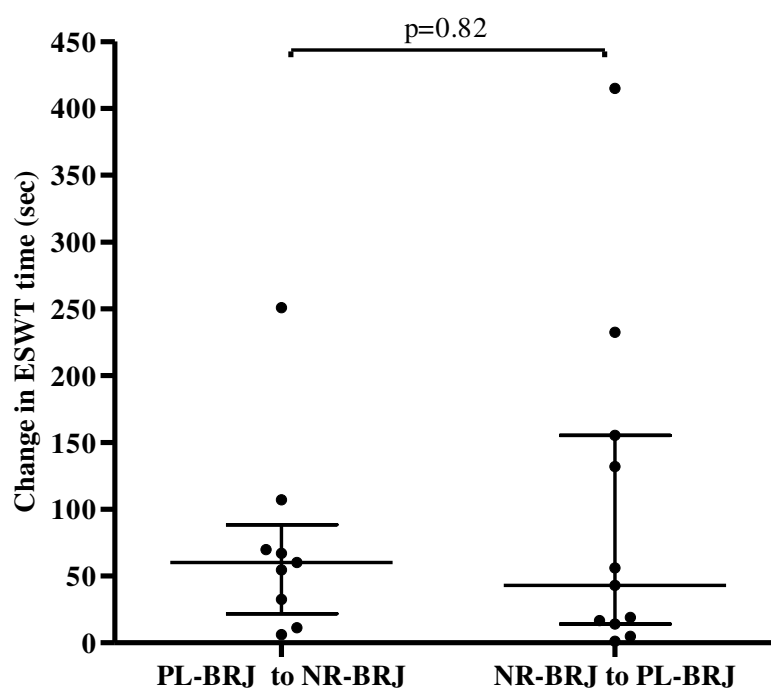
Table E4. Fractional Exhaled Nitric Oxide (FeNO)

	F_ENO post NR-BRJ (ppb)	FeNo post PL-BRJ (ppb)
Baseline	18.5 (15.0, 21.5)	18.0 (14.0, 22.5)
0 Minutes	19.5 (16.0, 22.5)	19.0 (15.0, 22.5)
30 Minutes	44.5 (27.0, 63.0)	21.0 (17.0, 25.5)
60 Minutes	49.5 (33.5, 78.5)	21.5 (17.0, 29.0)
90 Minutes	54.0 (26.5, 90.0)	19.0 (15.0, 27.5)
120 Minutes	49.0 (32.5, 56.0)	20.5 (16.0, 24.5)
150 Minutes	59.0 (33.5, 84.0)	20.0 (17.0, 32.5)
180 Minutes	55.0 (35.0, 76.5)	21.5 (15.5, 27.5)

F_ENO levels measured at baseline (visit 1) and subsequently at intervention visits (visits 3 and 4) at time point zero minutes prior to dosing with either PL-BRJ or NR-BRJ and subsequently every 30 minutes until 180 minutes post dosing. Data presented: median (IQR).

The AUC was calculated for each treatment group and compared to estimate the treatment effect using the Hodges-Lehman estimate. The median (IQR) AUC for when the subjects were on placebo was 3622.5 (3181.9, 4796.9) and the corresponding results for when the subjects were on Nitrate-rich beetroot juice was 9440.6 (6273.8, 11831.3) and the treatment effect with its 95% CI was 5407 (3096 to 7576), $p=0.0011$. The results suggest that the FeNO levels while the subjects were on Nitrate-rich beetroot juice were significantly higher than when they were on placebo

Abbreviations: F_ENO – Fractional Exhaled nitric Oxide; IQR – Interquartile Range; AUC – area under the curve; ppb – Parts Per Billion

Figure E2. Primary Outcome Order Effect

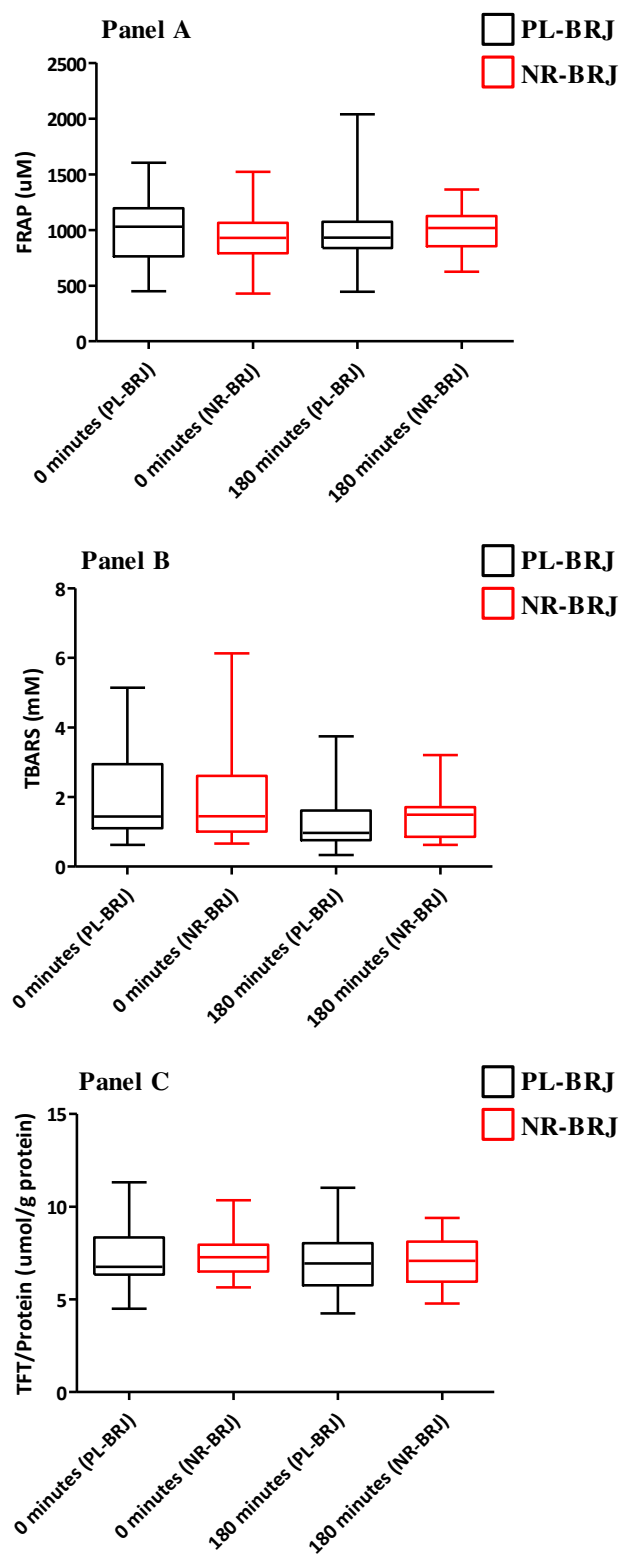
Change in ESWT time (seconds) when testing for intervention order effect, if PL-BRJ was applied first or NR-BRJ. Data presented median (line) and interquartile range (whiskers) with as individual data points (dots). Mann-Whitney U test, the median (IQR) change in ESWT time if PL-BRJ was applied first was 60.0 (21.8, 88.4) seconds, compared to 43.1 (14.03, 155.3) seconds, if NR-BRJ was applied first; $p = 0.82$.

Abbreviations: ESWT – Endurance Shuttle Walk Test; PL-BRJ – Placebo Beetroot Juice; NR-BRJ – Nitrate Rich Beetroot Juice

Results E1. Effect of dietary nitrate supplementation on endurance shuttle walk time

There is a clear outlier in this dataset. When this individual's data was removed from analysis, all individuals still walked further following consumption the NO₃⁻-rich BRJ. There was a statistically significant difference between the median (IQR) ESWT time with the outlier removed; NO₃⁻-rich BRJ 193.8 (145.5, 389.6) seconds vs PL 158.2 (121.6, 236.6) seconds; $p = 0.0001$. Regarding this specific individual at baseline assessment their best ISWT distance was 370 meters, using the ESWT conversion table the ESWT speed was calculated as 4.65 km/h which equates to ESWT level 11. All individuals undertook a practice ESWT, this individual's practice ESWT time was 599 seconds. The ESWT time that this individual achieved following consumption of the placebo beverage was 785 seconds. For both ESWT this individual reported peak Borg Dyspnoea scale of 8. This individual's data was included in the full analysis as it is felt to be a true representation of this individuals exercise endurance.

Figure E3. Measures of oxidative stress



Measures of oxidative stress for PL-BRJ and NR-BRJ Dosing conditions. Data presented as Data presented are median and interquartile range (box) with whiskers representing minimum to maximum values. Plasma samples were measured at baseline (zero minutes) and 180 minutes after dosing. Wilcoxon sign-rank test was used to compare change in measures of oxidative stress between intervention groups. Mann-Whitney U test was used to compare change in measures of oxidative stress between treatment conditions.

Panel A. Ferric reducing ability of plasma (FRAP)

There was no statistically significant difference between interventions for baseline and post intervention FRAP. Baseline FRAP PL-BRJ: 1028 (762.9, 1195) μM vs FRAP NR-BRJ: 927.7 (790.2, 1064) μM ; $p = 0.7$. Post intervention FRAP PL-BRJ: 930.2 (836.8, 1073) μM vs FRAP NR-BRJ: 1018 (853.0, 1125) μM ; $p = 1.0$. (Wilcoxon sign-rank test)

There was no statistically significant difference between baseline FRAP levels and post dosing levels with either PL-BRJ and NR-BRJ. FRAP PL-BRJ: 1028 (762.9, 1195) μM vs post dosing 930.2 (836.8, 1073) μM ; $p = 0.9$. FRAP NR-BRJ: 927.7 (790.2, 1064) μM vs post dosing 1018 (853.0, 1125) μM ; $p = 0.3$. (Mann-Whitney U test).

Panel B. Thiobarbituric acid-reactive substance (TBARS)

There was no statistically significant difference between interventions for baseline and post intervention TBARS. Baseline TBARS PL-BRJ: 1.443 (1.102, 2.940) mM vs TBARS NR-BRJ: 1.450 (1.007, 2.613) mM; $p = 0.8$. Post intervention TBARS PL-BRJ: 0.971 (0.766, 1.614) mM vs TBARS NR-BRJ: 1.499 (0.855, 3.209) mM; $p = 0.4$. (Wilcoxon sign-rank test)

There was no statistically significant difference between baseline TBARS levels and post dosing levels with either PL-BRJ and NR-BRJ. TBARS PL-BRJ: 1.443 (1.102, 2.940) mM vs post dosing 0.971 (0.766, 1.614) mM; $p = 0.8$. TBARS NR-BRJ: 1.450 (1.007, 2.613) mM vs post dosing 1.499 (0.855, 3.209) mM; $p = 0.3$. (Mann-Whitney U test).

Panel C. Total free thiols (TFT) per protein

There was no statistically significant difference between interventions for baseline and post intervention TFT per protein. Baseline TFT per protein PL-BRJ: 6.754 (6.328, 8.342) $\mu\text{mol.g}^{-1}$ protein vs TFT per protein NR-BRJ: 7.284 (6.508, 7.960) $\mu\text{mol.g}^{-1}$ protein; $p = 0.9$. Post intervention TFT per protein PL-BRJ: 6.942 (5.768, 8.026) $\mu\text{mol.g}^{-1}$ protein vs TFT per protein NR-BRJ: 7.079 (5.961, 8.115) $\mu\text{mol.g}^{-1}$ protein; $p = 0.5$. (Wilcoxon sign-rank test)

There was no statistically significant difference between baseline TFT per protein levels and post dosing levels with either PL-BRJ and NR-BRJ. TFT per protein PL-BRJ: 6.754 (6.328, 8.342) $\mu\text{mol.g}^{-1}$ protein vs post dosing 6.942 (5.768, 8.026) $\mu\text{mol.g}^{-1}$ protein; $p = 0.1$. TFT per protein NR-BRJ: 7.284 (6.508, 7.960) $\mu\text{mol.g}^{-1}$ vs post dosing 7.079 (5.961, 8.115) $\mu\text{mol.g}^{-1}$ protein; $p = 0.4$. (Mann-Whitney U test).

Abbreviations: FRAP - Ferric Reducing Ability of Plasma; TBARS - Thiobarbituric Acid-Reactive Substance; TFT – Total Free Thiols; PL-BRJ – Placebo Beetroot Juice; NR-BRJ – Nitrate-rich Beetroot Juice; mM - millimole

REFERENCES

1. Cumpstey AF, Hennis PJ, Gilbert-Kawai ET, Fernandez BO, Poudevigne M, Cobb A, Meale P, Mitchell K, Moyses H, Pöhl H, Mythen MG, Grocott MPW, Feelisch M, Martin DS. Effects of dietary nitrate on respiratory physiology at high altitude - Results from the Xtreme Alps study. *Nitric oxide : biology and chemistry / official journal of the Nitric Oxide Society* 2017; 71: 57-68.
2. Kleinbongard P, Dejam A, Lauer T, Rassaf T, Schindler A, Picker O, Scheeren T, Godecke A, Schrader J, Schulz R, Heusch G, Schaub GA, Bryan NS, Feelisch M, Kelm M. Plasma nitrite reflects constitutive nitric oxide synthase activity in mammals. *Free radical biology & medicine* 2003; 35(7): 790-796.
3. Rassaf T, Bryan NS, Kelm M, Feelisch M. Concomitant presence of N-nitroso and S-nitroso proteins in human plasma. *Free radical biology & medicine* 2002; 33(11): 1590-1596.
4. Benzie IF, Strain JJ. The ferric reducing ability of plasma (FRAP) as a measure of "antioxidant power": the FRAP assay. *Analytical biochemistry* 1996; 239(1): 70-76.
5. Zeb A, Ullah F. A Simple Spectrophotometric Method for the Determination of Thiobarbituric Acid Reactive Substances in Fried Fast Foods. *Journal of analytical methods in chemistry* 2016; 2016: 9412767.
6. Banne AF, Amiri A, Pero RW. Reduced level of serum thiols in patients with a diagnosis of active disease. *Journal of anti-aging medicine* 2003; 6(4): 327-334.
7. Frenay AS, de Borst MH, Bachtler M, Tschopp N, Keyzer CA, van den Berg E, Bakker SJL, Feelisch M, Pasch A, van Goor H. Serum free sulfhydryl status is associated with patient and graft survival in renal transplant recipients. *Free radical biology & medicine* 2016; 99: 345-351.
8. Koning AM, Meijers WC, Pasch A, Leuvenink HGD, Frenay AS, Dekker MM, Feelisch M, de Boer RA, van Goor H. Serum free thiols in chronic heart failure. *Pharmacological research* 2016; 111: 452-458.
9. Thijssen DH, Black MA, Pyke KE, Padilla J, Atkinson G, Harris RA, Parker B, Widlansky ME, Tschakovsky ME, Green DJ. Assessment of flow-mediated dilation in humans: a methodological and physiological guideline. *Am J Physiol Heart Circ Physiol* 2011; 300(1): H2-12.
10. Rodriguez-Miguel P, Seigler N, Harris RA. Ultrasound Assessment of Endothelial Function: A Technical Guideline of the Flow-mediated Dilation Test. *Journal of visualized experiments : JoVE* 2016(110).
11. Schindelin J, Arganda-Carreras I, Frise E, Kaynig V, Longair M, Pietzsch T, Preibisch S, Rueden C, Saalfeld S, Schmid B, Tinevez JY, White DJ, Hartenstein V, Eliceiri K, Tomancak P, Cardona A. Fiji: an open-source platform for biological-image analysis. *Nat Methods* 2012; 9(7): 676-682.